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Velazquez-Marrero, Cristina M.; Wynne, Patricia M.; Bernardo, Alexandra; Palacio, Stephanie; Martin, Gilles E.; and Treistman, Steven N., "The relationship between duration of initial alcohol exposure and persistence of molecular tolerance is markedly nonlinear" (2011). GSBS Student Publications. 1743.  
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Neurobiology of Disease

The Relationship between Duration of Initial Alcohol Exposure and Persistence of Molecular Tolerance Is Markedly Nonlinear

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The neuronal calcium- and voltage-activated BK potassium channel is modulated by ethanol, and plays a role in behavioral tolerance in vertebrates and invertebrates. We examine the influence of temporal parameters of alcohol exposure on the characteristics of BK molecular tolerance in the ventral striatum, an important component of brain reward circuitry. BK channels in striatal neurons of C57BL/6j mice exhibited molecular tolerance whose duration was a function of exposure time. After 6 h exposure to 20 mM (0.09 mg%) ethanol, alcohol sensitivity was suppressed beyond 24 h after withdrawal, while after a 1 or 3 h exposure, sensitivity had significantly recovered after 4 h. This temporally controlled transition to persistent molecular tolerance parallels changes in BK channel isoform profile. After withdrawal from 6 h, but not 3 h alcohol exposure, mRNA levels of the alcohol-insensitive STREX (stress axis-regulated exon) splice variant were increased. Moreover, the biophysical properties of BK channels during withdrawal from 6 h exposure were altered, and match the properties of STREX channels exogenously expressed in HEK 293 cells. Our results suggest a temporally triggered shift in BK isoform identity. Once activated, the transition does not require the continued presence of alcohol. We next determined whether the results obtained using cultured striatal neurons could be observed in acutely dissociated striatal neurons, after alcohol administration in the living mouse. The results were in remarkable agreement with the striatal culture data, showing persistent molecular tolerance after injections producing 6 h of intoxication, but not after injections producing only 3 h of intoxication.

Introduction

The progression from recreational to compulsive alcohol use is influenced by many factors such as increased craving, loss of control, and acquired tolerance. Neural adaptations such as functional tolerance contribute to the development of alcoholism by either permitting or causing increased levels of alcohol consumption. These adaptations reduce the effects of alcohol, such that higher doses are required to maintain the same response. Importantly, longitudinal studies have shown that enhanced behavioral acute tolerance is one of the best predictors of future alcohol problems (Schuckit, 1994). Acquired alcohol tolerance and dependency have long been used as criteria for a diagnosis of alcoholism (Kalant, 1998).

Large conductance calcium-activated potassium (BK) channels may provide insights into the bridge between tolerance at the molecular and behavioral levels. BK has been shown to be a key component of behavioral tolerance to alcohol in invertebrates (Scholz et al., 2000; Ghezzi et al., 2004; Cowmeadow et al., 2005, 2006). In mammalian neurons, molecular tolerance of BK to alcohol includes reduced sensitivity to the drug, which occurs within minutes, and a slower developing declustering and internalization, measured in hours (Pietrzykowski et al., 2004). Use of genetically altered mice in which the β4 subunit of the BK channel is knocked out indicate that this subunit influences alcohol acute molecular tolerance at the single-channel and action potential levels, as well as influencing behavioral tolerance, and remarkably, the propensity of mice to drink alcohol. Thus, as in human studies, the presence of acute tolerance is a predictor of drinking behavior (Martin et al., 2008).

The transition from drug abuse to compulsive drug use is influenced by parameters of drug exposure (Ahmed and Koob, 1998), including concentration and temporal pattern (Lê and Kalant, 1992). For example, escalation of cocaine intake has been linked to drug exposure protocol in rats. Six hours appears to represent a critical time point, beyond which drug access leads to escalating drug intake (Ahmed and Koob, 1998). Here, we ask whether there are individual molecular transitions, selectively tripped by different temporal patterns of exposure, which influence the characteristics of BK molecular tolerance observed during withdrawal from alcohol. A number of reports in the literature describe changes in biochemical pathways involving, e.g., cyclic nucleotidare and CREB alterations, or in the persistent
activation of genes such as ΔFosB, that outlast the presence of the initiating drug of abuse, and so would qualify as drug-activated switches (Kelz et al., 1999; Nestler et al., 2001; Hyman et al., 2006; Renthal et al., 2008). In particular, we examine whether the transition observed in the persistence of alcohol molecular tolerance between 3 and 6 h of exposure might be related to a recently described mechanism whereby alcohol, via an epigenetic mechanism involving microRNA (miRNA), mediates a rapid reorganization of BK α isoforms (Pietrzykowski et al., 2008). Those isoforms that predominate after 6 h drug exposure are relatively resistant to alcohol’s actions. Our results suggest that a process of isoform reorganization does, indeed, represent an important component of the transition to persistent molecular tolerance (PMT) observed between 3 and 6 h exposure times.

