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D-Serine and D-Cycloserine Reduce Compulsive Alcohol Intake in Rats

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There is considerable interest in NMDAR modulators to enhance memory and treat neuropsychiatric disorders such as addiction, depression, and schizophrenia. D-serine and D-cycloserine, the NMDAR activators at the glycine site, are of particular interest because they have been used in humans without serious adverse effects. Interestingly, D-serine also inhibits some NMDARs active at hyperpolarized potentials (HA-NMDARs), and we previously found that HA-NMDARs within the nucleus accumbens core (NAcore) are critical for promoting compulsion-like alcohol drinking, where rats consume alcohol despite pairing with an aversive stimulus such as quinine, a paradigm considered to model compulsive aspects of human alcohol use disorders (AUDs). Here, we examined the impact of D-serine and D-cycloserine on this aversion-resistant alcohol intake (that persists despite adulteration with quinine) and consumption of quinine-free alcohol. Systemic D-serine reduced aversion-resistant alcohol intake (that persists despite adulteration support of quinine-free alcohol or saccharin with or without quinine. Importantly, D-serine within the NAcore but not the dorsolateral striatum also selectively reduced aversion-resistant alcohol drinking, without altering consumption of mPFC–NAcore terminals in alcohol-drinking rats, similar to reported effects of the NMDAR blocker AP5. Further, D-serine preexposure occluded AP5 inhibition of mPFC-evoked EPSCs, suggesting that D-serine reduced EPSCs by inhibiting HA-NMDARs. Systemic D-cycloserine also selectively reduced intake of quinine-adulterated alcohol, and D-cycloserine inhibited NAcore HA-NMDARs *in vitro*. Our results indicate that HA-NMDAR modulators can reduce aversion-resistant alcohol drinking, and support testing of D-serine and D-cycloserine as immediately accessible, FDA-approved drugs to treat AUDs.

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INTRODUCTION

The compulsion to drink alcohol despite significant adverse social, physical, and legal consequences is a pernicious obstacle to treating alcohol use disorders (AUDs) (Anton, 2000; Koob and Volkow, 2010; Modell *et al*, 1992). Because existing pharmacotherapies are moderately efficacious or act only in a subset of AUD patients (Spanagel, 2009), there is a critical need to better understand the mechanisms driving compulsive intake in order to improve treatment. Procedures in which animals voluntary self-administer alcohol despite pairing it with aversive stimuli provide models to study and potentially intervene in this aversion-resistant, compulsion-like drinking. For example, we found that, following ~ 3 months of intermittent alcohol intake, rats continue to drink alcohol despite pairing intake with footshocks or the

bitter-tastant quinine (Hopf *et al*, 2010; Seif *et al*, 2013). This aversion-resistant intake is considered to model compulsive aspects of human AUDs (Everitt and Robbins, 2005; Hopf and Lesscher, 2014; Hopf *et al*, 2010; Lesscher *et al*, 2010; Seif *et al*, 2013; Vengeliene *et al*, 2009).

The nucleus accumbens core (NAcore) integrates motivationally relevant stimuli to drive activity (Everitt and Robbins, 2005; Koob and Volkow, 2010) including aversion-related behavior (Roitman et al, 2005; Schwienbacher et al, 2004). We recently found that long-term, intermittent alcohol consumption in rats leads to appearance of unusual NMDA receptors that are active at hyperpolarized potentials (HA-NMDARs) within the NAcore (Seif et al, 2013). These HA-NMDARs are only observed under cortical inputs to NAcore, specifically from the medial prefrontal cortex (mPFC) and insula (Seif et al, 2013), regions that regulate addiction- and aversion-related behaviors (Everitt and Robbins, 2005; Koob and Volkow, 2010; Naqvi et al, 2014). Importantly, this HA-NMDAR/cortical-NAcore signaling is critical for promoting aversion-resistant alcohol drinking, but does not regulate alcohol drinking that is not explicitly paired with aversive consequences (Seif et al, 2013). In this regard, some clinical investigators (Naqvi et al, 2014; Tiffany

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and Conklin, 2000) have suggested that cortical regions are selectively recruited by the conflict associated with aversionresistant, compulsive intake, that is, between motivation to drink alcohol and the desire to avoid aversive consequences. In contrast, alcohol intake without overt adverse consequences would involve the striatum, with less requirement for cortical input. Thus, we propose that NAcore HA-NMDARs play a role in overcoming conflict associated with aversion-resistant intake, and thus represent a novel therapeutic target to counteract the compulsion to drink alcohol.

There has long been interest in using NMDAR modulators to address a number of human conditions, for example, to suppress the enhanced glutamatergic function associated with protracted alcohol consumption (Gass and Olive, 2008; Krystal et al, 2003, 2011; Spanagel, 2009), and to enhance memory as well as overcome decreased NMDAR function that may contribute to schizophrenia, Parkinson's disease, and cocaine addiction (D'Ascenzo et al, 2014; Heresco-Levy et al, 2013; Martineau et al, 2014; Sani et al, 2012; Tsai and Lin, 2010). Modulators of the NMDAR glycine site, such as D-serine and D-cycloserine (DCS), are of particular interest because of their use in many patient populations with few side effects or safety issues noted (D'Ascenzo et al, 2014; D'Souza et al, 2013; Heresco-Levy et al, 2013; Kantrowitz et al, 2010; Tsai and Lin, 2010; but see Krug et al, 2007), and because the moderate NMDAR inhibition these reagents afford is considered more favorable than stronger NMDAR inhibitors that can have significant adverse effects (cf, Heresco-Levy et al, 2013; Millan, 2005).

