Chronic ethanol and withdrawal effects on kainate receptor–mediated excitatory neurotransmission in the rat basolateral amygdala

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Abstract

Withdrawal (WD) anxiety is a significant factor contributing to continued alcohol abuse in alcoholics. This anxiety is extensive, long-lasting, and develops well after the obvious physical symptoms of acute WD. The neurobiological mechanisms underlying this prolonged WD-induced anxiety are not well understood. The basolateral amygdala (BLA) is a major emotional center in the brain and regulates the expression of anxiety. New evidence suggests that increased glutamatergic function in the BLA may contribute to WD-related anxiety following chronic ethanol exposure. Recent evidence also suggests that kainate-type ionotropic glutamate receptors are inhibited by intoxicating concentrations of acute ethanol. This acute sensitivity suggests potential (KA-R) contributions by these receptors to the increased glutamatergic function seen during chronic exposure. Therefore, we examined the effect of chronic intermittent ethanol (CIE) and WD on KA-R-mediated synaptic transmission in the BLA of Sprague–Dawley rats. Our study showed that CIE, but not WD, increased synaptic responses mediated by KA-Rs. Interestingly, both CIE and WD occluded KA-R-mediated synaptic plasticity. Finally, we found that BLA field excitatory postsynaptic potential responses were increased during CIE and WD via a mechanism that is independent of glutamate release from presynaptic terminals. Taken together, these data suggest that KA-Rs might contribute to postsynaptic increases in glutamatergic synaptic transmission during CIE and that the mechanisms responsible for the expression of KA-R–dependent synaptic plasticity might be engaged by chronic ethanol exposure and WD. © 2009 Elsevier Inc. All rights reserved.

Keywords: Basolateral amygdala; Electrophysiology; Kainate; Anxiety; Withdrawal

Introduction

Withdrawal (WD) from chronic alcohol exposure causes long-term alterations in numerous behavioral outcomes. Although several of these alterations are believed to contribute significantly to treatment efficacy in human alcoholics, WD-related anxiety makes a significant contribution to relapse in humans (Cohn et al., 2003; Linnoila, 1989; Lucht et al., 2002; Verheul et al., 2005). The neurophysiological mechanisms governing these long-term changes in anxiety-related behaviors are not well understood. Importantly, numerous rodent models of chronic ethanol exposure have long-term changes in anxiety-like behavior during WD as a common characteristic (Borlikova et al., 2006; Breese et al., 2005; Klithermes, 2005; Lack et al., 2007; Santucci et al., 2008; Valdez et al., 2002; Zhao et al., 2007). The neural circuitry and neurophysiological mechanisms governing anxiety-like behavior in rodents have been relatively well studied. Importantly, it has recently been demonstrated that these same brain areas appear to mediate increased anxiety following WD from chronic ethanol exposure (Funk and Koob, 2007; Knapp et al., 2007; Lack et al., 2007).

The basolateral amygdala (BLA) is a central component of the neural circuitry governing anxiety-related information and is also involved with alcohol WD-related behaviors. The flow of information through the amygdala starts with cortical and thalamic input into the lateral and basolateral nuclei, then proceeds from there through efferent projections to the central nucleus of the amygdala, bed nucleus of the stria terminalis, and the nucleus accumbens (De Olmos et al., 1985). Alcohol interacts with neurotransmitter systems mediating both fast excitatory and inhibitory synaptic transmission in the amygdala. Importantly, the balance of these neurotransmitter systems can be disrupted to alter anxiety-related behavior (Lack et al., 2007; Sanders and Shekhar, 1995). Recent work has shown that chronic intermittent ethanol (CIE) and WD produce a significant
upregulation of the glutamatergic system in the BLA that may ultimately influence anxiety-like behavior (Lack et al., 2007).

