Role of amygdala and hippocampus in the neural circuit subserving conditioned defeat in Syrian hamsters

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We examined the roles of the amygdala and hippocampus in the formation of emotionally relevant memories using an ethological model of conditioned fear termed conditioned defeat (CD). Temporary inactivation of the ventral, but not dorsal hippocampus (VH, DH, respectively) using muscimol disrupted the acquisition of CD, whereas pretraining VH infusions of anisomycin, a protein synthesis inhibitor, failed to block CD. To test for a functional connection between the VH and basolateral amygdala (BLA), we used a classic functional connectivity design wherein injections are made unilaterally in brain areas either on the same or opposite sides of the brain. A functional connection between the BLA and VH necessary for the acquisition of CD could not be found because unilateral inactivation of either BLA alone (but not either VH alone) was sufficient to disrupt CD. This finding suggested instead that there may be a critical functional connection between the left and right VH. In our final experiment, we infused muscimol unilaterally in the VH and assessed Fos immunoreactivity on the contralateral side following exposure to social defeat. Inactivation of either VH significantly reduced defeat-induced Fos immunoreactivity in the contralateral BLA. These experiments demonstrate for the first time that whereas the VH is necessary for the acquisition of CD, it does not appear to mediate the plastic changes underlying CD. There also appears to be a critical interaction between the two VHs such that bilateral activation of this brain area must occur in order to support fear learning in this model, a finding that is unprecedented to date.

Our laboratory has taken a novel approach to examine the behavioral and physiological changes that accompany social experience by studying a striking behavioral response that is exhibited following social defeat. When a Syrian hamster is paired with a larger, more aggressive opponent and is defeated, it subsequently becomes highly submissive and fails to defend its own home cage even against a smaller, nonaggressive intruder. We call this change in the behavior of the defeated hamster conditioned defeat (CD) (Portegal et al. 1993) and believe that it is a valuable model with which to study neural and behavioral plasticity following exposure to a biologically relevant stressor.

One of the critical structures subserving CD is the amygdala; temporary inactivation of its major subnuclei, including the basolateral amygdala (BLA), blocks the acquisition of CD (Jasnow and Huhman 2001). Together with the findings that protein synthesis inhibition in the BLA effectively disrupts CD (Markham and Huhman 2008) and that overexpression of cAMP response element binding protein (CREB) in the BLA enhances CD (Jasnow et al. 2005), the data support the hypothesis that the BLA is a critical site for plasticity related to CD.

One brain region that we have largely overlooked, but which receives considerable attention for its role in learning and memory, is the hippocampus. Several groups have now gathered anatomical and behavioral data demonstrating functionally specific dissociation between the dorsal (DH) and ventral (VH) regions of the hippocampus (Risold and Swanson 1996; Moser and Moser 1998; Bannerman et al. 2004; McEown and Treit 2009). While the DH is critical for spatial relationships (O’Keefe and Nadel 1978; Moser et al. 1993; Eichenbaum 1996) and has been shown to play an important role in social recognition in hamsters (Lai et al. 2005), the VH appears to be involved in the production of behaviors produced in response to aversive stimuli (Trivedi and Coover 2004; Pentkowski et al. 2006).

Considering how critically important the hippocampus and amygdala are in relation to fear and memory, some studies are beginning to suggest that these areas may functionally interact to modulate memory function (Akirav and Richter-Levin 2002; McGaugh et al. 2002; McGaugh 2004; Vouimba et al. 2007). The BLA projects to the hippocampus (Amaral and Insausti 1992), and high-frequency stimulation of the BLA combined with tetanic stimulation of the perforant pathway facilitates hippocampal long-term potentiation (LTP) (Ikegaya et al. 1996). Additionally, electrolytic lesions of the VH produce a deficit in the acquisition of fear to a contextual conditioned stimulus, and NMDA lesions of the BLA cause a nonselective deficit in the acquisition of fear to both contextual and acoustic conditioned stimuli (Maren and Fanselow 1995). Although our laboratory has previously demonstrated that the BLA is critically involved in the acquisition of CD (Jasnow and Huhman 2001), the role of the hippocampus has yet to be investigated. The aim of the present study was to examine whether the VH and DH are involved in mediating CD and also to determine whether there is a functional interaction between the hippocampus and the amygdala in the acquisition of CD.