Although D-serine is a canonical positive modulator of NMDARs (Cull-Candy and Leszkiewicz, 2004; Millan, 2005), it can also inhibit HA-NMDARs (Chatterton et al, 2002; Takarada et al, 2009, 2012). Here, we examined whether D-serine or DCS systemically, or D-serine within the NAcore, could suppress aversion-resistant alcohol intake. We also combined in vitro electrophysiology with optogenetics to address the molecular mechanism of D-serine and DCS. Together, our results support D-serine and DCS as immediately accessible pharmacological therapies to reduce compulsive drinking associated with AUDs.

MATERIALS AND METHODS

Animal Handling

All procedures followed the Guide for Care and Use of Laboratory Animals provided by the National Institutes of Health, and approval of the institutional animal care and use committee of UCSF. Adult male Wistar rats (~450-550 g at surgery) were individually housed with ad libitum access to food and water.

Alcohol Drinking

Rats learned to voluntarily consume alcohol through twobottle, home cage, intermittent access to 20% alcohol (IAA) as previously described (Hopf et al, 2010; Seif et al, 2013). Starting on Monday, Wednesday, and Friday late afternoons, rats had 16-20 h access to two bottles, one containing alcohol (20% v/v with water) and the other water. After ~ 10 weeks of IAA, rats were switched to drink alcohol versus water under

two-bottle choice for 20 min/day, 5 day/week in the late afternoon. Rat blood alcohol concentrations were 48.5 ± 9.4 mg/dl for 0.97 ± 0.09 g/kg alcohol intake (n = 10), measured 30 min after starting access to alcohol (methods as in Carnicella et al, 2009). After ~4 weeks of 20 min/day access, and after intake had <20% variability for 5 days, we tested alcohol intake with or without the bitter-tastant quinine (10 mg/l, 0.001%, ~ 25 μ M) (20 min/day, 5 day/week) or performed in vitro electrophysiology.

To minimize the possible contribution of habituation to the aversive properties of quinine across time, rats first had minimal initial habituation to alcohol+quinine (2-3 days). Importantly, rats then experienced the different experimental conditions (alcohol drinking with or without quinine, and with vehicle versus D-serine or DCS) in a randomized, counterbalanced manner using a Latin-Square design. Rats were therefore as likely to receive D-serine during the first alcohol-quinine session as they were to receive vehicle, and thus D-serine changes in alcohol drinking were unlikely to represent order effects or habituation to quinine across time. Also, rats always had at least 1 day of drinking alcohol without quinine and without drug treatment between experimental test days, avoiding any concerns related to drinking quininealcohol across successive days. D-serine, DCS, or saline was injected 15-20 min before access to alcohol.

Saccharin Intake

After ~10 weeks of IAA followed by ~4 weeks of access to alcohol for 20 min/day, 5 day/week, rats were switched to drinking saccharin ± quinine versus water under two-bottle choice access for 20 min/day, 5 day/week in the late morning (1100–1200 h). These rats also drank alcohol (20 min/day) in the late afternoon. For the first experiment, rats drank one of four cocktails: 300 mg/l (0.03%, 1.65 mM) saccharin, 1330 mg/l saccharin (0.13%, 7.3 mM), 1330 mg/l saccharin +10 mg/l quinine, or 1330 mg/l saccharin+30 mg/l quinine (Seif et al, 2013). For the second experiment, rats drank 150 mg/l saccharin (0.015%, 0.83 mM) or 300 mg/l saccharin +10 mg/l quinine. Different conditions were randomized across and within rats, and across days, using a Latin-square design. Rats drank each concentration of saccharin \pm quinine at least 4 days before D-serine/DCS experiments.

Surgery and Histology

Surgical methods were as previously described (Seif et al, 2013). Rats were infected in the mPFC (for NAcore recording: AP +3.20, ML \pm 1.20, DV -3.00, 10° tilt; for DLS recording: same except DV -2.00) with 0.6 µl/side of AAV-Camk2a-ChR2-EYFP ~ 8 weeks before in vitro experiments. Cannulae bilaterally targeted the medial NAcore (AP +2.20, ML ±2.70, DV -6.18, 8° tilt) or DLS (AP +1.0, ML \pm 3.6, DV – 4.0), with the injector 1 mM below cannulae tip during D-serine injection (in $0.6 \,\mu$ l, over 60 s, then 60 s diffusion before removing the injector).