There are three major subtypes of ionotropic glutamate receptors: NMDA, AMPA, and kainate-type (KA-R). Although we are beginning to understand the effects of alcohol on BLA, AMPA, and NMDA receptors, very little is known about the effects of alcohol exposure on KA-R glutamate receptors. These receptors are structurally similar to the better-studied AMPA-type glutamate receptors and are typically found in both pre- and postsynaptic compartments throughout the forebrain (Jin et al., 2006; Weiner et al., 1999; West et al., 2007). KA-Rs can consist of five potential subunit combinations. GLUR 5 et al., 1999; West et al., 2007). KA-Rs can consist of five potential subunit combinations. GLUR5–7 are needed for functional channels and the KA1–2 subunits confer unique pharmacological and biophysical properties to the native receptor. GLUR3,6 and KA2 subunits are highly expressed in the BLA, where they contribute to glutamatergic excitatory postsynaptic potentials (EPSPs) in this brain region (Li and Rogawski, 1998).

Until recently, the close similarities between AMPA- and KA-R glutamate receptors had prevented a systematic understanding of the distinct roles these receptors play in various physiological processes. However, the development of selective antagonists allows the separation of AMPA- and KA-R-mediated responses (reviewed in Pinheiro and Mulle, 2006). For example, we now know that KA-Rs contribute to a significant postsynaptic response in the BLA (Li and Rogawski, 1998) and also mediate a form of long-lasting heterosynaptic plasticity in this brain region (Li et al., 2001). Recent evidence also suggests that the acute sensitivity of KA-Rs to ethanol may play a prominent role in regulating plastic changes in BLA synaptic transmission (Lack et al., 2008). Furthermore, our data with the GluR2-selective antagonist UBP296 support suggestions that GluR2-containing KA-Rs in the BLA may be potential therapeutic targets for anxiety-related behaviors (Aroniadou-Anderjaska et al., 2007) and provide strong rationale for determining the effects of CIE and WD on KA-Rs in the BLA.

Materials and methods

Animals

All animal procedures were performed in accordance with protocols approved by the Wake Forest University School of Medicine Animal Care and Use Committee and were consistent with the National Institutes of Health Guide for the Care and Use of Laboratory Animals. Male Sprague–Dawley rats (120–140 g) were obtained from Harlan at the beginning of the experiments described in the following section. All animals were housed in an animal care facility at 23°C with a 12-h light/dark cycle and given food and water ad libitum. Rats were weighed daily to ensure that at least 80% of their free-feeding weight was maintained during vapor chamber ethanol exposure.

Chronic ethanol exposure

Ethanol exposure was accomplished via an ethanol vapor chamber similar to that used in other studies (Lack et al., 2007). Briefly, animals in their home cages were placed into airtight, Plexiglas enclosures and exposed to either ethanol vapor (~33 mg/L) or room air (CON) during the light cycle (12 h/day) for 10 consecutive days. Animals receiving the chronic intermittent treatment were divided into those euthanized while still intoxicated at the end of the last exposure (CIE) and those euthanized 24 h after the last exposure (WD). Tail blood was taken periodically during the exposure to monitor blood–ethanol concentration and trunk blood was collected on the day of sacrifice from subjects in the CIE group. Blood ethanol levels at the time the CIE animals were sacrificed were 225 ± 32 mg/dL.

Preparation of brain tissue

Animals were anesthetized with isoflurane and euthanized by decapitation. Coronal brain slices (400 μm) were prepared as described previously (Lack et al., 2007) using modified aCSF (in mM: 180 sucrose, 30 NaCl, 4.5 KCl, 1 MgCl2, 6H2O, 26 NaHCO3, 1.2 NaH2PO4, and 10 glucose) containing 100-μM ketamine. Slices from all experimental groups were transferred and stored in 0.5-L standard oxygenated aCSF solution (in mM: 126 NaCl, 3 KCl, 1.25 NaH2PO4, 2 MgSO4, 26 NaHCO3, 10 glucose, and 2 CaCl2) at room temperature for at least 1 h and up to 6 h.