Results

Histological analysis

Figure 1A shows the cannula placements for animals infused with muscimol or vehicle into the DH (Experiment 1). Location of the needle tips was confined primarily to the dorsal regions of the DH. However, five animals were removed from the study due to misplaced or occluded cannula. Figure 1B shows the location of the cannula for animals receiving bilateral infusions into the VH (Experiments 2 and 3). Needle tips were confined to the CA1, CA2,
and CA3 subregions of the VH, while avoiding the dentate gyrus (DG) as well as the polymorph layer of the DG. Animals with misplaced or occluded cannula (nine in Experiment 2; two in Experiment 3) were removed from the study. Finally, Figure 2 shows the placements of the cannula aimed at the VH (Fig. 2A) and BLA (Fig. 2B; Experiments 4 and 5). Subjects with misplaced (n = 17; VH = 5; BLA = 12), damaged (n = 6), or occluded (n = 4) cannula were not included in the statistical analysis.

**Experiment 1: Muscimol infused into the DH does not reduce the acquisition of conditioned defeat**

A total of 36 animals were used in the statistical analysis: vehicle (n = 9), 1.1 nmol (n = 7), 2.2 nmol (n = 12), and 3.3 nmol (n = 8). Infusion of muscimol into the DH immediately before defeat did not reduce the display of submissive behaviors (F(3,31) = 0.072; P > 0.05) (Fig. 3). In addition, there were no significant differences observed in aggressive, social, and nonsocial behaviors (Fig. 3). Information about animals dropped from each experiment is given in the Histological analysis section. Additionally, no animals had to be removed from any experiment due to injury.

**Experiment 2: Muscimol infused into the VH reduces the acquisition of conditioned defeat**

A total of 32 animals were used in the statistical analysis: vehicle (n = 8), 1.1 nmol (n = 8), 2.2 nmol (n = 9), and 2.7 nmol (n = 7). Infusion of muscimol immediately before training significantly reduced the display of submissive behaviors during subsequent testing (F(3,31) = 4.096; P < 0.05) (Fig. 4) without altering the behavior of either the resident aggressors or subjects during training. Post-hoc analysis showed that infusion of muscimol into the VH reduced submissiveness during testing at three doses when compared with animals injected with vehicle (P < 0.05 for each comparison). Infusion of muscimol also significantly increased nonsocial behavior at all three doses, which consisted of normal locomotor activity and grooming (F(3,31) = 7.895; P < 0.05). Finally, there were no significant differences in aggressive and social behaviors, indicating that the inhibition of submissive behavior by muscimol was not due to a nonspecific decrease in agonistic or social behaviors.

**Experiment 3: Anisomycin infused in the VH does not reduce the acquisition of conditioned defeat**

A total of 30 animals were used in the statistical analysis: vehicle (n = 12) and anisomycin (n = 18). As shown in Figure 5, infusion of anisomycin 30 min prior to defeat did not cause a significant reduction in submissive behaviors when animals were tested against a nonaggressive intruder (NAI) 24 h later (F(1,29) = 0.395; P > 0.05). This dose of anisomycin was identical to that used in previous studies in which this concentration was effective in impairing contextual fear conditioning in the hippocampus (Rudy and Matus-Amat 2005) and acquisition of CD in the BLA (Markham and Huhman 2008).