In vitro Electrophysiology

Methods, including slice preparation, were as previously described (Seif et al, 2013). Internal solution for excitatory postsynaptic currents (EPSCs) contained (in mM): 120

D-serine and D-cycloserine reduce compulsive alcohol intake T Seif et al



Figure 1 Systemic D-serine only reduced aversion-resistant alcohol consumption. (a) D-serine (i.p., 300 mg/kg) reduced drinking of quinine-adulterated alcohol but not quinine-free alcohol (quinine: $F_{1,7} = 20.675$, p = 0.003; D-serine: $F_{1,14} = 17.799$, p = 0.001; interaction: $F_{1,14} = 4.294$, p = 0.035). *p < 0.05 for quinine/D-serine-300 vs quinine/vehicle (n = 8). D-serine at 600 mg/kg reduced intake of both quinine-adulterated and quinine-free alcohol (*p < 0.05 quinine/free vs quinine conditions; *p < 0.05 D-serine vs vehicle). (b, c) D-serine decreases in quinine-adulterated alcohol drinking were not accompanied by changes in water intake (b) (quinine: $F_{1,7} = 1.926$, p = 0.208; D-serine: $F_{1,14} = 1.210$, p = 0.327; interaction: $F_{1,14} = 3.268$, p = 0.068) but there were significant decreases in preference (alcohol intake/(alcohol+water intake)) (c) (quinine: $F_{1,7} = 1.3862$, p = 0.007; D-serine: $F_{1,14} = 12.561$, p < 0.001; interaction: $F_{1,14} = 3.313$, p = 0.066). (d) D-serine treatment did not reduce alcohol intake the day after D-serine treatment (quinine: $F_{1,7} = 0.099$, p = 0.763; D-serine: $F_{1,14} = 1.589$, p = 0.239; interaction: $F_{1,14} = 1.194$, p = 0.332). (e) D-serine (300 mg/kg) did not alter intake of saccharin $\pm quinine$ (n = 13; saccharin-quinine doses: $F_{3,36} = 27.837$, p < 0.001; D-serine: $F_{1,36} = 0.213$, p = 0.653; interaction: $F_{3,36} = 2.074$, p = 0.121). Quinine significantly reduced intake of 1330 g/l saccharin $\binom{30}{2}$ duinine flow s without quin; $^{+}p < 0.01$, sacc with 10 vs 30 mg/l quin). (f) D-serine (300 mg/kg) did not reduce, but instead slightly increased, consumption of lower saccharin concentrations (150 mg/l) or saccharin-quinine (300 mg/l) sacc+10 mg/l quin) (n = 14; saccharin $\pm quinine: F_{1,13} = 1.3.886$, p = 0.003; interaction: $F_{1,13} = 0.035$, p = 0.854). (g, h) After only 6 or 8 weeks of intermittent alcohol intake (n = 17), quinine adulteration (30 mg/l) signific

cesium-methanesulfonate, 20 HEPES, 0.4 EGTA, 2.8 NaCl, 5 TEA-Cl, 2.5 Mg-ATP, and 0.25 Na-GTP, pH 7.2–7.3. ACSF contained (in mM): 126 NaCl, 2.5 KCl, 1.2 NaH₂PO4, 1.2 MgCl₂, 2.4 CaCl₂, 18 NaHCO₃, and 11 glucose, pH 7.2–7.4, 31–32 °C. EPSCs were recorded using Clampex 10.1 and

Axon 700A amplifier (Molecular Devices, Foster City, CA), filtered at 2 kHz and digitized at 10 kHz. Series and input resistance were monitored with a 4-mV (50 ms) depolarizing step 800 ms after every EPSC. Electrically evoked currents were elicited using a bipolar stimulating electrode $\sim 200 \,\mu\text{M}$

dorsal to the patched neuron. ChR2-evoked EPSCs were elicited with an LED (Prizmatrix, 480 nm, 5 ms). EPSC data were expressed as percent change from baseline, averaged during 5 min before drug application. For statistics, drug response was the average current across the last 3 min of drug application.

Reagents

AAV2-*CamK2*-ChR2-eYFP was from University of North Carolina, Chapel Hill Vector Core. Quinine hydrochloride, D-serine, and DCS were from Sigma-Aldrich (St Louis, MO). Saccharin sodium hydrate was from Acros Organics (Morris Plains, NJ).

Statistical Analysis

All data are shown as mean \pm SEM. D-serine and DCS behavioral results were normally distributed and evaluated by two-way repeated-measures ANOVA with repeated measures for \pm quinine and \pm D-serine treatment, and Tukey's *post hoc* test. *In vitro* experiments were analyzed using an unpaired *t*-test for normally distributed data, and Mann–Whitney test or Friedman repeated-measures ANOVA on ranks for results that were not normally distributed. Statistical analyses were performed with Sigma-Plot v13 (Systat Software, San Jose, CA).

RESULTS

Systemic Administration of D-Serine Reduced Aversion-Resistant Alcohol Intake

D-serine is an agonist at most NMDAR subunits, but can also inhibit HA-NMDARs (Chatterton et al, 2002; Takarada et al, 2009, 2012). We recently showed that inhibiting NAcore HA-NMDARs selectively decreases aversion-resistant alcohol drinking, with no effect on intake not explicitly paired with aversive consequences (Seif et al, 2013). Thus, we hypothesized that D-serine would selectively suppress consumption of alcohol adulterated with 10 mg/l quinine. D-serine at 300 mg/kg (i.p.) significantly reduced drinking of quinineadulterated alcohol, with no effect on quinine-free alcohol intake (Figure 1a, p < 0.05 post hoc). A higher D-serine dose (600 mg/kg) reduced alcohol consumption irrespective of whether or not quinine was present (Figure 1a), indicating more general effects at this dose (see also Lockridge et al, 2012). Decreased aversion-resistant drinking was not accompanied by significant changes in concurrent water intake (Figure 1b), or a significant quinine/D-serine interaction for alcohol preference (Figure 1c, see figure legend). In addition, alcohol drinking the day after testing was not different among groups, indicating no lasting effects of D-serine exposure on alcohol drinking (Figure 1d).