Materials and Methods

**Primary striatal culture**

 Cultures of dissociated rat striatal neurons were prepared using a modification of a previously described protocol (Leveque et al., 2000). Briefly, postnatal day 8 Sprague Dawley rat pups were decapitated, and brains removed and immersed in ice-cold PBS. Next, the striatum was dissected out, transferred to HBSS (Invitrogen) with 10 mM HEPES, and dissociated using a fire-polished Pasteur pipette. Dissociated cells were transferred to the plating medium (a 1:1 mix of F12 nutrient mixture (Invitrogen) and DMEM medium supplemented with GlutaMAX; 2.0 mM glutamine final concentration), 2% fetal bovine serum (HyClone), 2% B-27 (Invitrogen), 1% penicillin/streptomycin (Invitrogen). Cells in the plating medium were seeded onto 35 mm Petri dishes (Nunc) coated successively with 0.01% (w/v) poly-L-ornithine (Sigma) and 33 μg/ml mouse laminin (Invitrogen). Twenty-four hours later, the medium was replaced with serum-free medium supplemented with 2% B-27, 2.0 mM glutamine and 1% penicillin/streptomycin. Thereafter, the medium was replaced every 3–4 d. Neuronal cultures were maintained in a 5% CO2, humidified incubator (99% relative humidity) at 37°C. All experiments were performed on neurons that were 14–21 d in culture.

For initial alcohol exposures, culture media was replaced with media containing 20 mM EtOH and incubated for 1, 3, or 6 h. For withdrawal periods, the dishes were washed 4 times with ethanol (EtOH)-free medium. Alcohol concentrations in the media at the end of the 1, 3, or 6 h exposure and after washing were measured using a GM7Analyser (Analox Instruments Inc.). Control dishes received media changes at the same time as the ethanol exposed dishes. Media changes did not have an effect on ethanol sensitivity of the channel. For electrophysiological experiments medium spiny neurons were identified by morphology (small to medium cell bodies and multiple thin processes) (Meredith et al., 1992).

**HEK cell transfection**

Cells from human embryonic kidney cell lines (HEK 293 cells) were cultured in DMEM medium supplemented with 10% heat-inactivated fetal bovine serum, 1% penicillin/streptomycin, and 25 mM HEPES (Invitrogen), and plated onto 25 cm² flasks. The cells were maintained in a 5% CO2, humidified incubator (99% relative humidity) at 37°C. Before transfection HEK293 cells were split from confluent cultures and plated on 60 mm Petri dishes. For all experiments, 50–70% confluent cells were transfected using PolyFect transfection reagent (Qiagen) complexed with BK α variant (BK-Insertless, BK-ALCOREX, BK-STREX) and BK-β4 subunit (when required) cDNAs in pVAX vector (Invitrogen) together with the expression plasmid (pTRE3-CD-8) encoding the α subunit of the human CD-8 lymphocyte surface antigen (GenBank M12824). To identify transfected cells, CD-8 antibody-coated beads were used (Dynal/Invitrogen). Before electrophysiological recordings, 0.5 μg/ml of the CD-8 antibody-coated beads was added to each culture dish for 30 min. Unattached beads were washed out with regular Locke’s solution and recordings obtained from clearly identified cells with attached beads.

**Electrophysiological recordings**

Recording electrodes were pulled on a horizontal puller (Sutter Instruments), coated with Sylgard (Dow Corning), and fire-polished from borosilicate thin-wall capillary glass (Drummond) to a final resistance of 4–8 MΩ. Currents were recorded in voltage-clamp mode with a HEKA EPC 10 amplifier at a sampling rate of 5 kHz and 10 kHz for whole-cell and single-channel recordings respectively, and low-pass filtered at 3 and 2 kHz, respectively. Leak currents were subtracted on-line using a conventional P/n (4) protocol. In whole-cell mode, series resistance did not exceed 20 MΩ and was 60% compensated. Potentials and currents were digitized and stored using PatchMaster acquisition and analysis software version 2.05 (HEKA Elektronik).