Whole cell patch clamp electrophysiology

Methods for whole-cell “blind” patch-clamp recordings from principle BLA projection neurons within slices were similar to those reported previously (Lack et al., 2007). Electrodes were filled with an intracellular pipette solution containing the following (in mM): 122 CsOH, 17.5 CsCl, 1 MgCl2, 6H2O, 26 NaHCO3, 1.2 NaH2PO4, and 10 glucose) containing 100-μM ketamine. Slices from all experimental groups were transferred and stored in 0.5-L standard oxygenated aCSF solution (in mM: 126 NaCl, 3 KCl, 1.25 NaH2PO4, 2 MgSO4, 26 NaHCO3, 10 glucose, and 2 CaCl2) at room temperature for at least 1 h and up to 6 h. Compound glutamatergic events were pharmacologically isolated using 10 μM bicuculline to inhibit fast GABAergic transmission. In addition, kainate currents were isolated using 50-μM DL-2-amino-5-phosphono-pentanoic acid (APV), an NMDA receptor antagonist, and 50-μM GYKI 53655 (1-(4-aminophenyl)-3-methylcarbamyl-1-4-methyl-3, 4-dihydro-7,8-methylenedioxo-5H-2,3-benzodiazepine), an...
AMPA receptor–specific noncompetitive antagonist. Recordings were acquired with an Axoclamp 2B amplifier (Axon Instruments, Foster City, CA) and digitized with a Digidata 1200B (Axon Instruments).

**Paired pulse ratio**

The paired pulse ratio during whole-cell recordings was measured using pairs of electrical stimuli of equal intensity at 25, 50, or 250 m interstimulus intervals. Ratios of the amplitudes of the evoked EPSCs were calculated as the amplitude of the second event minus the amplitude of the first event divided by the amplitude of the first event. All values were expressed as mean ± standard error of mean (S.E.M.), and data were subjected to a one-way analysis of variance (ANOVA) and Newman–Keuls post-test with \( P < .05 \) considered statistically significant.

**Field EPSP recordings**

Slices were placed in the recording chamber and were continuously superfused in aCSF (2 mL/min) warmed to approximately 30°C (Lack et al., 2008). Electrodes were filled with an extracellular pipette solution containing 150-mM NaCl. Ensemble postsynaptic potentials were evoked every 30 s by brief (0.2 ms) square-wave electrical stimulation within the external capsule (EC) along the lateral border of the BLA. These synaptic responses are referred to as field excitatory postsynaptic potentials (fEPSPs) throughout the study and were completely sensitive to both the AMPA/KR antagonist DNQX (20 \( \mu \)M) and tetrodotoxin (1 \( \mu \)M; not shown). For the ATPA–induced plasticity experiments, maximal fEPSP responses were determined in every slice; the stimulus intensity that evoked 50% maximal amplitude was used for the baseline period. All field experiments were recorded in the presence of 10-\( \mu \)M bicuculline methiodide. Baseline fEPSPs were recorded for 10 min prior to application of the KA-R agonist ATPA ((RS)-2-amino-3-(3-hydroxy-5-tert-butylisoxazol-4-yl) propanoic acid; 5 \( \mu \)M) for 15 min; the slice was then washed for 40 min in drug-free aCSF. fEPSP recordings were acquired with an Axoclamp 2B amplifier (Axon Instruments, Foster City, CA) in the current-clamp mode and digitized with a Digidata 1200B (Axon Instruments). All values were expressed as mean percent control ± S.E.M. Data were subjected to a one-way ANOVA or a two-way ANOVA with \( P < .05 \) considered statistically significant.
Input—output recordings

Input/output recordings were measured for both KA-R mediated EPSCs and fEPSPs. For both types of recording, input/output values were collected using increasing stimulus intensities for each cell or slice. At each stimulus intensity, EPSC amplitude or fEPSP slope was measured and compared across the treatment conditions. All values were expressed as mean response area (for EPSCs) or percent control (for fEPSPs) ± S.E.M., and data were subjected to a two-way ANOVA with stimulus intensity and fEPSP slope as the main dependent variables with Bonferroni post hoc analyses; \( P < .05 \) was considered statistically significant.