Alcohol can be perceived as sweet or sweet/bitter combined (Goodwin *et al*, 2000). As decreased consumption of quinine-adulterated alcohol might reflect changes in palatability, we next determined whether D-serine reduced intake of saccharin alone or combined with quinine. Quinine decreased intake of saccharin (1330 mg/l) in a dosedependent manner, but D-serine (300 mg/kg) did not alter consumption of saccharin-quinine (Figure 1e). In a separate experiment, D-serine did not alter drinking of lower concentrations of saccharin-quinine or saccharin alone (Figure 1f) where drinking volumes were more similar to those during alcohol intake (~5-6 ml/kg). Similar to alcohol sessions, concurrent water intake was low during saccharinquinine sessions, resulting in high preference values (not shown). Although 10 mg/l quinine more moderately reduced intake of 1330 mg/l saccharin, quinine decreased consumption of a lower saccharin dose (300 mg/l) by ~ 50% (p < 0.001), confirming the aversive nature of this quinine concentration. In agreement, quinine adulteration significantly reduced alcohol drinking after only 6 or 8 weeks of intermittent alcohol intake (Figure 1g and h), concurring with our previous results that showed that aversion-resistant alcohol drinking develops after 12 weeks but not 6 weeks of intermittent intake (Hopf et al, 2010). Furthermore, quinine in water is highly aversive to alcohol-drinking rats at 1 mg/l quinine or greater (Hopf et al, 2010). Together, these results confirm that quinine concentrations used here were aversive to intermittent-drinking rats, supporting the conclusion that systemic D-serine specifically reduced aversion-resistant alcohol intake without altering taste palatability or general motor function.

D-Serine within the NAcore Reduced Aversion-Resistant Alcohol Consumption

We recently showed that inhibiting NAcore HA-NMDARs with the NMDAR blocker AP5, or by shRNA knockdown of the HA-NMDAR subunit GluN2c, selectively reduces aversion-resistant alcohol drinking (Seif et al, 2013). Thus, we locally infused D-serine (50 or 100 µg/side; Fiorenza et al, 2012; Kretschmer and Schmidt, 1996) to determine whether the NAcore could mediate systemic D-serine effects on drinking. Intra-NAcore D-serine significantly reduced intake of guinine-adulterated alcohol but not guinine-free alcohol (Figure 2a), without significant changes in preference for the alcohol-quinine solution (Figure 2c and d); the lack of significant changes in preference likely reflects the low water intake during the 20 min intake session. D-serine decreased consumption of quinine-adulterated alcohol by $39.9 \pm 11.9\%$ (for 50 μ g) and 61.5 \pm 7.6% (for 100 μ g) relative to quininefree alcohol, indicating a substantial effect of NAcore D-serine on aversion-resistant intake. Alcohol drinking the day after testing did not differ among groups, indicating no lasting effects of D-serine on intake (Figure 2e). Also, in contrast to its effects in the NAcore, bilateral D-serine injection within the dorsolateral striatum (DLS) (50 µg/side) had no effect on intake of alcohol ± quinine (Figure 2f and g). Thus, the selective impact of systemic D-serine on aversion-resistant alcohol drinking likely required the NAcore but not the DLS.

D-Serine Reduced NMDAR Function at -70 mV Resting Potential *In vitro*

HA-NMDARs in the NAcore of alcohol-drinking rats promote aversion-resistant alcohol intake (Seif *et al*, 2013). As D-serine can inhibit HA-NMDARs (Chatterton *et al*, 2002; Takarada *et al*, 2009, 2012), we determined whether D-serine inhibited NAcore HA-NMDARs *in vitro* in alcoholdrinking rats. In particular, we examined EPSCs generated at -70 mV by ChR2 stimulation of mPFC–NAcore inputs

D-serine and D-cycloserine reduce compulsive alcohol intake T Seif et al



Figure 2 D-serine within the NAcore reduced aversion-resistant alcohol intake. (a) Infusion of D-serine within the NAcore (50 or 100 μ g/side) significantly reduced consumption of quinine-adulterated alcohol but not quinine-free alcohol (n = 9) (quinine: $F_{1,16} = 13.761$, p = 0.006; D-serine: $F_{2,16} = 7.543$, p = 0.005; interaction: $F_{2,16} = 3.917$, p = 0.041). *p < 0.05 for no quinine vs quinine conditions. (b) Histological placement of cannulae targeting the NAcore. Co, NAcore; Sh, NAcb shell. (c, d) Decreased quinine-adulterated alcohol drinking was not accompanied by changes in water intake (c) (all p > 0.1) or alcohol preference (d), although quinine did decrease preference (quinine: $F_{1,16} = 13.275$, p = 0.007; D-serine: $F_{2,16} = 0.444$, p = 0.788; interaction: $F_{2,16} = 2.173$, p = 0.146). (e) Intra-NAcore D-serine did not reduce alcohol intake the day after D-serine treatment (all p > 0.3). (f) Infusion of D-serine within the DLS (50 μ g/side) had no impact on quinine-free or quinine-adulterated alcohol intake (n = 8) (all p > 0.15). (g) Histology of cannulae targeting the DLS. Numbers in (b) and (g) indicate distance from Bregma. D-seri, veh, vehicle.

