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Alcohol-Seeking Triggered by Discrete Pavlovian Cues is Invigorated by Alcohol Contexts and Mediated by Glutamate Signaling in the Basolateral Amygdala

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The environmental context in which a discrete Pavlovian conditioned stimulus (CS) is experienced can profoundly impact conditioned responding elicited by the CS. We hypothesized that alcohol-seeking behavior elicited by a discrete CS that predicted alcohol would be influenced by context and require glutamate signaling in the basolateral amygdala (BLA). Male, Long-Evans rats were allowed to drink 15% ethanol (v/v) until consumption stabilized. Next, rats received Pavlovian conditioning sessions in which a 10 s CS (15 trials/session) was paired with ethanol (0.2 ml/CS). Entries into a port where ethanol was delivered were measured. Pavlovian conditioning occurred in a specific context (alcohol context) and was alternated with sessions in a different context (non-alcohol context) where neither the CS nor ethanol was presented. At test, the CS was presented without ethanol in the alcohol context or the non-alcohol context, following a bilateral microinfusion (0.3 μ l/hemisphere) of saline or the AMPA glutamate receptor antagonist NBQX (2,3-dioxo-6-nitro-1,2,3,4-tetrahydrobenzo[f]quinoxaline-7-sulfonamide disodium salt) in the BLA (0, 0.3, or 1.0 μ g/0.3 μ l). The effect of NBQX (0, 0.3 μ g/0.3 μ l) in the caudate putamen (CPu) on CS responding in the non-alcohol context. NBQX in the BLA reduced CS responding in both contexts but had no effect in the CPu. These data indicate that AMPA glutamate receptors in the BLA are critical for alcohol-seeking elicited by a discrete CS and that behavior triggered by the CS is strongly invigorated by an alcohol context.

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INTRODUCTION

The seminal hypothesis that environmental contexts associated with drugs of abuse influence drug use, abuse, and relapse (Hinson *et al*, 1986; Siegel, 1988; Powell, 1995) has gained renewed traction from emerging preclinical evidence that drug-associated contexts impact drug-seeking behavior (Crombag and Shaham, 2002; Zironi *et al*, 2006; Chaudhri *et al*, 2008; Hamlin *et al*, 2008). We have reported that Pavlovian alcohol-seeking, defined herein as alcohol-seeking behavior elicited by a discrete conditioned stimulus (CS) that predicts alcohol, is invigorated in a context where alcohol was previously consumed, relative to a context where alcohol intake had never occurred (Remedios *et al*, 2014). These results may, however, have been attributable to a longer history of exposure to the alcohol context. The first objective of the present research was to address this potential limitation in a new behavioral assay developed to examine the impact of context on Pavlovian alcohol-seeking.

Environmental cues that predict alcohol can have profound and lasting effects on behavior (Ciccocioppo *et al*, 2001; Veilleux and Skinner, 2015). For example, sensory stimuli associated with alcohol can become established as Pavlovian conditioned stimuli (Field and Duka, 2002) and trigger physiological, behavioral, and psychological responses in abstinent individuals (Cooney *et al*, 1984; Field and Duka, 2002). Craving for alcohol is one such conditioned response, and levels of cue-induced alcohol craving measured in the laboratory predict the likelihood of relapse in abstinent alcoholics (Litt *et al*, 2000; Sinha, 2011). Such findings highlight a critical link between cue-induced craving and relapse—yet the neural mechanisms that engender and maintain this link are poorly understood.

The glutamatergic system has been strongly implicated in cue-induced, alcohol-seeking behavior. Systemic injections of either ionotropic (Backstrom and Hyytia, 2004b) or metabotropic (Backstrom *et al*, 2004a) glutamate receptor antagonists reduce cue-induced reinstatement of responding on an

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alcohol-associated lever in rats. In addition, enhancing the activity of α -amino-3-hydroxy-5-methyl-4-isoxazolepropionic acid (AMPA) glutamate receptors facilitates operant alcohol self-administration and cue-induced reinstatement of alcohol-seeking behavior (Cannady et al, 2013). Topiramate, a drug that acts partly through blocking AMPA receptors, also decreases drinking in alcohol-dependent people (Johnson et al, 2007), suggesting that AMPA receptors may be an important new target for the treatment of alcohol use disorders.

Glutamate in the basolateral amygdala (BLA) appears to have an essential role in alcohol-seeking behaviors that were acquired through instrumental conditioning and reinforced by stimuli that were previously paired with alcohol. For example, cue-induced reinstatement of operant alcoholseeking, but not food-seeking behavior, is associated with an increase in BLA glutamate (Gass et al, 2011), and blocking mGluR5 glutamate receptors in the BLA attenuates cueinduced reinstatement of alcohol-seeking (Sinclair et al, 2012). These findings suggest that BLA glutamate is needed to promote relapse-like behavior after a period of abstinence when responding is not reinforced by alcohol or associated cues. Less is known about processes that underlie conditioned responses elicited by Pavlovian cues that predict alcohol.

The BLA has an established role in the formation and expression of Pavlovian associations between conditioned and unconditioned stimuli (Wilensky et al, 1999; Meil and See, 1997; Cardinal et al, 2002). Interestingly, this region has also been implicated in alcohol-seeking triggered by conditioned stimuli that predict alcohol (Chaudhri et al, 2013; Marinelli et al, 2010; Gremel and Cunningham, 2008). Within the BLA, AMPA receptors contribute to synaptic plasticity that underpins conditioned reinforcement (Mead and Stephens, 2003) and cue-reward learning (Tye et al, 2008). Consequently, we hypothesized that AMPA receptors in the BLA would be needed for the expression of alcoholseeking elicited by a discrete CS and anticipated that if this was a critical process then blocking AMPA receptors in the BLA would reduce alcohol-seeking elicited by a discrete CS in whichever context the CS was experienced.