**Single-channel recordings.** Conventional cell-attached patch-clamp mode was used, to maintain the intracellular milieu and secondary signaling pathways during recordings to assess ethanol sensitivity of single channels. Control (no alcohol) and ethanol-containing solutions were contained in 50 ml syringes and expelled from hemocrit tubes. Each cell served as its own control. To determine baseline activity (defined here as open probability, NPo) before alcohol application, BK channel activity was recorded 3 times, 20 s each, at 1 min intervals. For alcohol perfusion, the ethanol containing hemocrit tube was positioned close to the cell. Then, alcohol was applied and BK channel activity recorded in successive blocks of 20 s at 1 min intervals for up to 10 min. Control experiments were performed in which patches were perfused with an ethanol-free solution for up to 15 min. In control experiments, there was no significant deviation in baseline channel activity throughout the 15 min recording period (data not shown).

Cell-attached patch-clamp mode was also used to determine mean open/closed times. In these experiments, BK channel activity was recorded 3 times, 20 s each, with 1 min intervals. For all cell-attached experiments, the voltages given represent the potential at the intracellular side of the membrane.

To determine whether BK channels were present in the membrane of cultured striatal neurons we recorded BK recorded single channels in the inside-out and outside-out patch-clamp configuration. We recorded in these configurations to precisely control the free intracellular calcium concentration to which BK channels were exposed. For all experiments, imposed voltages correspond to the potential at the intracellular side of the membrane. To measure the activation rate of channels in striatal culture, macroscopic currents were compiled by summing 100 consecutive single-channel traces obtained by stepping the membrane of an outside-out patch from a holding potential of −60 mV to a potential eliciting an NPo of −0.5, between +30 and +60 mV, in the presence of 10 μM free Ca2+ in the recording pipette. Leak currents were subtracted on-line using the P/n (4) protocol. To yield the macroscopic current, traces were summed and the activation kinetics fit using FitMaster software (HEKA Elektronik). The activation rate (milliseconds) versus voltage was plotted and linearly fit. The slope of the linear fit was used as the measure of voltage dependence.

**Whole-cell recordings.** Using the standard whole-cell patch-clamp recording method (Hamill et al., 1981), the membrane was depolarized to various potentials for 500 ms from a holding potential of −60 mV. Mean BK current amplitude was measured at steady state, 450–490 ms after the beginning of the voltage step. The recording pipette routinely included in mM: 0.1 leupeptin, 12 phosphocreatine, 2 K-ATP, and 0.2 Na-GTP to prevent run-down of the Ca2+ current (Kittler et al., 2005). The inclusion of leupeptin serves to block proteases while phosphocreatine and K-ATP allow the regeneration of ATP. When applying pharmacological agents such as 4-aminopyridine (4-AP), ibetixin (Ibx), tetraethylammonium chloride (TEA-Cl), and 8-bromo-cAMP hemiacid tubes containing the appropriate solution were juxtaposed to the cell.

**In vivo ethanol exposure**

For these experiments, we used P25–P33 male C57BL/6j mice. Hourly EtOH intraperitoneal injections were administered. The first injection (1.8 g/kg, i.p.) was followed by booster injections (1.2 g/kg, i.p.) every hour for two or five subsequent injections. BEC measurements were obtained from mice (naïve, or immediately before a 3 or 6 h EtOH injection protocol). Trunk blood was collected in heparinized capillary tubes, centrifuged (1500 × g for 5 min) and analyzed using an alcohol oxidase-based assay. We measured blood alcohol levels on a GM7 MicroStat Analyzer (Analox Inst Ltd.).
In vivo alcohol exposure and acute dissociation of striatal neurons

Procedures for slice preparation and freshly isolated striatal neurons have been described in detail by Martin et al. (2008). Briefly, mouse brains (P25–P33) were sliced (350 μm) using a Vibratome 3000 (Vibratome) and incubated for up to 6 h at room temp (20–22°C) in a gassed (95% O2 and 5% CO2) saline solution. Following enzymatic digestion with protease XIV (1 mg/ml), the tissue was mechanically triturated using fire-polished Pasteur pipettes, and cells were plated onto a 35 mm Petri dish.

Neuron survival assay

After rinsing the dissociated neurons with HBSS (Sigma-Aldrich), cells were stained with 7.5 μg/ml fluorescein diacetate and 2.3 μg/ml propidium iodide in HBSS. After 20 min at room temperature, the coverslips were mounted on slides and imaged immediately. With blue excitation, live neurons fluoresce green and dead cells fluoresce red (Breuer et al., 1993). The measure of survival was reported: percentage survival = [live cells/(live cells + dead cells)] × 100. In all acutely dissociated neuronal preparations, we observed >84% viable neurons.