Results

CIE, but not WD, increases KA-R function at BLA glutamatergic synapses

Previous research in our laboratory indicated that KA-Rs are inhibited by acute ethanol (Lack et al., 2008). These results suggested that KA-Rs might then be upregulated during CIE and WD in response to this acute sensitivity. Therefore, we determined the contribution made by KARs to glutamatergic synaptic responses during CIE and WD. To examine this, we used whole-cell patch clamp electrophysiology to measure the input/output (stimulus–response) relationship for KA-R via stimulation of the EC (Li and Rogawski, 1998; Li et al., 2001). Because KA-R-mediated responses are small relative to AMPA-mediated synaptic currents, we measured evoked kainate EPSCs using stimulus trains of three pulses (100 Hz, 10 ms interstimulus interval) in the presence of a blocker cocktail containing the NMDA receptor antagonist APV, the GABAA receptor antagonist bicuculline, and the noncompetitive AMPA receptor antagonist GYKI 53655 (Fig. 1A, C).

Across discrete stimulus intensities, we found that the charge carried by the GYKI-resistant, KA-R-mediated synaptic responses were significantly increased in CIE neurons, but not in cells recorded from CON and WD slices (Fig. 2B, C). Likewise, the amplitude of the GYKI-resistant KA-R synaptic response was significantly greater in CIE neurons than in either CON or WD neurons at both the 25-μA stimulus (CIE, 27.3 ± 7.7 pA, \( n = 7 \); CON, 10.1 ± 2.2 pA, \( n = 7 \); WD, 11.2 ± 1.5 pA, \( n = 8 \); \( P < .05 \) with one-way ANOVA and Newman–Keuls post-test) and the 35-μA stimulus intensity (CIE, 56.4 ± 12.2 pA; CON, 23.0 ± 4.5 pA; WD, 27.55 ± 5.64 pA; \( P < .05 \) with one-way ANOVA and Newman–Keuls post-test). These data suggest that the KA-R contribution to BLA synaptic responses following EC electrical stimulation is increased during CIE relative to CON and returns to CON levels during WD.

CIE and WD occlude ATPA—induced synaptic plasticity in the BLA

BLA KA-Rs have been shown to mediate a form of glutamatergic synaptic plasticity in principal neurons, potentially through recruitment of additional excitatory synapses (Li and Rogawski, 1998). We have recently shown that this synaptic plasticity, induced by the selective agonist ATPA, can be inhibited by both acute application of ethanol and the GluR3 competitive antagonist, UBP 296 ((RS)-1-(2-amino-2-carboxyethyl)-3-(2-carboxybenzyl)pyrimidine-2,4-dione; Lack et al., 2008). Given that CIE, but not WD, increased KA-R synaptic function, we hypothesized that ATPA—induced plasticity would be more readily established in CIE slices relative to both CON and WD. Following a 10 min baseline, we applied 5 μM ATPA for 15 min, followed by a 40 min washout period with drug-free aCSF. ATPA exposure caused a slowly developing increase in the fEPSP slope in CON slices (\( n = 10 \), Fig. 2A, B). This is consistent with previous findings (Lack et al., 2008; Li et al., 2001). Surprisingly, this time-dependent, ATPA—induced plasticity (Fig. 2B) was significantly attenuated in slices from both CIE (\( n = 8 \) and WD (\( n = 10 \)) with significant main effects of treatment (\( F = 45.94, P < .0001 \), two-way ANOVA), time (\( F = 11.73, P < .0001 \)), and a significant interaction between variables (\( F = 3.48, P < .0001 \)). Comparing these data from the baseline region (first 10 min) and the last 10 min of washout (Fig. 2C), ATPA increased fEPSP slope in CON slices (406.3% ± 29.11%) but was significantly less effective in slices from both CIE (177.5% ± 21.33%, \( P < .05 \), one-way ANOVA with Bonferroni’s post-test) and WD animals (207.4% ± 32.11%, \( n = 10 \), \( P < .05 \)). These data indicate that ATPA—induced synaptic plasticity at EC–BLA glutamatergic synapses is partially occluded by CIE and WD. Because KA-R synaptic function is upregulated in CIE slices (relative to control, Fig. 1), these data also suggest that the mechanisms involved with the expression of KA-R—dependent increases in synaptic function might be affected by these treatments.