In Experiment 1, we developed a new behavioral assay to assess the impact of alcohol-associated contexts on alcoholseeking elicited by a discrete CS that predicted alcohol. Experiment 2 examined the impact of bilateral microinfusions of the AMPA receptor antagonist NBQX (2,3-dihydroxy-6nitro-7-sulfamoyl-benzo[f]quinoxaline-2,3-dione)) in the BLA on alcohol-seeking elicited by a discrete CS in a context associated with alcohol intake (alcohol context) and in a context where alcohol had never previously been consumed (non-alcohol context). Experiment 3 investigated the possibility that the reduction in Pavlovian alcohol-seeking observed in Experiment 2 may have been attributable to upward diffusion into the caudate putamen (CPu).

MATERIALS AND METHODS

Subjects

Male, Long-Evans rats (220-275 g on arrival; Harlan Laboratories, Indianapolis, USA) were individually housed in polycarbonate cages in a controlled (21 °C) vivarium on a

12 h light-dark cycle (lights on at 0700 hours; procedures conducted during light phase) with unrestricted access to water and rat chow (Ralston Purina, Canada). Rats were weighed and handled for 7-10 days before behavioral procedures were initiated. Procedures were approved by the Animal Research Ethics Committee at Concordia University and concur with recommendations in the Guide for the Care and Use of Laboratory Animals (National Research Council, 1996).

Apparatus

Operant conditioning chambers and peripheral devices were obtained from Med-Associated (St Albans, VT, USA) and are described in detail elsewhere (Remedios et al, 2014; Sciascia et al, 2014).

Drugs and Solutions

Tap water was added to 95% ethanol to obtain a 15% ethanol solution (v/v). NBQX disodium (Abcam, Cambridge, MA, USA) was dissolved in sterile 0.9% sodium chloride (10 mg in 0.6 ml) to obtain a $5 \mu g/0.3 \mu l$ dose. Doses of $1.0 \mu g/0.3 \mu l$ and $0.3 \,\mu\text{g}/0.3 \,\mu\text{l}$ were made through serial dilution. Aliquots of each dose were stored at -20 °C until use. Doses were based on preliminary research and published studies (Walker and Davis, 1997). Odors for creating distinctive contexts (see below) were prepared by adding tap water to lemon oil (SAFC Supply Solutions, St Louis, MO, USA) or to benzaldehyde (almond odor; OMEGA Chemical Company, Levis, QC, Canada) to obtain 10% (v/v) solutions.

Home-Cage Ethanol Exposure

Rats were acclimated to ethanol in their home-cage using a 24-h, intermittent-access schedule (Wise, 1973; Simms et al, 2008; Sparks et al, 2014). Access to water was provided for 7 days per week. However, on Monday, Wednesday, and Friday a 100-ml graduated cylinder containing ethanol was placed on the lid of the home-cage. Before each session, ethanol cylinders, water bottles, and rats were weighed, and 24 h later, ethanol cylinders and water bottles were re-weighed to record intake. The placement of ethanol and water containers on different sides of the lid was alternated in each session. Spillage was accounted for by subtracting ethanol or water lost from bottles in empty cages from consumption during corresponding sessions.

Surgery

Upon completing home-cage ethanol exposure, rats in Experiments 2 and 3 were implanted with bilateral, 26-gauge guide cannulae (Plastics One, Roanoke, VA) targeting the BLA (AP: -2.8; ML: ± 5.1 ; DV -5.4) or CPu (AP: -2.8; ML: ± 5.1 ; DV -4), respectively. Standard stereotaxic procedures were used (Chaudhri et al, 2013). Guide cannulae were anchored to the skull using dental cement and metal screws. To maintain hydration, 1 ml of 0.9% sterile saline was injected subcutaneously (s.c.) 1 h into surgery and upon completion. Following surgery, rats received a single injection of buprenorphine (0.1 ml/kg, s.c.) and sweetened,

softened chow to encourage feeding. Weight gain was monitored over a 7-20-day recovery period.

Intracranial Microinfusions

Bilateral microinfusions were conducted in the behavior testing room using 33-gauge injectors (Plastics One, Roanoke, VA, USA) that extended below the guide cannula (Experiment 2; 3-mm protrusion, DV at injector tip = -8.4: Experiment 3; 2-mm protrusion, DV at injector tip = -6.0). The procedure for conducting microinfusions is described elsewhere (Chaudhri *et al*, 2013). A volume of 0.3 µl/hemi-sphere was infused at a rate of 0.3 µl/min, followed by 2 min for diffusion. Session onset occurred 5–25 min after the first microinfusion, based on the number of rats tested in a given cohort.

General Behavioral Procedures

Habituation. Rats were first individually handled in the behavioral testing room for 1 min. The next day, each rat was habituated a designated testing chamber for 20 min, during which a white chamber light located near the ceiling on the left wall of each chamber was illuminated and entries into a fluid port located 2 cm above the floor on the right wall of each chamber were recorded via interruptions of a photo beam situated across the entrance of the port. Each chamber was set up at Context 1, created by the addition of distinctive visual, olfactory, and tactile stimuli to the chamber. Specifically, Context 1 comprised of black walls, a smooth Plexiglas floor, and a lemon odor. This step was repeated 24 h later, at which time each chamber was set up as Context 2, which comprised of clear Plexiglas walls, a floor made up of 1-inch metal grids, and an almond odor. Odors were applied to the waste pan beneath chamber floors.