Immunocytochemistry

After acute dissociation, neurons plated on glass coverslips were fixed for 30 min with formaldehyde 3.7%, permeabilized for 10 min with 0.1% Triton X-100 and blocked for 30 min with 10% goat serum in Dulbecco’s PBS (DPBS) (Invitrogen) at room temperature. Samples were incubated overnight at 4°C with primary antibody, rabbit anti-glutamic acid decarboxylase 65/67 (GAD) (1:500; Sigma-Aldrich); followed by a 2 h incubation at room temperature with a secondary antibody mixture of Alexa Fluor 594 goat-anti-rabbit (1:300; Invitrogen) and mouse anti-α tubulin-Alexa 488 (1:300; Invitrogen). The nuclei of the cells were stained with 1 μg/ml Hoechst 33342 (Invitrogen) for 5 min at room temperature. Every treatment was followed by 2 PBS washes, 10 min each. The coverslips were mounted onto slides with Vectashield mounting medium (Vector Laboratories) and sealed with nail polish. Pictures were acquired through an LSM 5 Pascal Confocal Microscope (Zeiss) using a 40× oil-immersion objective. Immunostaining results indicate 83% of cells from the acutely dissociated preparation were markedly stained for GAD.

Data analysis

Data were analyzed using Tac X4.1.5 and TacFit X4.1.5 software (Bruxton). NP0 values were calculated from all-points amplitude histograms by fitting the histogram with a sum of Gaussian functions (Bruxton). NPo values were calculated from all-points amplitude histograms by measuring the distance between the modes corresponding to the closed state and the first opening level. The unitary conductance (γ) was taken as the slope of the unitary current amplitude–voltage relationship. In single-channel patches, durations of open and closed times were measured with half-amplitude threshold analysis. A maximum-likelihood minimization routine was used to fit curves to the distribution of open and closed times. Determination of the minimum number of terms for adequate fit was established using a standard F statistic table (significance level, p < 0.01).

To compute Gmax/60 mV, a series of macroscopic currents were obtained in the presence of 1 mM 4-AP to block Iγ current. Current traces were evoked from a holding potential of −60 mV to +180 mV or 360 mV in 20 mV increments with 1 or 10 μM free Ca2+ in the recording pipette, respectively. The conductance (G) is the mean amplitude taken from the sustained portion of the macroscopic current, 450–490 ms after the beginning of the voltage step. Gmax is the mean amplitude taken at the minimum voltage that elicits maximal current. Depolarizing steps greater than the minimum voltage do not elicit a significant amount of additional current.

Experimental solutions

Regular Locke’s solution contained the following (in mM): 2 KCl, 142 NaCl, 2 MgCl2, 2 CaCl2, 13 glucose, and 15 HEPES. High potassium pipette solution contained the following (in mM): 135–140 K-glucmate, 0–4 HEDTA [N-(2-hydroxyethyl)ethylendiaminetriacetic acid], 0–4 EGTA, 15 HEPES, 1 MgCl2, and 0.5–2.2 CaCl2. HEDTA, EGTA, and CaCl2 concentrations were adjusted to obtain the desired concentrations of free calcium, ranging from 1 to 10 μM free Ca2+. Free Ca2+ concentrations were determined by Sluder’s software and confirmed with a Kwik-Tip calcium probe (World Precision Instruments).

Chemicals

Ethanol, HEPES, and MgCl2 were obtained from American Bioanalytical. BaCl2 and CaCl2 were from Fisher Scientific. Potassium gluconate, glucose, HEDTA, EGTA, TEA-Cl, 4-AP, Ibtx, 8-bromo-cAMP, leupeptin, phosphocreatine, KATP, and Na-GTP were obtained from Sigma-Aldrich. NaCl and KCl were from EM Science.

Reverse transcription PCR

Total RNA was isolated from the striatal culture dishes with TRizol (Invitrogen). Manufacturer’s instructions were followed for RNA isolation using an RNeasy Protect Mini extraction kit (Qiagen) and Glycol Blue (Ambion) to aid in the visualization of RNA precipitate. RNA quality (A260/280) and concentration was determined by a ND-1000 Spectrophotometer (NanoDrop). After extraction, total RNA aliquots were treated with 20 U of RNase-Free DNase (Promega) for 15 min at 37°C to minimize the risk of genomic DNA contamination. First strand cDNA was reverse transcribed from the DNase-treated aliquots using the iScript cDNA Synthesis Kit (Bio-Rad) and stored at −20°C.