**Experiment 4: Inactivation of the BLA-VH circuit inhibits the acquisition of conditioned defeat**

A total of 66 animals were used in the statistical analysis. Because there was no evidence of laterality (no significant differences between injections of the left and right hemispheres), groups were collapsed over side of injection and combined for subsequent analysis: Control Group: vehicle in contralateral BLA and VH (n = 11); Ipsilateral Group: muscimol in BLA and VH in the same hemisphere (n = 13); Contralateral Group: muscimol in contralateral BLA and VH (n = 12); Unilateral BLA Group: muscimol on one side of the BLA and vehicle in the contralateral VH (n = 14); and Unilateral VH Group: muscimol on one side of the VH and vehicle in the contralateral BLA (n = 16).

There was a significant effect of treatment on the display of submissive (F(4,61) = 5.754; P < 0.001) and social (F(4,61) = 9.795; P < 0.0001) behaviors during testing with the NAI 24 h later (Fig. 6). Treatment did not affect the initial defeat training experience as indicated by the fact that the durations of aggressive, social, and nonsocial behavior of the resident aggressor toward the experimental
animal were similar among the groups regardless of treatment condition (data not shown). A Tukey-Kramer multiple comparison post-hoc test indicated that unilateral inactivation of the BLA, regardless of group (or other area injected), significantly reduced the duration of submissive behaviors compared to vehicle and VH-only inactivation groups ($P < 0.01$). No instances of aggression toward the NAI were observed, and there were no significant differences among groups in nonsocial behaviors.

**Experiment 5: Unilateral inactivation of BLA suppresses Fos activity in contralateral BLA**

A total of 16 animals were used in the statistical analysis: Vehicle Defeat ($n = 5$), Muscimol Defeat ($n = 6$), and Muscimol No Defeat ($n = 5$). Because there were no significant differences in Fos immunoreactivity (Fos-IR) based on the side of muscimol injection (left or right BLA), these groups were combined. As shown in Figure 7, there was a significant effect of treatment on Fos-IR in the contralateral BLA ($F_{2,13} = 24.66; P < 0.0001$). Post-hoc tests showed that both the Muscimol/No Defeat and Muscimol Defeat groups had significantly lower levels of Fos-positive cells compared to the Vehicle/Defeat animals in the BLA opposite the injection site. No significant differences were found between the Muscimol/No Defeat and Muscimol/Defeat groups.

In order to ensure that muscimol infusions alone did not cause a global or nonspecific inhibition of neural activity, Fos-positive cells were also counted in brain areas in close proximity to the BLA, including the CeA, MeA, BNST, VMH, LSv, and the Arcuate nucleus. Table 1 shows that aside from the BLA and LSv, no other structure showed significant differences in the number of Fos-positive cells when comparing the Saline/Defeat and Muscimol/Defeat groups.

**Discussion**

These experiments demonstrate that the ventral hippocampus is a component of the neural circuit subserving CD in that temporary inactivation of the ventral but not the dorsal hippocampus blocks the acquisition of conditioned defeat. Additionally, it first appeared that disrupting the VH–BLA pathway also significantly impairs the acquisition of CD; however, this attenuation appears to be due to the fact that unilaterally inactivating the BLA alone (originally performed as a control group) is sufficient to inhibit the acquisition of CD. This finding suggests the possibility that there is functional connectivity between the left and right BLA such that neural activity in both nuclei must occur in order to mediate the behavioral plasticity associated with a defeat experience. We began to examine this possibility in our final experiment wherein we demonstrated that unilateral inactivation of the BLA suppresses Fos-immunoreactivity (Fos-IR) in the contralateral BLA following social defeat. Our findings thus provide strong evidence to support the hypothesis that there is a functional connection between the left and right BLA such that inactivating one side of the BLA effectively inhibits the other.