Behavioral training and test. A schematic illustration of the behavioral training and test sessions is provided in Figure 1a. Behavioral training (Monday-Friday; 73.5-min sessions) comprised of Pavlovian conditioning sessions in one context, alternated with sessions of exposure to the same chambers that were set up as a different context. In each Pavlovian conditioning session, the chamber light was illuminated 2 min after initiating the Med-PC program, to indicate session onset. Each session comprised of 15 trials of an auditory CS (10 s; continuous white noise) that occurred with an inter-trial interval (ITI) that varied around an average of 240 s (comprised of 180-, 240-, and 360-s intervals). This variable ITI did not include 10s intervals before (PreCS) or after (PostCS) each CS. At 4s after CS onset, 0.2 ml of ethanol was delivered over 6 s into the fluid port for oral consumption (Figure 1b). This short-delay conditioning procedure has been shown to produce reliable Pavlovian conditioning with ethanol (Chaudhri et al, 2008; Remedios et al, 2014). A total of 3 ml of ethanol was delivered per session, and ports were checked at the end of each session to ensure that it had been consumed. Pavlovian conditioning was conducted in Context 1 for half the rats and Context 2 for the remainder. Assignment to each context was counterbalanced according to ethanol intake averaged across the last three sessions of home-cage ethanol exposure.

The context in which Pavlovian conditioning sessions were conducted is referred to as the 'alcohol context'.

Pavlovian conditioning sessions were alternated with 73.5min sessions in the same chamber that was set up with the context that was not used for Pavlovian conditioning (referred to as the 'non-alcohol context'). The chamber light was illuminated after a 2-min delay, but neither the CS nor ethanol was presented. The pump was activated for 6 s at similar intervals as during Pavlovian conditioning sessions, but no syringes were mounted on the pumps.

At 24 h after the last training session (see below), port entries elicited by the CS were tested in the alcohol context or the non-alcohol context. At test, the CS was presented as during Pavlovian conditioning but without ethanol.

Experiment 1: Effect of context on alcohol-seeking elicited by the CS. Rats (N=21) received 12 sessions of home-cage ethanol exposure, after which 5 rats with the lowest ethanol intake values (1.70-3.8 g/kg) were dropped from the study. The remainder (n=16) received 10 Pavlovian conditioning sessions alternated with 10 sessions of exposure to the nonalcohol context. At 24 h after the last training session, alcohol-seeking elicited by the CS without ethanol was tested in the alcohol context for half the rats and in the non-alcohol context for the remainder. The test was repeated 24 h later in the alternate context for each rat, based on a within-subjects design.

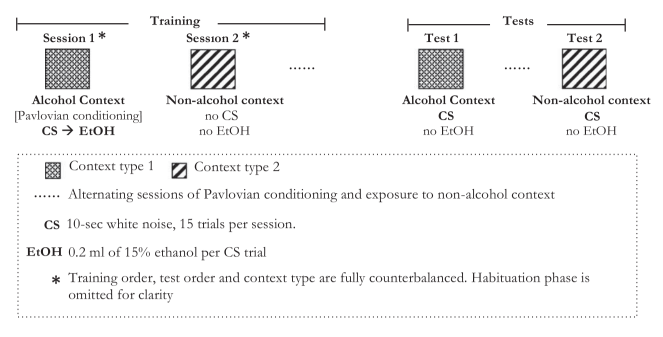
Experiment 2a: Effect of NBQX in the BLA on alcoholseeking elicited by the CS. Rats (N=42) received 15 sessions of home-cage ethanol exposure, after which 6 rats were dropped from the study because of low ethanol intake (n=2; <0.26 g/kg) or aggressive behavior (n=4). The rest (n=36) were implanted with guide cannulae targeting the BLA. Following recovery, 11 Pavlovian conditioning sessions occurred, which were alternated with 11 sessions of exposure to the non-alcohol context. At 24 h after the last training session, alcohol-seeking elicited by the CS without ethanol was tested in the alcohol context or the non-alcohol context. Next, three additional Pavlovian conditioning sessions and three sessions of exposure to the non-alcohol context were conducted, followed by a second, identical test in the alternate context.

Before each test, a bilateral microinfusion of NBQX (0, 0.3, or $1.0 \mu g/0.3 \mu l$) was administered into the BLA, with dose as a between-subjects factor that remained constant across tests in each context. Assignment to each dose was counterbalanced according to the average number of port entries elicited by the CS across the last three Pavlovian conditioning sessions that occurred before test 1. A sham microinfusion using injectors cut to the same length as the guide cannula was conducted before session 7, and a saline sham microinfusion was conducted before session 9 to acclimate rats to these procedures.

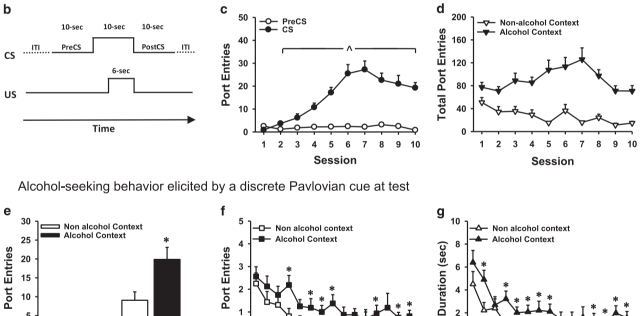
Experiment 2b: Effect of NBQX in the BLA on port entries elicited by the CS when it was paired with ethanol. Following Experiment 2a, five consecutive Pavlovian conditioning sessions were conducted. Before session 5, a bilateral infusion of NBQX (0, 0.3, or $1.0 \,\mu\text{g}/0.3 \,\mu\text{l}$) was administered into the BLA. Dose was between-subjects, and

а Schematic of behavioral assay

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Acquisition of Pavlovian alcohol-seeking behavior