A 6-carboxy-fluorescein reporter dye-based real-time PCR was used to quantify expression changes of BK β1 and STREX (stress axis-regulated exon) mRNA in treated striatal cultures compared with control cultures and performed in triplicate. β-actin mRNA is unaffected by acute and chronic alcohol exposure in neuronal tissue and was used as an endogenous control to accurately calculate mRNA expression. No-reverse transcriptase and no-template controls were routinely included. The following pairs of primers were used: KCNMB1, ATCAAGGACCAGGAAAGCTG (5’ primer) and CTCATCTGCAGGCAAGAC (3’ primer); and STREX AGGCCGCCCCAAAGATGT (5’ primer) and ATGCGCCGACGTCACTGCTCA (3’ primer). All amplicons (the primers and probes) were checked for lack of secondary structure formation ensuring optimal PCR efficiency (IDT BioTools). Probes were synthesized to have their 5’ end labeled in 25 μl final volume, using Real Time Master Mix Probe (Eppendorf), on an ABI Prism 7500 PCR System (Applied Biosystems). The reverse-transcription reactions were performed at 42°C for 45 min, followed by 35 cycles of PCR amplification (15 s at 94°C, 30 s at 56°C, 15 s at 72°C). To verify that the signals detected were not caused by genomic DNA contamination, Reverse transcription PCRs were also performed for each pair of primers without reverse transcriptase. Significance was determined using one-way ANOVA.

Results

Cultured striatal neurons express functional BK channels

We first determined the basic electrophysiological properties, including voltage sensitivity and conductance, of single BK channels recorded from inside-out patches pulled from cultured striatal neurons (Fig. 1A). Single-channel currents were elicited by depolarizing the membrane from −80 mV to +80 mV in 20 mV increments, while exposing the intracellular surface to 10 μM free Ca2+. Single-channel activity between −60 mV and +40 mV is shown in Figure 1B. At −60 mV, the channel displays a low open probability (NP0 = 0.10), and shows increasing activity (NP0 = 0.96) as the membrane is depolarized to +40 mV. Figure 1C shows a plot of the current amplitude versus the membrane potential of a striatal BK channel. The current–voltage relationship is well fitted by linear regression (r = 0.99) yielding a slope conductance of 230 pS, similar to BK channels recorded in other preparations (McManus, 1991; Vergara et al., 1998, 1999). In addition, current reversed at 0 mV in symmetrical potassium conditions, [K+]i = [K+]o indicating the channels are selective for potassium.

The relative contribution of BK current to the total macroscopic current was determined by pharmacologically dissecting the total K+ current. In Figure 1D, the largest trace is the total macroscopic...
K+ current, recorded in normal Locke’s solution containing 2.2 mM calcium. Addition of the $I_h$ channel inhibitor 4-AP (1 mM) blocks the fast inactivating $I_h$ and indicates that 43 ± 5% of the total current is due to $I_h$. Perfusion with 100 nM Ibtx specifically blocks BK channels, leaving a 4-AP- and Ibtx-resistant current, which indicates that the BK channel contributes 27 ± 7% of the total K+ current. Finally, addition of 100 mM TEA-Cl, a blocker of voltage-dependent potassium channels, leaves a small residual current (smallest trace). In all subsequent figures, $I_h$ is subtracted from the macroscopic current by inclusion of 1 mM 4-AP.

Figure 1. Conductance, voltage dependence, and pharmacology of BK channels in cultured striatal neurons. A. Digitally captured image of rat P8 striatal neurons 1 week in culture. B. Time spent in the open state (inside-out configuration; 10 μM free Ca$^{2+}$) increases as the membrane is depolarized. C and O represent the closed and open states, respectively. C. Plot of BK channel current amplitude as a function of membrane potential. A linear fit of this relationship ($r = 0.99$) gave a BK channel unitary conductance of 230 pS. D. Pharmacological dissection of the macroscopic potassium current in striatal neurons. Macropscopic currents were evoked by stepping from a holding potential of −60 mV to +80 mV. The largest trace is the total current, recorded in normal Locke’s solution containing 2.2 mM calcium. Addition of 1 mM 4-AP to the bath solution leaves a small residual current. The remaining current, $I_h$, is subtracted from the macroscopic current by inclusion of 1 mM 4-AP.