The data from a recent study by Lai and colleagues (2005) showing that temporary inactivation of the DH attenuates the
avoidance by defeated hamsters of an animal that had previously defeated them led us to hypothesize that the DH would also be involved in CD. Instead, the results of Experiment 1 showed that muscimol infusion in the DH fails to inhibit the acquisition of CD, suggesting that the DH is not a critical component of the neural circuit mediating conditioned defeat. In the Lai et al. (2005) study, hamsters were exposed to the familiar, dominant opponent. In contrast, in our studies the stimulus animal (the nonaggressive intruder) is very different in size, age, and behavior from the resident aggressor used during the initial defeat training. Thus, it appears that the DH may be involved in behavioral responses to a particular dominant opponent, but it does not appear to be involved in the acquisition of generalized behavioral changes that are produced in response to a defeat experience, per se. The differences between the two models may therefore explain why inactivation of the DH failed to inhibit the acquisition of CD. We do agree that the DH is an integral component of fear conditioning circuitry subserving some of the more traditional models, particularly wherein specific context cues are relevant. Another possible explanation for our failure to observe a role for the DH is that muscimol infusions may have caused an increase in general activity levels during defeat and that this increase interfered with the normal pattern of defeat. Indeed, damage to the hippocampus has been shown in some studies (Yoon and Otto 2007) to increase locomotor activity. Careful analysis of the defeat sessions, however, revealed no such locomotor changes.

Our finding that infusion of muscimol into the VH inhibits the acquisition of CD is consistent with several other studies showing that this area is important in the production of anxiety-like behaviors. For example, electrolytic or neurotoxic lesions of the VH have been shown to have anxiolytic effects, including a reduction in the avoidance of the open arms of an elevated T-maze (Trivedi and Coover 2004), reduced avoidance of a male conspecific in the social interaction test (Deacon et al. 2002), reduced anxiety-like behaviors in a social interaction test (McHugh et al. 2004), and an impairment of risk assessment behaviors (a behavioral index of anxiety) in rats confronted with an ethologically relevant threat stimulus (Pentkowski et al. 2006). In addition to its role in regulating anxiety-like behaviors, a number of recent studies have demonstrated that the VH may also be critically involved in certain forms of conditioned fear (Rogers et al. 2006; Yoon and Otto 2007; Czerniawski et al. 2009). The study by Czerniawski et al. (2009) demonstrated a double-dissociation between the DH and VH as temporary inactivation of the ventral, but not dorsal, hippocampus blocked the acquisition and expression of trace fear conditioning, whereas inactivation of the dorsal, but not ventral, hippocampus impaired the behavioral response to a delayed reinforced alternation task.

Although the previously mentioned findings, combined with our own work, provide strong evidence for the involvement of the VH in conditioned fear responses, few if any studies have specifically examined whether actual plasticity and memory consolidation occurs in the VH. Although there has been some recent controversy concerning the use of protein synthesis inhibitors to disrupt learning (Gold 2008), this technique is still widely used and accepted. We have previously demonstrated that infusion of anisomycin into the BLA blocks memory formation necessary for CD (Markham and Huhman 2008). Infusion of this same dose of anisomycin into the VH in the present study was ineffective. The fact that we found anisomycin to be effective in the BLA but not the VH argues against a nonspecific effect of anisomycin on the well-being and function of neurons explaining the effectiveness of the treatment in blocking memory formation for CD. Together, our data suggest that while the VH is an integral component of the neural circuit mediating the acquisition of CD, actual plasticity related to the defeat experience likely occurs elsewhere (e.g., the BLA). Indeed, the VH has reciprocal connections to multiple brain regions that are critical in fear, anxiety, and stress responsivity (Moser and Moser 1998; Bast et al. 2001; Petrovich et al. 2001) such as the amygdala. We must also acknowledge, however, the possibility that the VH is also a critical site for plasticity underlying CD but that this plasticity is not dependent on protein synthesis.