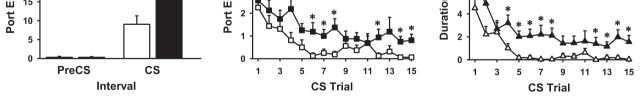


Figure I A new behavioral assay revealed that presentations of a discrete CS without ethanol elicited stronger alcohol-seeking in an alcohol context, compared with a context where alcohol was never consumed. Experiment 1, n = 16, within-subjects design. (a) Schematic illustration of the behavioral assay. (b) Depiction of the temporal relation between the CS and US (ethanol) and of intervals of intervals of interest. (c) Mean (± SEM) number of port entries during the CS (filled circles) and 10 s PreCS intervals (open circles) across 10 Pavlovian conditioning sessions in which each CS trial was paired with 0.2 ml of 15% ethanol. (d) Mean (± SEM) total number of port entries (sum of PreCS, CS, PostCS, and ITI) made during each Pavlovian conditioning sessions in an alcohol context (filled triangles) and during alternating sessions in a different, non-alcohol context where neither the CS nor ethanol were presented (open triangles). (e) Mean (± SEM) number of port entries during PreCS and CS intervals in the non-alcohol context (open bars) and alcohol context (filled bars) during tests in which the CS was presented without ethanol. (f) Mean (± SEM) number of port entries during each CS trial at test in both contexts. (g) Mean (± SEM) amount of time spent (seconds) in the port during each CS trial at test in both contexts. $^p < 0.05$, CS > PreCS. *p < 0.05, alcohol context > non-alcohol context.

assignment into each dose was based on behavior averaged across the prior four Pavlovian conditioning sessions. Each dose included some rats from each of the prior doses in Experiment 2a.

Following Experiment 2, data were dropped from one rat for failure to learn that the CS predicted ethanol and from nine rats with incorrect cannula placements (Supplementary Figure S1). In addition, two rats from Experiment 2a became too aggressive to be included in Experiment 2b. Final samples sizes after histological analyses were n = 26 for Experiment 2a and n = 25 for Experiment 2b.

Experiment 3: Effect of NBQX in the CPu on alcoholseeking elicited by the CS. Rats (N=12) received 15 sessions of home-cage ethanol exposure, followed by 11 Pavlovian conditioning sessions alternated with 11 sessions of exposure to the non-alcohol context. At test, alcoholseeking elicited by the CS without ethanol was tested in the non-alcohol context. A second, identical test was administered after four additional sessions each of Pavlovian conditioning and exposure to the alternate context.

Before each test, a bilateral microinfusion of NBQX (0 or $0.3 \mu g/0.3 \mu l$) was administered into the CPu, according to a within-subjects design. Data from one rat were dropped following the study because of failure to learn the CS-ethanol association (final n = 11).

A time line of experimental events and treatment groups for each experiment is provided in Table 1.

Histology

Rats were anesthetized with isoflurane, given a microinfusion of fast green dye through the guide cannulae, and decapitated. Brains were removed from the skull, postfixed in formalin for 24 h and 25% sucrose for 7 days, and then sectioned (60 µm, coronal) on a cryostat (-19 °C). Sections were mounted onto glass slides, stained with cresyl violet, and analyzed using light microscopy. Data from rats were included if the most ventral point of injector tips touched or was within the boundary of the BLA or CPu (Paxinos and Watson, 1997).

Data Analysis

During home-cage ethanol exposure, ethanol intake (g/kg; g of ethanol consumed/body weight in kg) and ethanol preference (%; (g of ethanol consumed/sum of water and ethanol intake in g)*100) were assessed in each session.

During Pavlovian conditioning sessions, port entries were recorded during: each 10 s CS trial (CS); 10 s intervals before each CS trial (PreCS); 10 s intervals after each CS trial (PostCS); and the variable ITI (total port entries minus the sum of PreCS, CS, and PostCS). The total number of port entries, which represents the sum of port entries made during the PreCS, CS, PostCS, and ITI, was also recorded.

Data were analyzed using mixed two- or three-way ANOVA. Homogeneity of variance was assessed using Mauchly's test of sphericity and adjusted using the Huynh-Feldt correction. Statistically significant main effects and interactions were pursued using t-tests for paired or independent sample. The significance level for ANOVA was $\alpha = 0.05$. ANOVA results that were not statistically significant are indicated as 'NS' (all F-values < 2.18 and all

Table I Timel	Table I Timeline of Experimental Events and Treatment Groups	nts and Tre	atment Groups			
	Home-cage ethanol exposure	Surgery	Surgery Behavioral training (alternating sessions in each context)	Test	Re-training	Test
Exp. 1 (<i>n</i> = 16)	12 sessions		CS → EtOH Pavlovian conditioning alcohol context (10 sessions) No CS, no EtOH Non-alcohol context (10 sessions)	CS, no EtOH Alcohol context Non-alcohol context		CS, no EtOH Non-alcohol context Alcohol context
Exp. 2a (<i>n</i> =26)	I.5 sessions	BLA	CS → EtOH Pavlovian conditioning, alcohol context (11 sessions) No CS, no EtOH Non-alcohol context (11 sessions)	CS, no EtOH Alcohol context ^a Non-alcohol context ^a	CS → EtOH Pavlovian conditioning, alcohol context (3 æssions) No CS, no EtOH Non-alcohol context (3 æssions)	CS, no EtOH Non-alcohol context ^a Alcohol context ^a
Exp. 2b (<i>n</i> =25)			Same rats as in Exp. 2a		CS → EtOH Pavlovian conditioning, alcohol context (4 sesions)	$CS \rightarrow EtOH$ Alcohol context ^a (1 session)
Exp. 3 (n=11)	I.5 sessions	CPu	CS → EtOH Pavlovian conditioning, alcohol context (11 sessions) No CS, no EtOH Non-alcohol context (11 sessions)	CS, no EtOH Non-alcohol context ^b (I session)	CS → EtOH Pavlovian conditioning, alcohol context (4 sessions) No CS, no EtOH Non-alcohol context (4 sessions)	CS, no EtOH Non-alcohol context ^b (I session)
^a Intra-BLA microin ^b Intra-CPu microin	$^{\rm h}$ htra-BLA microinfusion of NBQX before test (0, 0.3, or 1.0 $\mu g/0.3 ~\mu l)$ $^{\rm b}$ htra-CPu microinfusion of NBQX before test (0 or 0.3 $\mu g/0.3 ~\mu l)$: (0, 0.3, or 1. t (0 or 0.3 μg	0 µg/0.3 µl). 0.3 µl).			