CIE and WD increase field EPSP responses in the BLA

Previous studies have shown dramatic increases in glutamatergic function at local BLA synapses after CIE and WD (Lack et al., 2007). This suggests that CIE and WD might occlude ATPA—induced plasticity by engaging expression-associated mechanisms at EC–BLA synapses. To test this, we measured stimulus–response relationships for fEPSPs in BLA slices from CON–, CIE–, and WD–treated rats. We electrically stimulated the external capsule using a range of intensities, from 5 to 40 μA at intervals of 5 μA (Fig. 3A, B). fEPSP slope in CIE and WD slices significantly differed from CON at most stimuli >15 μA (Fig. 3A) with two-way ANOVA indicating significant main effects of treatment (\( F = 28.8, P < .0001 \)) and stimulation intensity (\( F = 19.5, P < .0001 \)) but no interaction between these factors. These results are complementary to previous findings showing that there is increased AMPA—type glutamate receptor function in the BLA in response to CIE treatment and 24 h of WD (Lack et al.,...
Importantly, the findings suggest that the occlusion of ATPA-induced plasticity is related to the increased synaptic responses in CIE and WD slices. These ethanol/WD-dependent effects closely parallel the cellular responses that characterize activity-dependent synaptic plasticity in this brain region.

**Increased synaptic responses in CIE and WD BLA does not involve presynaptic release-related mechanisms**

fEPSPs represent action potentials within a population of neurons. However, we previously showed that CIE and WD can increase presynaptic function at local BLA glutamatergic synapses (Lack et al., 2007). To examine whether similar effects might be responsible for increased synaptic responsiveness at EC—BLA synapses, we carried out a paired pulse facilitation experiment to measure presynaptic release probability indirectly with postsynaptic responses to pairs of electrical stimuli. We used interpulse intervals of 25, 50, and 250 ms to determine the ratio of the second synaptic response to the first across times that yield information about release probability (25 and 50 ms; Andreasen and Hablitz, 1994; Katz et al., 1993) as well as autoreceptor-mediated decreases in presynaptic function (250 ms; Brucato et al., 1992). We compared the ratios of the CIE— WD—treated slices to those of control slices. We found no significant difference (one-way ANOVA) between paired pulse ratios in the CIE (n = 10) or WD (n = 10) neurons compared with CON (n = 10) at any of the intervals tested (Fig. 4A, B). This suggests that the increased response at EC—BLA synapses seen in CIE and WD with the fEPSP measures does not involve a significant change in presynaptic release probability.

**Discussion**

Our research suggests that KA-Rs in the BLA, such as NMDA and AMPA receptors, actively participate in the overall increase in glutamatergic function in response to chronic ethanol exposure. Specifically, CIE increases KA-R-mediated synaptic responses in the BLA. In fact, this ethanol-dependent increase in KA-R function during CIE might contribute to glutamatergic synaptic plasticity during either CIE or WD. This is consistent with previous findings that repeated ethanol WD causes synaptic strengthening during which 5 μM ATPA was perfused on the slices. fEPSP slope from each slice was normalized to the mean CON values measured during the 10-min baseline period. *P < .05, **P < .01, ***P < .001; two-way ANOVA followed by Bonferroni’s test, CON relative to CIE (*) and WD (#). (C) Summary of ATPA—induced facilitation of the fEPSP slope. fEPSP slope is expressed as a percent increase during the 10-min baseline period measured during the last 10 min of the washout period. ATPA—induced increases in fEPSP slope were significantly attenuated by the CIE and WD treatments. *P < .05, one-way analysis of variance with Newman—Keuls post-test.
in the amygdala and occludes BLA long-term plasticity (Stephens et al., 2005). Saturation of long-term plasticity has been shown to impair fear learning (Moser et al., 1998) but leads to generalized increases in anxiety-like behavior. We have recently shown that BLA-dependent processes contribute to this WD-associated anxiety (Lack et al., 2007). Therefore, occlusion of ATPA-induced synaptic plasticity by CIE and WD may suggest that KA-R-mediated plasticity could contribute to WD-related anxiety.