We were initially surprised at the finding in Experiment 4 that temporary inactivation of the VH–BLA circuit in both the ipsilateral...
and contralateral groups caused a significant reduction in the acquisition of CD. This suggested to us that the projections between these structures may, in fact, be both ipsilateral and contralateral instead of mainly ipsilateral, which had been suggested in the literature (Packard et al. 1994). In fact, it appears that this surprising observation can actually be explained by the equally unexpected finding that unilateral inactivation of the BLA (but not VH) significantly inhibits the acquisition of CD. Several key conclusions can be based on these results. First, these findings indicate that, at least with the conditioned defeat model, the left and right VH can function independently to support learning while the left and right BLA cannot. Second, there does not appear to be any laterality of function within the hamster amygdala, or, alternatively, the left and right BLA are connected such that any laterality of function is masked. Indeed, there is some evidence for this connection, and it has been suggested that it occurs via the anterior commissure, at least in rats (Martinez-Lorenzana et al. 2004). Our final experiment sought to indirectly test the existence of this connection by unilaterally inactivating one BLA and examining Fos-IR in the contralateral BLA. As reported above, this inactivation caused a suppression of Fos opposite to the side of the infusion, suggesting that unilateral inactivation of the BLA is sufficient to disrupt CD because it also inhibits activation of the contralateral BLA. Importantly, this suppression was specific to the BLA, as Fos counts in the neighboring regions in muscimol-infused animals (with the exception of the ventral lateral septum; LSV) were not significantly different from the counts obtained from vehicle-infused hamsters. These findings seem to contradict previous reports showing that conditioned fear can be supported even after unilateral lesions of the amygdala (LaBar and LeDoux 1996; Goosens and Maren 2001). There are other examples, however, where bilateral activation of a brain region is required for the expression of a particular behavior. For example, Ferris and colleagues (1994) demonstrated that bilateral activity of the anterior hypothalamus (AH) is critical for the expression of flankmarking behavior in hamsters.

In conclusion, the results of the present series of experiments, combined with previous findings from our laboratory (Jasnow and Huhman 2001; Huhman et al. 2003; Jasnow et al. 2005; Markham and Huhman 2008), indicate that while several neural sites are important mediators of CD, the BLA appears, thus far, to be the key site wherein plasticity related to the defeat experience occurs. Additionally, we provide strong indirect evidence that the left and right BLA form a functional unit that is necessary for the mediation of CD.

Table 1. Counts of Fos-positive cells

| Region | Bregma (mm) | Saline/Defeat (n = 5) | Muscimol/Defeat (n = 6) | Muscimol/No Defeat (n = 5) |
|--------|-------------|-----------------------|------------------------|--------------------------|
| BLA    | −1.50       | 167 ± 24              | 75 ± 9<sup>a,c</sup>   | 28 ± 9<sup>b</sup>       |
| CeA    | −1.20       | 28 ± 7                | 23 ± 6                 | 2 ± 0.9<sup>b</sup>      |
| MeApv  | −1.50       | 245 ± 60              | 193 ± 16               | 80 ± 12<sup>b</sup>      |
| MeApd  | −1.50       | 130 ± 43              | 93 ± 8                 | 28 ± 5<sup>b</sup>       |
| BNSTpm | +0.5        | 83 ± 12               | 67 ± 14                | 26 ± 5<sup>b</sup>       |
| BNSTpi | +0.5        | 109 ± 22              | 96 ± 24                | 28 ± 5<sup>b</sup>       |
| LSV    | +0.9        | 396 ± 60              | 290 ± 55               | 108 ± 41<sup>b</sup>     |
| VMHdm  | +2.0        | 83 ± 17               | 67 ± 7                 | 24 ± 8<sup>b</sup>       |
| Arc    | −2.0        | 116 ± 43              | 103 ± 10               | 40 ± 8<sup>b</sup>       |

The region of interest was counted using the adjoining Nissl-stained sections.

<sup>a</sup> p < 0.05 versus saline/defeat animals.
<sup>b</sup> p < 0.01 versus muscimol/defeat animals.
<sup>c</sup> p < 0.01 versus muscimol/no defeat animals.