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p-values > 0.145 for non-significant comparisons). For *post-hoc t*-tests, a Bonferroni-corrected α was calculated by dividing $\alpha = 0.05$ by the number of comparisons in each analysis. Only statistically significant *t*-test comparisons are reported. Analyses were conducted using SPSS (Version 18).

RESULTS

In all the three experiments, ethanol intake and preference increased across sessions of home-cage ethanol exposure (Supplementary Figure S2).

Experiment 1: Effect of Context on Alcohol-Seeking Elicited by the CS

We investigated the capacity of context to modulate alcoholseeking elicited by a discrete CS that predicted alcohol using a newly developed assay that equated exposure to an alcohol context and a non-alcohol context before test.

Rats learned to associate a discrete auditory CS with ethanol, as indicated by a comparison between port entries made during the CS and a PreCS intervals (Figure 1c). Port entries during the CS, but not the PreCS, increased across sessions (Session, F(9,135) = 17.65, p < 0.001; Interval, F(1,15) = 59.28, p < 0.001; Session × Interval, F(9,135) = 25.09, p < 0.001).

The total number of port entries made during each session (Figure 1d) was greater during Pavlovian conditioning compared with sessions in the non-alcohol context (Context, F(1,15) = 77.75, p < 0.001). Total port entries decreased across session (Session, $F(9,135) = 3.46 \ p = 0.001$), with a more pronounced reduction in the non-alcohol context (Context × Session $F(9,135) = 3.91 \ p < 0.001$). Paired samples *t*-tests (Bonferroni-corrected α , p = 0.025) verified that in the non-alcohol context total port entries in session 1 were higher than in session 10 (t(15) = 3.67, p = 0.002) but the same in sessions 1 and 10 in the alcohol context. The across-session reduction in total port entries in the non-alcohol context suggests that this context became associated with the absence of ethanol. Similar results across acquisition were obtained for Experiments 2 and 3 (Supplementary Figure S3).

At test, alcohol-seeking elicited by the CS without ethanol was examined in the alcohol context and the non-alcohol context. The CS alone elicited port entries in the nonalcohol context, and this behavior was invigorated in the alcohol context (Figure 1e-g). ANOVA on CS and PreCS port entries at test (Figure 1e) indicated higher overall responding in the alcohol context (Context, F(1,15) = 39.67, p < 0.001) and during the CS interval (Interval, F(1,15) = 10.86, p = 0.005). Importantly, port entries during the CS but not PreCS were elevated in the alcohol context, compared with the non-alcohol context (Context \times Interval, F(1,15) = 12.39, p = 0.003). Paired-samples *t*-test (Bonferroni-corrected) α , p = 0.013) verified that more port entries occurred during the CS than the PreCS in the non-alcohol context (t(15) =3.93, p = 0.001) and the alcohol context (t(15) = -6.22), p < 0.001). There was no difference in PreCS responding across contexts but more CS port entries in the alcohol context than the non-alcohol context (t(15) = 3.41), p = 0.004). There was no effect of context on port entries during PostCS or ITI intervals (Supplementary Figure S4).

To determine whether context had an impact on withinsession responding to the CS, the number of port entries elicited by each CS trial at test was examined (Figure 1f). Overall, more port entries occurred in the alcohol context than the non alcohol context (Context, F(1,15) = 10.52, p = 0.005), and there was a reduction in the number of port entries per CS trial (Trial, F(14,210) = 7.68, p < 0.001) that occurred in both contexts (Trial × Context = NS). Planned comparisons to test the hypothesis that CS presentations in the alcohol context would elicit greater responding on a trial-by-trial basis indicated that port entries were higher in the alcohol context compared with the non-alcohol context on trials 4, 6, 7, 8, 12, 14, and 15 (all ps < 0.05).

The amount of time spent in the port during each CS trial (Figure 1g) was higher in the alcohol context (Context, F(1,15) = 17.19, p = 0.001) and decreased across trials (Trial, F(14,210) = 12.69, p < 0.001) in both contexts (Trial × Context = NS). Planned comparisons indicated that more time was spent in the port in the alcohol context relative to the non-alcohol context on trials 2, 4, 5, 6, 7, 8, 12, 13, 14, and 15 (all ps < 0.05).

Experiment 2a: Effect of NBQX in the BLA on Alcohol-Seeking Elicited by the CS

As in Experiment 1, presentations of the CS without ethanol elicited more alcohol-seeking in the alcohol context than the non-alcohol context. This statement is based on data from saline-treated rats (Figure 2a), who made more port entries during the CS in the alcohol context at test, with no difference in PreCS responses across contexts (Context, F(1,11) = 5.12, p = 0.045; Interval, F(1,11) = 61.97, p < 0.001; Context × Interval, F(1,11) = 6.96, p = 0.023).