CIE, but not WD, increases KA-R synaptic function in the BLA

Previous research in our laboratory showed that BLA KA-Rs and KA-R-mediated synaptic plasticity are inhibited by acute ethanol (Lack et al., 2008). This suggests that the upregulation of KA-R-mediated synaptic responses in chronic ethanol-exposed neurons arises from this acute sensitivity. Although the mechanism responsible for this upregulation is not currently known, it was apparently transient because KA-R-mediated synaptic responses returned to baseline following 24 h of WD. This rapid return would suggest that changes in gene or protein expression levels are less-likely potential mechanisms. Unfortunately, there is little guidance in the literature with respect to the effects of chronic ethanol/WD on KA-Rs. Previous findings have been inconsistent and shown either no effect of chronic ethanol on KA-R subunit protein levels (Chandler et al., 1999; Ferreira et al., 2001) or up-regulation during WD (Carta et al., 2002). These contrasting findings likely reflect the diversity of experimental approaches (e.g., native vs. in vitro systems) and ethanol exposures.

The behavioral consequences of increased KA-R-mediated synaptic function during chronic intermittent ethanol exposure are not precisely defined. However, we have previously shown that anxiety-like behavior is not increased in CIE animals (Lack et al., 2007). This suggests two possible interpretations: (1) CIE-dependent increases in KA-R function have no behavioral relevance or (2) KA-R-dependent contributions during CIE are not behaviorally manifest until WD. Supporting this latter interpretation, ATPA microinjection into ethanol-naïve BLA increases anxiety-like behavior (Lack et al., 2008). Because AMPA—, NMDA—, and KA-R—dependent synaptic function is increased in both CIE and, in some cases, WD animals, the absence of any significant increase in anxiety-like behavior in CIE animals presumably reflects ethanol-sensitive contributions by other neurotransmitter systems.

CIE and WD occlude ATPA—induced synaptic plasticity in the BLA

The KA-R agonist ATPA has been shown to increase synaptic strength in BLA principal neurons via an unknown, calcium-dependent mechanism (Li et al., 2001). However, in the CA1 region of the hippocampus, long-term ATPA application led to an enduring increase in the number of glutamatergic synapses in that region (Vesikansa et al., 2007). Using 5 μM ATPA, a concentration that does not affect AMPA receptors (Stensbol et al., 1999), we were able to replicate the ATPA—induced synaptic plasticity in this work. We have recently shown that this facilitation can be blocked

Fig. 3. CIE and WD increase fEPSP responses to electrical stimulation of EC—BLA synapses. (A) Input/output measurements of fEPSP slope versus stimulation intensity show significant increases after CIE and WD treatment across a range of intensities. (B) Exemplar traces show the increase in fEPSP slope for a representative control, CIE and WD slice and are averaged from 20 baseline traces (first 10 min) and the last 20 traces (last 10 min) of washout. (*# P < .05; ## P < .01, one-way *analysis of variance followed by Dunnett’s test, relative to control.)

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by GluR$_5$ antagonist UBP 296 and by acute ethanol (Lack et al., 2008). In this study, both CIE and WD occluded ATPA--induced synaptic plasticity in the BLA. This was an unexpected finding considering that the function of the KA-Rs was elevated in CIE neurons and went back to baseline during WD. One possible explanation is that extrasynaptic KA-R, and not the synaptic receptors measured (Fig. 1), are responsible for ATPA-induced plasticity. However, the initiation of synaptic plasticity at EC--BLA synapses by low-frequency electrical stimulation is dependent on synaptic activation of KA-Rs; and, this process parallels ATPA--induced plasticity in this same brain region (Li et al., 2001). An alternative explanation is that KA-R may be important only for the initiation of the synaptic plasticity at some undefined point during the CIE exposure. Regardless, we have recently shown that AMPA receptors are functionally upregulated during both CIE and WD (Lack et al., 2007). Because AMPA receptors have been shown to maintain both NMDAR--dependent and NMDAR--independent long-term synaptic potentiation in the hippocampus (Grover, 1998) and in the BLA (Maren, 2005; McKernan and Shinnick-Gallagher, 1997; Rogan et al., 1997), our findings suggest that CIE engages the mechanisms related to the expression/maintenance of synaptic plasticity, ultimately resulting in the occlusion of ATPA--induced plasticity at the time of our measures. This hypothesis parallels previous findings showing repeated ethanol/WD leading to the occlusion of NMDA--dependent forms of synaptic plasticity in the BLA (Stephens et al., 2005).