The relationship between duration of initial alcohol exposure and persistence of molecular tolerance is markedly nonlinear. First, we determined the mRNA levels of various BK channel isoforms, such that STREX, the predominant isoform, predominates. We took a threefold approach, to determine whether the 3 h versus 6 h transition to persistent ethanol tolerance can be attributed to changes in the predominant α subunit isoform. First, we determined the mRNA levels of different BK channel isoforms during withdrawal from 3 and 6 h ethanol exposure. Second, we exogenously expressed various splice variants in HEK 293 cells and compared their biophysical properties with those observed for BK in striatal culture. Finally, we used pharmacological tools to discriminate between splice variant populations based on their differential regulation by secondary signaling pathways (Chen et al., 2005). There are no commercially available antibodies to differentiate BK splice variants. Moreover, antibody studies will be problematic because the majority of BK splice sites are located intracellularly in the transscripts, especially STREX, encode alcohol-resistant BK channel proteins (Pietrzynkowski et al., 2008). We hypothesized that the time-dependent changes in BK properties observed in striatal neurons might be explained by a time-dependent shift in α subunit isoforms, such that STREX, the EtOH-insensitive variant, predominates. We took a threefold approach, to determine whether the 3 h versus 6 h transition to persistent ethanol tolerance can be attributed to changes in the predominant α isoform. First, we determined the mRNA levels of different BK channel isoforms during withdrawal from 3 and 6 h ethanol exposure. Second, we exogenously expressed various α splice variants in HEK 293 cells and compared their biophysical properties with those observed for BK in striatal culture. Finally, we used pharmacological tools to discriminate between splice variant populations based on their differential regulation by secondary signaling pathways (Chen et al., 2005). There are no commercially available antibodies to differentiate BK splice variants. Moreover, antibody studies will be problematic because the majority of BK splice sites are located intracellularly in the
C-terminal tail (Xia et al., 2002; Krishnamoorthy et al., 2005). Thus, a description of which splice variants are functionally present in the membrane would be difficult using morphologic techniques. This necessitated this correlative approach, rather than a more direct one, such as determining protein levels using immunoblotting procedures.

**STREX mRNA is upregulated during withdrawal from 6 h but not 3 h EtOH**

The BK ALCOREX, Insertless, and STREX α subunit isoforms are all present in alcohol naïve striatal neurons (Pietrzykowski et al., 2008). ALCOREX and Insertless are relatively sensitive to 50 mM EtOH, while STREX is resistant (Pietrzykowski et al., 2008). Therefore, we postulated that during withdrawal from 6 h alcohol exposure STREX becomes dominant. Specific primers to STREX were designed (see Materials and Methods) to test this. STREX was upregulated after 6 h of exposure to 20 mM EtOH (3.11 ± 0.18-fold compared with control; n = 6, experiments performed in triplicate); there was no similar upregulation after 3 h EtOH exposure (Fig. 3). Furthermore, this upregulation was sustained, returning to baseline only after 24 h (Fig. 3). We also included primers for the auxiliary BK subunit β1, which is detected at low levels in the striatum (Martin et al., 2004) and other brain regions (Jiang et al., 1999), and has been reported to confer alcohol resistance (Feinberg-Zadek and Treistman, 2007). β1 mRNA, which was present at low levels, did not significantly differ from control during withdrawal from either 3 or 6 h EtOH exposure (one-way ANOVA, p = 0.97).

**Neuronal BK characteristics during withdrawal from 3 versus 6 h EtOH exposure**

The calcium sensitivity, activation rate, and mean open and closed times of BK were determined during withdrawal.

**Gating kinetics of BK are altered after withdrawal from 6 h but not 3 h alcohol exposure**

To study BK current kinetics we summed 100 repetitively evoked single-channel sweeps into a cumulative current trace, rather than using whole-cell macroscopic analysis. The cumulative current trace resembles the classical macroscopic current recorded in the whole-cell patch-clamp configuration, but assures the absence of even very small contaminating currents that might influence kinetic analysis using whole-cell macroscopic methodology. The potential of an outside-out patch was stepped, in the presence of 1 mM 4-AP to block K+ channels preexposed to 20 mM EtOH for 3 h and with ethanol withdrawn for 4 h (A), preexposed to 20 mM EtOH for 6 h and with ethanol withdrawn for 4 h (B), or preexposed to 20 mM EtOH for 6 h and with ethanol withdrawn for 24 h (C). All traces were recorded at a holding potential of +60 mV. C and O represent the closed and open state, respectively. D, Plot of aggregate single-channel recording data. Note: In D, a value of 1 on the ordinate represents maximal tolerance, while higher numbers represent recovery from tolerance (i.e., return of potentiation).