Glutamate in the BLA is implicated in cue-induced reinstatement of operant alcohol-seeking behavior (Gass et al, 2011). Here we found that blocking AMPA glutamate receptors in the BLA reduced Pavlovian alcohol-seeking, and this effect was not dependent on the test context. ANOVA comparing port entries during CS and preCS intervals as a function of NBQX dose (Figure 2a) indicated that port entries were higher in the alcohol context than the nonalcohol context (Context, F(1,23) = 8.65, p = 0.007) and that more port entries were made during the CS than the PreCS (Interval, F(1,23) = 67.04, p < 0.001). Port entries during the CS, but not the PreCS interval, were elevated in the alcohol context, compared with the non-alcohol context (Context× Interval, F(1,23) = 10.69, p = 0.003). NBQX reduced port entries during the CS (Dose, F(2,23) = 14.61, p < 0.001; Interval × Dose, F(2,23) = 14.84, p < 0.001) in both contexts (Context \times Dose = NS; Context \times Interval \times Dose = NS). Separate analyses conducted on PreCS and CS responding confirmed that port entries during the CS were elevated in the alcohol context (Context, F(1,23) = 9.66, p = 0.005) and significantly reduced by NBQX (Dose, F(2,23) = 14.77, p < 0.001) in both contexts (Context × Dose = NS). To further investigate the significant main effect of Dose, Bonferroni post-hoc tests were conducted to compare CS port entries at each dose on data collapsed across context. This analysis indicated that, compared with saline, both doses of NBQX reduced CS responses $(0.3 \,\mu\text{g}/0.3 \,\mu\text{l}, p = 0.005: 1 \,\mu\text{g}/0.3 \,\mu\text{l},$ p < 0.001). There was no difference between the two NBQX

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Bilateral microinfusion of NBQX into the BLA before test in the non-alcohol context and the alcohol context

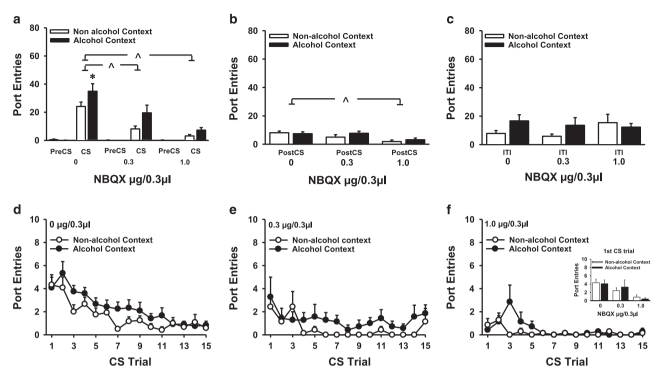


Figure 2 NBQX infused bilaterally into the basolateral amygdala reduced alcohol-seeking elicited by a discrete CS in both the non-alcohol context (open bars and symbols) and the alcohol context (filled bars and symbols). Experiment 2a ($0 \mu g/0.3 \mu$ l, n = 12; $0.3 \mu g/0.3 \mu$ l, n = 7; $1.0 \mu g/0.3 \mu$ l, n = 7). (a) Mean (\pm SEM) number of port entries during PreCS and CS intervals. (b) Mean (\pm SEM) number of port entries during PostCS intervals. (c) Mean (\pm SEM) number of port entries during the variable ITI, calculated by subtracting port entries during PreCS, CS, and PostCS intervals from total port entries. *p < 0.05, alcohol context. $^p < 0.05$, saline > NBQX on data collapsed across context. (d–f) Mean (\pm SEM) number of port entries per CS trial at each dose of NBQX. Inset depicts the mean (\pm SEM) number of port entries at the first CS trial for each dose in both contexts.

doses. NBQX had no effect on PreCS port entries (main effects and interaction = NS).

There was no impact of context (Context = NS) on port entries made during the 10 s PostCS interval (Figure 2b). However, NBQX reduced PostCS responding (Dose, F(2,23) = 7.46, p = 0.003) in both contexts (Context × Dose = NS). Bonferroni *post-hoc* analyses on data collapsed across context indicated a significant reduction in PostCS responses after 1.0 µg/0.3 µl NBQX compared with saline (p = 0.003), with no differences between saline and 0.3 µg/0.3 µl or 0.3 µg/0.3 µl and 1.0 µg/0.3 µl doses.

Port entries made during the variable intertrial interval between CS presentations were unaffected by NBQX (Figure 2c; main effects and interaction = NS).

The number of port entries made during each CS trial at test in both contexts was examined to evaluate the effect of NBQX on within-session responding (Figure 2d–f). A Dose × Trial × Context ANOVA indicated that this measure decreased across trials (Trial, F(14,322) = 10.32, p < 0.001), with significantly less responding after NBQX relative to saline (Dose, F(2,23) = 14.79, p < 0.001; Trial × Dose, F(28,322) = 2.72, p < 0.001). Responding in the alcohol context was elevated compared with the non-alcohol context (Context, F(1,23) = 9.81, p = 0.005), and this difference did not vary as a function of trial or dose (Context × Dose = NS; Trial × Context = NS; Trial × Context × Dose = NS). To further investigate the significant Trial × Dose interaction, Bonferroni comparisons were conducted to compare port entries made during the first CS trial across doses on data collapsed across context. This analysis indicated that, compared with saline, the first CS trial elicited fewer port entries following the $1.0 \ \mu g/0.3 \ \mu l$ dose (p = 0.005) but not the $0.3 \ \mu g/0.3 \ \mu l$ dose. There was no difference between the two doses of NBQX. The inset in Figure 2f depicts port entries at the first CS trial for each dose in both contexts. Similar outcomes were revealed for the amount of time spent in the port during each CS trial at test (Supplementary Figure S5).

Experiment 2b: Effect of NBQX in the BLA on Port Entries Elicited by the CS When it was Paired With Ethanol

To determine whether blocking AMPA glutamate receptors in the BLA affected the capacity to make a port entry response, NBQX was administered into the BLA before a Pavlovian conditioning session in which the CS was paired with ethanol. This manipulation had no impact on port entries (Figure 3). More port entries occurred during the CS than the PreCS interval at each NBQX dose tested (Figure 3a; Interval, F(1,22) = 118.52, p < 0.001; Dose = NS; Interval × Dose = NS). Furthermore, the number of port entries elicited by each CS trial at test (Figure 3b) was unaffected by NBQX (main effects and interaction = NS).