Summary of Fos immunoreactivity in select brain regions. Data show the mean number of c-Fos immunoreactive cells located in representative sections of each brain region (SEM). The region of interest was counted using adjoining the Nissl-stained sections at 20X magnification. Significant differences are indicated by unshared letters.

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**Figure 7.** Photomicrographs of Fos-immunoreactive cells (dark dots) in the basolateral amygdala (BLA) and surrounding areas in the (A) saline/defeat, (B) muscimol/defeat, and (C) muscimol/no defeat groups.
Materials and Methods

Animals and housing conditions
Male Syrian hamsters (Mesocricetus auratus) weighing 120–140 g at the beginning of the experiment were obtained from Charles River Laboratories (Kingston, NY). Older males (>6 mo) weighing 160–180 g were housed individually and used as resident aggressors during the defeat training. Younger males (2 mo) that weighed 100–110 g were group housed (five to six hamsters per cage) and used as nonaggressive intruders during CD testing. All hamsters were housed in polycarbonate cages (20 × 40 × 20 cm) with wire mesh tops, and food and water were available ad libitum. Animals were kept in a temperature- and humidity-controlled room with a 14:10-h light-dark cycle (lights out at 1100 h). All procedures and protocols were approved by the Georgia State University Institutional Animal Care and Use Committee, and all methods were in accordance with the standards outlined in the National Institutes of Health Guide for Care and Use of Laboratory Animals.

Surgical procedures
Hamsters were deeply anesthetized with sodium pentobarbital (90 mg/kg, i.p.) and placed into a stereotaxic frame. All animals were implanted with 4-mm, 26-gauge stainless steel cannula (Plastics One). Lambda and bregma were leveled prior to placement of the cannula. Guide cannula were bilaterally implanted (with the exception of Experiments 4 and 5) and aimed at the DH (Experiment 1), VH (Experiments 2 and 3), the BLA and VH (Experiment 4), or the BLA (Experiment 5). Following surgery, dummy styllets were placed in the guide cannula to help maintain patency. Hamsters were allowed 7–10 d to recover from surgery prior to the start of behavioral testing. They were housed each day after surgery by gently restraining them and removing and replacing the dummy styllet in order to habituate the subjects to the injection procedure.

Drug infusion
Infusions into the DH, VH, and BLA were administered to freely moving hamsters over 2 min with a Hamilton syringe mounted on a syringe pump (Harvard Apparatus) connected to a 33-gauge injection needle via polyethylene tubing (Fisher Scientific). The needle was kept in place for an additional 1 min before being removed to ensure complete diffusion of the drug after which the dummy styllet was replaced. Hamsters were infused with either the GABA_A agonist muscimol (Sigma; Experiments 1, 2, 4, and 5), the protein synthesis inhibitor anisomycin (Sigma; Experiment 3), or vehicle (physiological saline; used as a control in all experiments).

Social defeat and testing
Subjects were tested in the conditioned defeat model, described in detail elsewhere (Huhman et al. 2003). All training and testing sessions were performed during the first 3 h of the dark phase of the light–dark cycle to minimize circadian variation in the behavior and physiology. Training consisted of one 15-min exposure to the resident aggressor (RA) in the aggressor's home cage, upon which the RA reliably attacked the experimental hamsters within 60 sec. We employ a caveat wherein any hamster that is bitten by the RA reliably attacked the experimental hamsters within 60 sec. We employ a caveat wherein any hamster that is bitten is removed from the study and is examined by a veterinarian, but none of the subjects were excluded for this reason. The following day, animals were paired in their own home cage for 5 min with a nonaggressive intruder (NAI). An animal was considered to show conditioned defeat if it exhibited an increase in submissive and defensive behaviors and no aggressive behaviors when the NAI was introduced into its home cage.