**Figure 2.** Persistence of acute tolerance is a function of ethanol exposure time. A–C, BK channel activity (NPo) in cell-attached patches before and after challenge with 50 mM EtOH in neurons that were preexposed to 20 mM EtOH for 3 h and with ethanol withdrawn for 4 h (A), preexposed to 20 mM EtOH for 6 h and with ethanol withdrawn for 4 h (B), or preexposed to 20 mM EtOH for 6 h and with ethanol withdrawn for 24 h (C). All traces were recorded at a holding potential of +60 mV. C and O represent the closed and open state, respectively. D, Plot of aggregate single-channel recording data. Note: In D, a value of 1 on the ordinate represents maximal tolerance, while higher numbers represent recovery from tolerance (i.e., return of potentiation).

**Figure 3.** Six, but not 3 h of 20 mM EtOH induces an upregulation of STREX mRNA during withdrawal. mRNA levels in cultured striatal neurons were measured by real-time PCR with primers to α-STREX and to the β1 subunit. Immediately after 6 h of exposure to 20 mM EtOH, STREX mRNA increased approximately threefold compared with control. Bar graphs represent the mean ± SEM for six independent experiments performed in triplicate. Statistical significance was determined using one-way ANOVA;∗ p < 0.05.

There is a shift in BK channel calcium sensitivity after 6 h, but not a 3 h, exposure to EtOH

G/G\textsubscript{max} was derived from a series of macroscopic currents obtained with either 1 or 10 μM free Ca\textsuperscript{2+} in the recording pipette, in the presence of 1 mM 4-AP to block IK\textsubscript{A}. We found a leftward shift in the G–V relationship (Fig. 5), indicative of increased Ca\textsuperscript{2+}-responsiveness, during withdrawal from a 6 h but not 3 h exposure to 20 mM EtOH (left and middle panels). As would be expected, this shift was not immediately apparent, presumably
Kinetic properties of BK channels differ from control during withdrawal from a 6 h, but not a 3 h exposure to 20 mM EtOH. A–C, A series of six consecutive BK traces evoked by depolarizing channels that were naive (A), preexposed to 20 mM EtOH for 3 h and with ethanol withdrawn for 24 h (B), or preexposed to 20 mM EtOH for 6 h and with ethanol withdrawn for 24 h (C). Channel activity was recorded in an outside-out patch with 10 μM free Ca²⁺ in the recording electrode. Cumulative current was compiled from 100 single-channel traces. Current activation was best fit with a single exponential. The use of different voltage steps used in A and B versus C reflects the need to maintain constant Pₒ values in light of the shift in the Pₒ–V curve after transition to PMT. Compiled currents shown in D–F were obtained at the same potentials as the traces shown in A–C.

Figure 5. There is a leftward shift in G–V relationship present after withdrawal from a 6 h, but not 3 h, exposure to 20 mM EtOH. Plots of G–V relationship for macroscopic currents obtained with 1 μM free Ca²⁺ (left graph), or 10 μM free Ca²⁺ (middle graph). Graph on right was obtained with 10 μM free Ca²⁺ in the recording pipette and 100 mM Ibtx in the bath. BK currents were evoked from a holding potential of −70 mV with +10 mV (left and middle graph) or +20 mV (right graph) incremental voltage steps. Macroscopic currents were measured at steady-state condition, 450–490 ms after the beginning of the voltage step. Bath contained 1 mM 4-AP to block I Calcium current.

Figure 4. Kinetic properties of BK channels differ from control during withdrawal from a 6 h, but not a 3 h exposure to 20 mM EtOH. A–C, A series of six consecutive BK traces evoked by depolarizing channels that were naive (A), preexposed to 20 mM EtOH for 3 h and with ethanol withdrawn for 24 h (B), or preexposed to 20 mM EtOH for 6 h and with ethanol withdrawn for 24 h (C). Channel activity was recorded in an outside-out patch with 10 μM free Ca²⁺ in the recording electrode. Cumulative current was compiled from 100 single-channel traces. Current activation was best fit with a single exponential. The use of different voltage steps used in A and B versus C reflects the need to maintain constant Pₒ values in light of the shift in the Pₒ–V curve after transition to PMT. Compiled currents shown in D–F were obtained at the same potentials as the traces shown in A–C.

Single-channel kinetics
Single-channel kinetics
Single-channel kinetics

BK characteristics in transfected HEK-293 cells
HEK-293 cells were transfected with EtOH-sensitive BK α subunit variants (Insertless and ALCOREX), and an EtOH-insensitive variant, STREX, and the same biophysical parameters were measured as had been measured in striatal neurons. Macroscopic current traces were obtained with 1 μM free Ca²⁺ in the recording pipette and 100 mM Ibtx in the bath. BK currents were evoked from a holding potential of −70 mV with +10 mV (left and middle graph) or +20 mV (right graph) incremental voltage steps. Macroscopic currents were measured at steady-state condition, 450–490 ms after the beginning of the voltage step. Bath contained 1 mM 4-AP to block I calcium current.
activity was evoked by stepping from a holding potential of -60 mV to a potential of +70 mV for the traces recorded in A and B or to +50 mV for the traces recorded in C. The dwell time distributions of single-channel patches are shown in the histograms below the current traces. Open and closed times were computed with half-amplitude threshold analysis. Curves were fitted using a maximum-likelihood minimization routine. The open times distribution could be well fitted with a single-component function while the closed time distribution was well fitted with a double exponential.