Bilateral microinfusion of NBQX into the BLA before a Pavlovian conditioning session

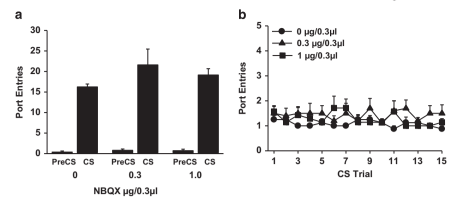


Figure 3 NBQX infused bilaterally into the basolateral amygdala had no impact on Pavlovian alcohol-seeking when the CS was paired with ethanol. Experiment 2b ($0 \mu g/0.3 \mu$ |, n = 8; $0.3 \mu g/0.3 \mu$ |, n = 10; $1.0 \mu g/0.3 \mu$ |, n = 7). (a) Mean (\pm SEM) number of port entries during PreCS and CS intervals. (b) Mean (\pm SEM) number of port entries per CS trial.

Experiment 3: Effect of NBQX in the CPu on Alcohol-Seeking Elicited by the CS

Results from Experiment 2a suggested that AMPA glutamate receptors in the BLA were needed for Pavlovian alcoholseeking, regardless of the context in which the discrete CS was experienced. However, the attenuation in Pavlovian alcohol-seeking produced by intra-BLA NBQX might have been caused by upward diffusion of NBQX into the CPu. To test this hypothesis, we examined the effect of NBQX in the CPu on alcohol-seeking elicited by the CS without ethanol in a non-alcohol context.

Blocking AMPA glutamate receptors in the CPu had no impact on alcohol-seeking elicited by the CS alone in a nonalcohol context (Figure 4). Port entries during the CS were higher than during the PreCS interval after both saline and NBQX (Figure 4a; Interval, F(1,10) = 78.54, p < 0.001; Dose = NS; Interval × Dose = NS). The number of port entries elicited by each CS trial (Figure 4b) decreased across session (Trial, F(14,70) = 6.33, p < 0.001), with no impact of NBQX on this measure (Dose = NS; Interval × Dose = NS). NBQX had no impact on port entries made during the PostCS interval (Figure 4c) or ITI (Figure 4c).

The location of injector tips within the BLA in Experiment 2 and CPu in Experiment 3 is shown in Figure 5.

DISCUSSION

We report that alcohol-seeking behavior elicited by a discrete, Pavlovian CS was invigorated in a context associated with prior alcohol consumption, compared with a context where alcohol had never been consumed. Blocking AMPA glutamate receptors bilaterally in the BLA attenuated this behavior in both contexts but had no impact on alcohol-seeking elicited by the CS when it was paired with alcohol. Moreover, blocking AMPA receptors in the CPu did not affect alcohol-seeking elicited by the CS when it was presented without alcohol in a non-alcohol context. These results indicate that the context in which a discrete alcohol CS is experienced can be an important determinant in the level of conditioned alcohol-seeking elicited by that CS. They also suggest that glutamate signaling via AMPA receptors in

the BLA is critical for the expression of alcohol-seeking elicited by a discrete alcohol CS, regardless of the context in which the CS is experienced.

In the behavioral procedure described herein, rats were trained to discriminate between one context where a CS was paired with alcohol, and a second context where neither the CS nor alcohol was presented. They received equal exposure to both contexts before test, thereby addressing a limitation of our prior research in which the invigoration of alcoholseeking in an alcohol context may have been attributable to a longer period of pretest exposure to the alcohol context, relative to the non-alcohol context (Remedios *et al*, 2014). The across session decrease in total port entries in the nonalcohol context suggests that rats learned to associate that context with the absence of alcohol.

At test, presentations of the CS without alcohol triggered alcohol-seeking in both the non-alcohol context and the alcohol context; however, this conditioned response was significantly invigorated in the alcohol context. These results corroborate a growing literature indicating that the context in which a discrete drug cue is experienced can have a profound impact on the level of drug-seeking behavior elicited by that cue (Crombag and Shaham, 2002; Tsiang and Janak, 2006; Zironi et al, 2006; Chaudhri et al, 2008; Remedios et al, 2014; Fuchs et al, 2008; Hamlin et al, 2008; Hamlin et al, 2007; Janak and Chaudhri, 2010). It is notable that the discrete CS elicited alcohol-seeking when presented for the first time without alcohol in the non-alcohol context. This experimental condition parallels clinical and human laboratory studies in which conditioned subjective and physiological responses to alcohol-predictive cues are assessed in laboratory environments that bear little resemblance to 'real world' drinking contexts. Interestingly, when alcohol consumption is assessed in a typical laboratory setting vs a bar, consumption in the latter context is enhanced (Billings et al, 1976; Strickler et al, 1979; Wigmore and Hinson, 1991). Researchers have recently begun to incorporate context into experimental designs in order to assess reactivity evoked by alcohol-predictive cues under more ecological valid experimental conditions (Nees et al, 2012; Paris et al, 2011). Results indicate that contexts associated with alcohol consumption

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Bilateral microinfusion of NBQX into the CPu before test in the non-alcohol context

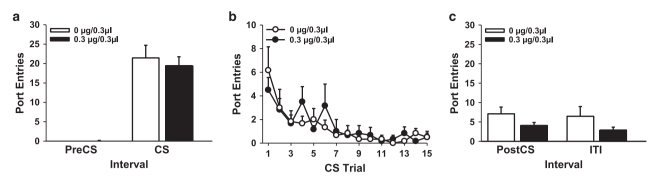


Figure 4 NBQX infused bilaterally into the caudate putamen had no impact on Pavlovian alcohol-seeking when the CS was presented without ethanol in a non-alcohol context. Experiment 3 (n = 11, within-subjects). (a) Mean (\pm SEM) number of port entries during PreCS and CS intervals. (b) Mean (\pm SEM) number of port entries during PostCS and ITI intervals.