without the AMPAR blocker DNQX, as in Seif *et al* (2013). Under these conditions, nearly all current is mediated by AMPARs (Seif *et al*, 2013), as canonical NMDARs are mostly inactive at hyperpolarized membrane potentials (Cull-Candy and Leszkiewicz, 2004). However, HA-NMDARs, when present, can be detected by a decrease in EPSC amplitude when adding NMDAR blockers such as AP5 (Seif *et al*, 2013). As HA-NMDAR currents are small, and direct measurement is technically challenging, we used EPSC

changes at -70 mV as a more robust indicator of the presence of HA-NMDARs (Seif *et al*, 2013).

D-serine (100 μ M, Chatterton *et al*, 2002; Junjaud *et al*, 2006; Wu *et al*, 1994) significantly reduced mPFC-evoked EPSCs in NAcore neurons from alcohol-drinking rats, with no effect in age-matched, alcohol-naive controls (Figure 3a and b). In contrast, D-serine had no impact on mPFC-ChR2-evoked EPSCs within the DLS (Figure 3c). In addition, we previously showed that NMDAR blockers do

2361



Figure 3 D-serine inhibited EPSCs evoked at -70 mV under mPFC-NAcore terminals in alcohol-drinking rats. (a) Cartoon showing that a 480 nm LED only evoked glutamate release (small black circles) from ChR2-filled mPFC terminals within the NAcore (dark gray). HC, hippocampus. (b) Sample traces and grouped data demonstrating that D-serine (100μ M) significantly reduced mPFC-ChR2-evoked EPSCs at -70 mV in neurons from alcohol-drinking rats (n = 9; $-32.9 \pm 10.1\%$ vs baseline) but not naive rats (n = 8; $-4.0 \pm 4.2\%$ vs baseline). Mann–Whitney test, U = 10; p = 0.011 alcohol vs control, *p < 0.05 alcohol vs control. (c) D-serine (100μ M) did not reduce mPFC-evoked EPSCs at -70 mV in the DLS of alcohol drinkers (n = 5; $-7.4 \pm 6.2\%$ vs baseline, $t_4 = 1.345$, p = 0.250 paired *t*-test). (d) Cartoon showing that electrical stimulation (lightning) evoked glutamate release (small black circles) from terminals originating from all regions (dark gray); only some of the known glutamate inputs to NAcore (Gerfen, 2004) are shown. (e) D-serine did not reduce electrically evoked EPSCs at -70 mV in NAcore neurons from alcohol-drinking (n = 6; $3.0 \pm 3.3\%$ vs baseline) or control rats (n = 9; $-1.8 \pm 5.5\%$ vs baseline). (f) D-serine enhanced NAcore NMDARs evoked at +40 mV, with no difference between alcohol and control (n = 4 alcohol, 4 control; $t_6 = 0.313$, p = 0.765). Ampl, amplitude.

not decrease EPSCs evoked by electrical field stimulation (Seif *et al*, 2013). Electrical stimulation activates all glutamatergic inputs, whereas HA-NMDARs in alcoholdrinking rats are present only under cortical-NAcore inputs (Seif *et al*, 2013), which are a small subset of the many glutamatergic inputs onto NAcore neurons (Gerfen, 2004) (Figure 3d). As predicted, D-serine did not alter electrically evoked EPSCs at -70 mV (Figure 3e). However, D-serine did enhance mPFC-evoked NAcore NMDAR currents at +40 mV (Figure 3f), showing that D-serine can exert its canonical action in the NAcore (Curcio *et al*, 2013), and the D-serine increase in NMDAR currents was not different between alcohol and control neurons (Figure 3f). Smaller D-serine enhancement of +40 mV NMDAR currents (Figure 3f) relative to that reported for NAc (Curcio *et al*, 2013) may reflect differences between adult (this study) and juvenile (Curcio *et al*, 2013). Most importantly, D-serine reduced the amplitude of mPFC-NAcore EPSCs evoked at hyperpolarized potentials only in alcohol-drinking rats.



Figure 4 D-serine inhibition of EPSCs occluded the ability of the NMDAR blocker AP5 to inhibit EPSCs. (a) Sample traces and (b, c) grouped data showing that D-serine significantly reduced ChR2-evoked EPSCs evoked at -70 mV ($-27.9 \pm 6.0\%$ vs baseline) and, importantly, occluded AP5 from producing any further decrease in EPSCs (n = 6), suggesting that D-serine acted by inhibiting HA-NMDARs. F_{2,10} = 10.15, p = 0.004, Friedman repeated-measures ANOVA on ranks, *post hoc* Holm–Sidak test *p < 0.05 for baseline vs D-serine vs D-serine +AP5; NS, for D-serine vs D-serine+AP5. D-ser, D-serine.