**CIE and WD increase fEPSP responses at EC--BLA synapses**

An alternative explanation for the occlusion of ATPA--induced plasticity during CIE and WD would be that these treatments have downregulated or blocked the mechanisms required for the expression of this type of “pharmacological LTP.” If this were the case, we would expect that the mechanisms associated with the expression of plasticity (e.g., AMPA receptors) would not be engaged by the treatments. However, our previous work has shown dramatic increases in AMPA receptor function after CIE and WD (Lack et al., 2007). Alternatively, the responses of BLA neurons themselves might be reduced during CIE and WD such that the increased AMPA receptor function was negated by reduced neuronal excitability. However, fEPSP responses measured after activation of the EC--BLA synapses were increased after CIE and WD. Thus, both glutamatergic synaptic transmission and the neurophysiological responses of BLA neurons to that transmission are increased during CIE and WD. These findings are inconsistent with any treatment-dependent inhibition of the mechanisms responsible for the expression of ATPA--related plasticity.

**CIE and WD do not affect presynaptic release probability at EC--BLA glutamatergic synapses**

Because our ATPA--fEPSP data suggested that the mechanisms responsible for the expression of synaptic plasticity were engaged by CIE and WD, we examined whether there was any treatment--specific alteration in presynaptic release of glutamate. This hypothesis was supported by our previous data that showed an increase in glutamate release from “local” glutamatergic synapses during CIE and WD (Lack et al., 2007). However, unlike at local glutamatergic synapses, we found no evidence of an increase in presynaptic probability of glutamate release at EC--BLA synapses after CIE and WD. The EC--BLA synapses contain cortical inputs for many divergent brain regions, including insular cortices (McDonald and Mascagni, 1996), the neighboring entorhinal cortex (McDonald and Mascagni, 1997), and executive-control areas, such as the anterior cingulate (McDonald and Mascagni, 1996). The “local stimulus” approach used in the Lack et al., 2007 study would have engaged these inputs as well as glutamatergic synapses arising from thalamus (Le Doux et al., 1991), hippocampus (Kishi et al., 2006), limbic and parietal cortex (McDonald and Mascagni, 1996), and intra--amygdala projections (Savander et al., 1995). The contrasting results found in this study and in the Lack et al. (2007) study suggest that the neural circuitry contributing to

![Fig. 4. Increased excitation of EC--BLA synapses was not associated with a change in paired pulse ratio. (A) Representative traces of paired pulse 50 responses for each treatment group (P > .05, one--way analysis of variance). (B) Bar graphs summarizing the effect of CIE and WD on the amplitude of pairs of AMPA EPSCs evoked at an interpulse interval of 25, 50, and 250 ms.](image-url)
distinct glutamatergic release pathways in the BLA might be differentially sensitive to chronic ethanol exposure and WD.

Conclusions

In summary, this work provides the first evidence of facilitation of KA-R synaptic function by chronic ethanol. This facilitation is contrasted with the occlusion of ATPA-mediated synaptic plasticity during both CIE and WD. The overall increase in neurophysiological responses during CIE and WD indicates that this occlusion results from a CIE/WD—dependent recruitment of the cellular mechanisms that govern the expression of synaptic plasticity within the BLA. This interpretation is supported by previous findings showing diminished activity-dependent synaptic plasticity in the BLA following alcohol exposure/WD (Stephens et al., 2005). With respect to downstream afferent areas, such as the central amygdala and the bed nucleus of the stria terminalis, the precise ramifications of this alcohol-dependent recruitment are not yet known. However, we hypothesize that the increased responsiveness of BLA neurons (evidenced by the fEPSP data) might be manifested as increases in glutamatergic transmission in these areas as well. This hypothesis is consistent with the increased anxiety-like behavior following ethanol exposure/WD (Lack et al., 2007). Likewise, the alcohol-dependent recruitment of mechanisms responsible for the expression of synaptic plasticity in the BLA is also consistent with the attenuation of amygdala-dependent fear learning following multiple WDs (Stephens et al., 2001). The present findings illustrate additional neurophysiological mechanisms by which chronic ethanol and WD influence anxiety-related neural circuitry.

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