Behavioral analysis
All training and testing sessions were videotaped via a camera mounted overhead. The video files were scored by an experimentally blind observer using the behavioral scoring program Hindsight (developed by Dr. Scott Weiss). The total duration of four classes of behavior were scored during the test session: (1) social behavior (stretch, approach, sniff, nose touching, and flank marking); (2) nonsocial behavior (locomotion, exploration, grooming, nest building, and sleeping); (3) submissive/defensive behaviors (flight, avoidance, tail up, upright, side defense, full submissive posture, stretch attend, head flag, attempted escape from cage); and (4) aggressive behaviors (upright and side offense, chase and attack, including bite). For all experiments, the behavior of the resident aggressors and subjects was also scored during training to ensure that the presence of a drugged subject during testing did not alter the behavior of the resident aggressor; (2) all animals received similar defeats; and (3) subjects were able to produce normal submissive behavior in response to attack.

Site verification
At the end of Experiments 1–4, hamsters were administered an overdose of sodium pentobarbital and infused with either 200 nl (BLA) or 300 nl (DH and VH) of India ink to verify the placement of the needle. The brains were removed and post-fixed in 10% buffered formalin for 3 d before being sectioned on a cryostat. Thirty-micrometer sections were taken and stained with neutral red and coverslipped with DPX (VWR International Ltd.). Sections were then examined under a light microscope for placement verification. Only animals with injection sites within 0.3 mm of the target structure were included in the statistical analysis.

Immunohistochemistry
In Experiment 5, animals were administered an overdose of sodium pentobarbital 60 min after the end of defeat and were infused with 200 nl of India ink to verify the placement of the needle. Subjects were then perfused transcardially with 0.1 M phosphate-buffered saline (PBS) and 4% paraformaldehyde. Brains were post-fixed in 4% paraformaldehyde overnight and switched to a 30% sucrose-paraformaldehyde solution and kept at 4°C for an additional 48 h. Brains were blocked on the coronal plane and sectioned on a cryostat to a thickness of 30 μm. Consecutive sections were placed sequentially across four vials filled with cyprotestant. The sections covered the areas between the anterior bed nucleus of the stria terminals (BNST) and the posterior division of the BLA. Sections were labeled for c-fos using primary antiserum directed against the protein product of the immediate early gene c-fos (rabbit anti-c-fos polyclonal antibody, 1:20000; Santa Cruz Biotechnology). All washes, rinses, and incubations were performed in 12-well tissue culture plates, which were gently shaken on an orbital shaker throughout the immunostaining procedure. Briefly, sections were rinsed in 0.1 M PBS and incubated in a 1% hydrogen peroxide solution for 15 min, followed by a PBS wash. Sections were then incubated for 1 h in the primary antibody at room temperature, and then for 48 h at 4°C. Following incubation with the primary antibody, the sections were rinsed in PBS and incubated for 1 h in a 0.4% solution of Triton X-100 in 0.1 M PBS containing the secondary antibody (biotinylated anti-rabbit IgG polyclonal antibody, 1:600; Vector Laboratories). They were then rinsed with PBS, followed by incubation for 1 h with an avidin-biotin complex reagent (Vectastain Elite ABC kit; Vector Laboratories). After rinsing with PBS, and sodium acetate solution, sections were incubated in a nickel-3,3'-diaminobenzidine (DAB) solution for 10 min in order to produce a blue-black-stained product and then washed again with sodium acetate and PBS in order to halt the DAB reaction. Finally, the sections were mounted on gelatin-coated slides, dehydrated, and coverslipped with DPX (VWR International).

Cell counting
An observer blind to the treatment conditions performed all cell counts using brightfield microscopy at 10–20× magnification. For the quantification of Fos labeling, areas of interest were first delineated using adjoining Cresyl violet-stained sections. Fos-labeled cells were counted within this standardized template for each brain region. The following areas were counted based on the hamster atlas of Morris and Wood (2001); BLA, medial amygdala (Ma), central amygdala (CeA), ventral lateral septum (LSv), the postero medial and postero intermediate bed nucleus of the stria
neuromodulatory terminals (BNSTp and BNSTp, respectively), ventromedial hypothalamus (VMH), and the arcuate nucleus (Arc).