To examine whether D-serine reduced mPFC–NACoreevoked EPSCs through inhibition of HA-NMDARs, we investigated whether D-serine could occlude the previously described AP5 inhibition of mPFC–NAcore EPSCs at – 70 mV in alcohol-drinking rats (Seif *et al*, 2013). D-serine significantly reduced the amplitude of mPFC-evoked EPSC, and adding AP5 after D-serine caused no further inhibition (Figure 4). Also, AP5 alone reduced EPSCs by – 32.4 ± 8.9% (data from Seif *et al*, 2013), whereas AP5 added after D-serine produced no change in EPSCs ($1.2 \pm 10.2\%$ change *vs* D-serine; $t_9 = 2.675$, p = 0.025 AP5 alone *versus* AP5 after D-serine). This D-serine occlusion of the effects of AP5 indicates that D-serine reduced mPFC-evoked EPSCs in alcohol-drinking rats by inhibiting HA-NMDARs.

D-Cycloserine Reduced Aversion-Resistant Alcohol Drinking

DCS is another FDA-approved, glycine-site modulator of NMDARs (Heresco-Levy et al, 2013; Tsai and Lin, 2010), and thus could modulate HA-NMDARs and represent a therapeutic intervention to reduce compulsion-like alcohol drinking. DCS (10 mg/kg, Moraes Ferreira and Morato, 1997; Torregrossa et al, 2010; Yaka et al, 2007) significantly reduced quinine-resistant alcohol intake, with no impact on quinine-free alcohol drinking (Figure 5a) or consumption of saccharin \pm quinine (Figure 5b). Thus, DCS selectively reduced aversion-resistant alcohol drinking, similar to D-serine (Figures 1 and 2) and other NMDAR inhibitors (Seif et al, 2013), but did not alter concurrent water intake (Figure 5c), alcohol preference (Figure 5d), or alcohol intake the following day (Figure 5e). In addition, in vitro DCS (10 µM, Henderson et al, 1990; Rouaud and Billard, 2003) significantly reduced mPFC-ChR2-evoked EPSCs in the NAcore at -70 mV in alcohol-drinking but not in control rats (Figure 5f) $(t_{19} = 3.507, p = 0.002$ alcohol vs control), with no effect on electrically evoked EPSCs (Figure 5g). These results are similar to those for D-serine and other NMDAR blockers, suggesting that DCS can inhibit HA-NMDARs under mPFC-NAcore inputs in alcoholdrinking rats and, in this way, suppress compulsion-like alcohol intake.

DISCUSSION

Our results suggest that D-serine and DCS represent novel therapeutic interventions to counteract the aversion-resistant, compulsive drives for alcohol that are a prominent feature of AUDs and a major obstacle to successful treatment. Systemic D-serine and DCS significantly reduced aversion-resistant alcohol intake, with no effect on concurrent water intake or drinking of quinine-free alcohol or saccharin \pm quinine. This suggests that decreased aversion-resistant drinking did not reflect changes in palatability or motor activity, in agreement with no effect of D-serine administered within the NAcore or systemically on locomotor activity or coordination (Curcio et al, 2013; Kretschmer and Schmidt, 1996; Lockridge et al, 2012). A selective effect on aversion-resistant alcohol intake also agrees with the lack of effect of D-serine and DCS on alcohol intoxication (Debrouse et al, 2013), and with our previous results (Seif et al, 2013) and theories proposed by several groups (Naqvi et al, 2014; Tiffany and Conklin, 2000) that conflict-processing cortical regions are recruited during compulsive intake, whereas noncompulsive intake has less requirement for cortical involvement (see Introduction). Importantly, D-serine within the NAcore but not DLS selectively reduced aversion-resistant alcohol drinking, identifying the NAcore as a critical region where D-serine regulates aversion-resistant consumption. Finally, we provide in vitro evidence that D-serine and DCS inhibited NAcore HA-NMDARs in alcohol-drinking rats, similar to other studies where D-serine inhibited HA-NMDARs (Chatterton et al, 2002; Takarada et al, 2009, 2012). D-serine and DCS had no effect on electrically evoked EPSCs, indicating that they did not affect AMPAR EPSCs in general. These results all agree with our previous work showing that NAcore HA-NMDARs promote aversion-resistant alcohol drinking (Seif et al, 2013), and indicate that D-serine and DCS might be immediately accessible pharmacotherapies for reducing the compulsion to drink alcohol.