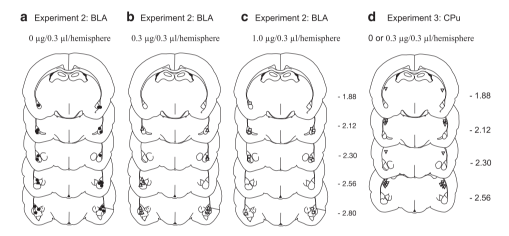


Figure 5 Location of injector tips in the basolateral amygdala (a–c, Experiment 2, dose is between-subjects) and the caudate putamen (d, Experiment 3, dose is within-subjects).

can influence cue reactivity in abstinent alcohol-dependent patients (Nees *et al*, 2012).

Localized BLA infusions of a selective AMPA glutamate receptor antagonist reduced alcohol-seeking elicited by the CS in both contexts. Although NBQX also reduced port entries during a PostCS interval, it had no impact on port entries made during the PreCS interval or variable ITI. It is thus unlikely that NBQX caused a generalized decrease in locomotion or impaired the capacity to make a port entry response. The finding that NBQX had no effect on CS port entries when the CS was paired with alcohol also supports this conclusion. A comparison across Experiments 2a and 2b suggests that AMPA receptors in the BLA are particularly important for alcohol-seeking that is guided by the memory of a CS-alcohol association. Interestingly, NBQX in the BLA reduced port entries elicited by the first CS trial at test in Experiment 2a, which suggests that AMPA receptors in this region may be needed for initiating alcohol-seeking in response to a discrete Pavlovian alcohol CS, as opposed to continued motivation to respond to the CS.

A neuroanatomical control study was conducted to rule out the possibility that the observed effects of NBQX were not attributable to actions in the caudal CPu, which would be accessible by upward diffusion through the cannula track. This region receives glutamatergic inputs from multiple sources, including frontal cerebral cortex (Fonnum *et al*, 1981), and expresses AMPA receptors (Albin *et al*, 1992). Blocking AMPA receptors in the CPu had no effect on CS port entries when the CS was presented without alcohol in a non-alcohol context. Moreover, data from rats in which cannulae failed to target the BLA (Supplementary Figure S1) indicate that NBQX outside the BLA had no systematic influence on alcohol-seeking.

The BLA is reciprocally connected with the prelimbic (PL) medial prefrontal cortex (McDonald, 1991, 1998; Sesack *et al*, 1989; Stefanik and Kalivas, 2013; Senn *et al*, 2014), and the PL is important for conditioned responding elicited by discrete, Pavlovian cues (Sierra-Mercado *et al*, 2011). Electrophysiological data obtained using a conditioned place preference (CPP) paradigm reveal that PL neurons are active during the acquisition, extinction, and recall of Pavlovian associations between morphine and a context where morphine is administered (Sun *et al*, 2011), and reductions in brain-derived neurotropic factor signaling in the PL cause

deficits in cocaine CPP (Choi et al, 2012). The PL also appears to be involved in conditioned responses elicited by Pavlovian cues that predict alcohol. Neuroimaging studies reveal increased brain activity in the prefrontal cortex of alcoholic subjects exposed to proximal alcohol cues (George et al, 2001), and self-reported craving correlates positively with neural activation in response to alcohol cues in the anterior cingulate and medial prefrontal cortex of patients across several phases of alcohol dependence (Fryer et al, 2013). Data obtained using an aversive Pavlovian conditioning assay suggest that projections from the PL to the BLA are involved in the expression of conditioned responses elicited by a discrete CS that predicts shock (Orsini et al, 2011). PL-to-BLA projections are also important for appetitive learning tasks such as the reinstatement of operant cocaineseeking behavior (Mashhoon et al, 2010), context-induced renewal of operant cocaine-seeking behavior (Fuchs et al, 2007), and risk-based decision-making in rats (St Onge et al, 2012). Future research should examine the role of PL-to-BLA projections in alcohol-seeking elicited by a discrete Pavlovian alcohol CS.

Another interesting way to advance the present research would be to characterize rats into sign-trackers or goaltrackers before behavioral training and testing (Robinson and Flagel, 2009). Rats that goal-track in response to a foodpredictive CS subsequently show greater levels of cocainecontext-induced hyperactivity and context-induced renewal of cocaine-seeking behavior than rats characterized as signtrackers (Saunders *et al*, 2014). Thus, the goal-tracking phenotype may be associated with a propensity to utilize contextual cues to motivate behavior. Based on this hypothesis, rats identified as goal-trackers may show greater context-mediated invigoration on Pavlovian alcohol-seeking behavior in the present assay, compared with the signtracking phenotype.

The present results have several important implications. They suggest that cue-induced alcohol craving measured in laboratory settings in humans likely underestimates craving levels that would be observed if craving was assessed in contexts associated with alcohol consumption (Litt and Cooney, 1999). They also imply that the combined experience of discrete and contextual alcohol cues may be a more potent trigger for relapse than either type of cue alone (Chaudhri et al, 2008; Remedios et al, 2014; Tsiang and Janak, 2006). Finally, they complement findings from instrumental conditioning procedures that implicate BLA glutamate in cue-induced alcohol-seeking (Gass et al, 2011; Backstrom et al, 2004a; Sinclair et al, 2012; Backstrom and Hyytia, 2004b) and extend this literature by identifying a novel role for AMPA glutamate receptors in the BLA in alcohol-seeking behavior elicited by discrete Pavlovian alcohol cues.

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AUTHOR CONTRIBUTIONS

NC and PHJ designed the experiments and obtained funding. RR and JMS collected and analyzed the data. NC and JMS wrote the manuscript and all authors provided feedback.

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Supplementary Information accompanies the paper on the Neuropsychopharmacology website (http://www.nature.com/npp)