Experiments 1 and 2: DH/VH acquisition of conditioned defeat

The purpose of Experiments 1 and 2 was to determine whether infusion of muscimol in the DH or VH, respectively, would reduce the acquisition of CD. Animals (Experiment 1: \( n = 41 \); Experiment 2: \( n = 41 \)) were randomly assigned to groups. Hamsters received either muscimol (1.1, 2.2, 2.7, or 3.3 nmol in 300 nL of saline) or vehicle (physiological saline) into the DH or VH immediately before being placed in the cage of a resident aggressor for 15 min. This range of dosages of muscimol is effective in blocking CD when given in the amygdala (Markham and Huhman 2008). On the following day, animals were tested for 5 min in their own home cage in the presence of a nonaggressive intruder, as described above.

Experiment 3: VH/protein synthesis inhibition

The aim of Experiment 3 was to determine whether the inhibition of protein synthesis via infusion of anisomycin in the VH before training would block the acquisition of CD. Animals (\( n = 32 \)) were again randomly assigned to groups after being weight matched. Anisomycin (125 \( \mu \)g/mL) or vehicle was infused in hamsters 30 min prior to initial defeat and tested with a resident aggressor for 15 min. The use of anisomycin given into the BLA blocks the acquisition of CD in hamsters (Markham and Huhman 2008). On the following day, animals were tested for 5 min in their own home cage in the presence of a nonaggressive intruder.

Experiment 4: VH/BLA acquisition of conditioned defeat

The purpose of Experiment 4 was to determine whether there is a functional interaction between the BLA and VH in the acquisition of CD. Animals (\( n = 93 \)) were randomly assigned to groups and infused with muscimol (2.2 nmol in 300 nL of saline in each injection site) or vehicle (physiological saline) 5 min before being placed in the cage of a resident aggressor for 15 min. On the following day animals were tested for 5 min in their own home cage in the presence of a nonaggressive intruder. The animals were divided into five major groups that were then balanced for side of injection: Control (contralateral saline infusions in either the left BLA and left VH), Contralateral (muscimol infusions in either the left BLA and left VH or right BLA and right VH), Contralateral (muscimol infusions in either the left BLA and right VH or right BLA and left VH), Unilateral BLA (muscimol infusion in either the right or left BLA and saline in the right or left VH), and Unilateral VH (muscimol infusion in either the right or left VH and saline in the right or left BLA).

Experiment 5: Unilateral muscimol BLA/Fos expression

The purpose of Experiment 5 was to further examine the novel possibility that there is a functional connection between the right and left BLA that is critical for the formation of CD. Animals (\( n = 18 \)) were randomly assigned to groups and unilaterally infused with muscimol (2.2 nmol in 200 nL of saline) or vehicle in the left or right BLA. Fifteen minutes after the injection, subjects were placed into the home cage of a resident aggressor and were defeated for 15 min (defeat group) or left undisturbed in their own home cages for the same amount of time (undefeated group). All animals were then left undisturbed for 1 h after which they were injected with an overdose of sodium pentobarbital (90 mg/kg i.p.) and transecradially perfused. Brains were removed and stored in 10% paraformaldehyde for 48 h after which they were cut into 30-μm sections on a cryostat.

Statistical analysis

The total duration (seconds) of each behavior displayed (Submissive/Defensive, Social, Nonsocial) was determined, and the mean total duration of each behavior was then compared using analysis of variance (ANOVA) with dose as the between-subjects factor. Statistically significant differences were further analyzed using a Tukey-Kramer multiple comparison post-hoc test to compare all pairwise differences among group means. The immunohistochemistry data were analyzed using two-way between-subjects ANOVAs, with drug injection and stimulus exposure as independent factors. Significant differences for all analyses were set at \( P < 0.05 \).

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