Long-term alcohol drinking leads to appearance of HA-NMDARs under cortical-NAcore inputs that promote aversion-resistant intake (Seif *et al*, 2013). Several lines of evidence suggest that D-serine inhibits these HA-NMDARs. Both D-serine (this study) and the NMDAR blocker AP5



Figure 5 D-cycloserine reduced aversion-resistant alcohol intake and EPSCs evoked at -70 mV *in vitro* under mPFC-NAcore terminals. (a) DCS (i.p., 10 mg/kg) reduced drinking of quinine-adulterated alcohol but not quinine-free alcohol (n = 13; quinine: $F_{1,12} = 3.501$, p = 0.086; D-serine: $F_{1,12} = 1.62$, p = 0.162; interaction: $F_{1,12} = 4.811$, p = 0.049). *p < 0.05 for DCS vs all other conditions. (b) Systemic DCS did not alter intake of saccharin \pm quinine (n = 14; saccharin-quinine doses: $F_{2,26} = 5.78$, p = 0.008; D-serine: $F_{1,26} = 0.001$, p = 0.996; interaction: $F_{2,26} = 0.088$, p = 0.916). (c-e) DCS reduction of aversion-resistant drinking was not accompanied by changes in concurrent water intake (c) (all p > 0.1), alcohol preference (d) (all p > 0.4), or alcohol intake the next day (e) (all p > 0.7). (f) DCS (10μ M) significantly reduced mPFC-ChR2-evoked EPSCs at $-70 \,$ mV in neurons from alcohol-drinking rats (n = 12; $-21.6 \pm 3.8\%$ vs baseline) but not naive rats (n = 9; $-5.2 \pm 3.8\%$ vs baseline). *p < 0.05 alcohol vs control. (g) DCS did not reduce electrically evoked EPSCs at $-70 \,$ mV in NAcore neurons from alcohol-drinking (n = 7; $-8.1 \pm 5.5\%$ vs baseline) or control rats (n = 14; $-3.8 \pm 6.4\%$ vs baseline). Veh, vehicle.

(Seif *et al*, 2013) within the NAcore reduce aversion-resistant but not quinine-free alcohol drinking. Similarly, both Dserine and DCS (this study) and AP5 (Seif *et al*, 2013) decrease mPFC-ChR2-evoked EPSCs at -70 mV in the NAcore of alcohol-drinking rats, with no effect in controls. Importantly, the *in vitro* D-serine inhibition of mPFC-NAcore EPSCs prevented AP5 from further decreasing mPFC-evoked EPSCs; this occlusion indicates that D-serine reduces EPSCs at -70 mV by inhibiting HA-NMDARs. Although D-serine is an NMDAR activator at the glycine site, it can also inhibit HA-NMDARs containing the GluN3 subunit (Chatterton *et al*, 2002; Takarada *et al*, 2009, 2012). Thus, NAcore HA-NMDARs in alcohol drinkers may contain GluN3 in addition to the HA-NMDAR subunit GluN2C described in Seif *et al* (2013). Furthermore, NMDARs containing only GluN1/3 but not GluN2 subunits are not AP5 sensitive (Smothers and Woodward, 2007), but NAcore HA-NMDARs in alcohol drinkers are blocked by AP5, consistent with the presence of GluN2C (Seif *et al*, 2013). Also, the $100 \,\mu$ M p-serine utilized here *in vitro* has

Neuropsychopharmacology

been used for testing effects of saturating concentrations of D-serine at the NMDAR glycine site (Zhang et al, 2008; Curcio et al, 2013), but GluN1/2D/3A HA-NMDARs are inhibited by D-serine only at higher concentrations ($\geq 100 \,\mu$ M, Takarada et al, 2009, 2012). Similarly, intra-NAcore D-serine can enhance NMDAR function at 4 µg (Curcio et al, 2013), but inhibited aversion-resistant alcohol drinking at 50-100 µg (Figure 2) but not 25 µg (not shown). Although dialysis studies estimate free D-serine in the forebrain at $\sim 6-8 \,\mu M$ (Matsui et al, 1995; Ciriacks and Bowser, 2006), basal levels could be several fold higher, and reach hundreds of µM with exogenous D-serine concentrations applied here (Hashimoto and Chiba, 2004; Zhang et al, 2008; Nishikawa, 2011). Together, these results support the idea that NAcore HA-NMDARs are GluN1/2C/3 heterotrimers. Overall, relatively little is known about such heterotrimers, although GluN2 subunits certainly alter GluN1/3 NMDAR properties (Chatterton et al, 2002; Das et al, 1998; Smothers and Woodward, 2007). Also, although the primary focus of our studies is the possible use of D-serine and DCS to reduce compulsion-like alcohol drinking, and we have demonstrated here that exogenous D-serine significantly and selectively reduced aversion-resistant intake in rats and thus may represent a novel intervention to suppress compulsive aspects of human AUDs, it would be interesting in future experiments to examine possible changes in endogenous D-serine regulation in alcohol-drinking individuals as it could interact with exogenous D-serine.

NMDAR modulators have long been considered as potent therapeutic agents to treat conditions ranging from schizophrenia, depression, and Parkinson's disease to drug and alcohol addiction (D'Ascenzo et al, 2014; Gass and Olive, 2008; Heresco-Levy et al, 2013; Krystal et al, 2003, 2011; Martineau et al, 2014; Sani et al, 2012; Spanagel, 2009; Tsai and Lin, 2010). However, previous studies with NMDAR antagonists have observed somewhat mixed results on alcohol-related behaviors in humans (Gass and Olive, 2008). Memantine, a noncompetitive NMDAR antagonist, has shown promise in some studies (Bisaga and Evans, 2004; Krupitsky et al, 2007; Muhonen et al, 2008) but not others (Evans et al, 2007). Acamprosate reduces alcohol drinking in some individuals, but can act on molecular targets other than NMDARs (Gass and Olive, 2008; Spanagel et al, 2014). D-serine, DCS, and related compounds are of particular interest because of their widespread use and a favorable safety profile, and also the moderate modulatory effects of such compounds that avoids more severe side effects of some NMDAR modulators (Heresco-Levy et al, 2013; Millan, 2005; Tsai and Lin, 2010). We note that nephrotoxicity is a potential concern with long-term, high-dose treatment of D-serine (Krug et al, 2007), but several longer-term D-serine studies have been performed in humans without adverse effects noted (D'Ascenzo et al, 2014; D'Souza et al, 2013; Heresco-Levy et al, 2013; Kantrowitz et al, 2010; Tsai and Lin, 2010). In addition, D-serine and DCS can enhance NMDAR activity and increase memory (Heresco-Levy et al, 2013; Martineau et al, 2014; Sani et al, 2012; Tsai and Lin, 2010). The interaction of this effect with HA-NMDAR inhibition may be complex, but could be synergistic, where D-serine or DCS increases memory for the suppression of compulsive drives for alcohol.

2365

D-serine and DCS reduction of aversion-resistant alcohol intake did not impact alcohol drinking the next day. In agreement, suppressing aversion-resistant intake (by inhibiting mPFC-to-NAcore inputs) did not reduce alcoholquinine intake the following day (Seif et al, 2013). Thus, we consider HA-NMDAR-mediated interventions useful for crisis management to acutely reduce compulsive drives for alcohol as an important part of a more comprehensive therapeutic approach including behavioral therapy and counseling. Also, although we have not directly tested whether D-serine regulates other forms of compulsion-like intake, there are several reasons to consider that it can. Most importantly, a similar cortico-NAcore circuit mediates both quinine-resistant and footshock-resistant alcohol drinking (Seif et al, 2013), and footshock-resistant intake for other rewards such as cocaine has been considered a robust model for aversion-resistant, compulsive aspects of human addiction (Deroche-Gamonet and Piazza, 2014; Everitt and Robbins, 2005). Furthermore, as HA-NMDARs regulate aversion-resistant drinking and are selectively inserted at cortical-NAcore synapses, the HA-NMDAR/cortico-NAcore pathway may represent a common mechanism to promote aversion-resistant alcohol drinking, with no impact on water or saccharin \pm quinine intake. Also, the BAC concentrations observed here (40-50 mg%, see also Seif et al, 2013; Simms et al, 2008) were below the level considered for binge intake (>80 mg%) which perhaps raises questions about the 'compulsivity' of the alcohol drinking we have studied. However, the similar circuit mechanism for quinine-resistant and especially footshock-resistant alcohol intake (Seif et al, 2013) suggests that rats drinking under the intermittent access model exhibited compulsion-like drinking. Our drinking paradigms are designed to examine consumption that persists despite the presence of an aversive consequence; such paradigms have been considered to model compulsive aspects of human AUDs (Everitt and Robbins, 2005; Hopf and Lesscher, 2014; Hopf et al, 2010; Lesscher et al, 2010; Seif et al, 2013; Vengeliene et al, 2009). In addition, the intermittent alcohol-drinking model exhibits other features that may mirror those seen in humans AUDs, including escalation of intake and sensitivity to compounds that can reduce human alcohol drinking (Hopf and Lesscher, 2014; Koob and Volkow, 2010; Simms et al, 2008). Also, other methods of quinine-resistant alcohol drinking with more moderate intake are associated with additional features sometimes observed with human addiction, including disrupted circadian patterns of intake and increased preference for higher concentrations of alcohol (Vengeliene et al, 2009; Wolffgramm et al, 2000). Thus, even with BAC levels of 40–50 mg%, there are many features of the intermittent access to alcohol and aversion-resistant drinking paradigms that provide reasonable confidence that these may model some aspects of pathological human alcohol drinking.

We note that there was an interaction trend (Figure 1e) and main effect of D-serine (Figure 1f) on consumption of saccharin, and thus some of the effect of D-serine on decreasing compulsion-like alcohol drinking may have occurred through nonspecific impacts on fluid intake. Future experiments with a broader range of D-serine doses may clarify this possibility. However, D-serine did not reduce intake of lower concentrations of saccharin+quinine, where drinking volume was more similar to those during alcohol intake. Also, DCS systemically had a similar, selective effect on aversion-resistant alcohol drinking as observed with D-serine, with little change in intake of saccharin+quinine, although a limited dose range of DCS was tested. Overall, our results support the possibility that D-serine and DCS selectively inhibited compulsion-like alcohol drinking.

In summary, our results suggest that D-serine and DCS suppressed aversion-resistant alcohol consumption through inhibition of HA-NMDARs within the NAcore. These findings identify D-serine and DCS as novel, immediately accessible and FDA-approved therapeutic interventions to counteract the compulsive aspects of alcohol drinking that represent a major clinical obstacle to successful treatment of AUDs.

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The authors declare no conflict of interest.

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