The hypothesis that the STREX isoform predominates after 6 h of exposure.

Striatal currents are inhibited by PKA activation during withdrawal from 6 h but not 3 h EtOH

Alternatively spliced α isoforms are differentially regulated by secondary signaling pathways, and we used this as a tool to further test the hypothesis that STREX isoforms predominate in striatal culture after 6, but not 3 h EtOH exposure, by assessing the actions of the PKA activator, 8-bromo-cAMP. Chen et al. (2005) have shown that STREX is inhibited by cAMP while other isoforms such as Insertless are potentiated (Chen et al., 2005). This differential response is attributable to an additional PKA site in the 58 aa STREX insert in the C-terminal tail (Chen et al., 2005). Figure 8 shows macroscopic currents evoked by steps to +80 mV from a holding potential of −60 mV (1 μm free Ca^{2+} in the pipette) after various ethanol exposures and withdrawal periods. Currents from naive neurons or neurons exposed to ethanol for 3 h and withdrawn for 24 h were potentiated by extracellular perfusion with 250 μM 8-bromo-cAMP (41.5 ± 8.43 and 35.94 ± 10.7%, respectively). In sharp contrast, currents from neurons exposed to ethanol for 6 h and withdrawn for 24 h were markedly inhibited (32.9 ± 15.1%) by 8-bromo-cAMP (Fig. 8F) consistent with studies showing that STREX channel activity is reduced by this agonist. Interestingly, immediately following either 3 or 6 h EtOH exposure, 8-bromo-cAMP did not have a significant effect (−4.0 ± 3.6 and −8.7 ± 6.2% respectively). This may reflect a change in the phosphorylation level of BK channels induced by alcohol that persists when the drug remains present.

The 3–6 h transition can be observed in vivo

We next asked whether the transition in persistence of tolerance observed between 3 and 6 h exposure in cultured striatal neurons in vitro could be observed after in vivo exposure in mice. One of the concerns with cultures of striatal neurons is that they are electrically silent because the cultures consist principally of GABAergic inhibitory neurons lacking excitatory inputs. Thus, it is important to show that the temporal shift in molecular tolerance is not simply a function of the cultures. For our in vivo studies, we used C57BL/6J mice. The rationale for use of mice rather than rats for this part of the study was predicated upon two considerations: (1) previous in vivo studies examining the role of BK in alcohol tolerance were performed with KO mice (Martin et al., 2008), providing the potential to use KO technology in future studies of BK and the transition to PMT, and (2) the results provide insight into the generality of the phenomenon between different rodent models.

We first determined a protocol of intraperitoneal injection of EtOH that mimicked the temporal and concentration characteristics used in the in vitro exposure protocol. In injected animals, ethanol is absorbed directly from the peritoneal cavity into the portal bloodstream, where it travels to the liver and is metabolized by hepatic alcohol dehydrogenase. Differences in metabolism of EtOH across gender, body weight, species and other factors determine the concentration of ethanol that actually reaches the brain. Crippens et al. (1999), used microdialysis to measure alcohol concentrations in the brain versus concentrations in the tail blood of adult rodents after intraperitoneal administration of ethanol, and consistently found that ethanol concentrations in the brain were significantly lower than in blood samples. Therefore, we selected an EtOH concentration that might better approximate the concentration range used for the in vitro experiments.
molecular tolerance that occurs between 3 and 6 h. Moreover, although our data are consistent with the upregulation of STREX as the causative agent of the transition to more persistent tolerance, we cannot rule out other routes, such as a time-dependent phosphorylation. Behavioral studies have shown that the development of tolerance after short alcohol exposures is dependent upon exposure protocol. For example, in mice and rats, a single intraperitoneal administration of EtOH reduces motor impairment and hypothermic response to a second dose administered 8–24 h after the first dose (Crabbe et al., 1979; Khanna et al., 1996). These studies also demonstrated that higher initial doses of alcohol produced the greatest degree of tolerance. Furthermore, the fact that the degree of tolerance was approximately equivalent after a single intraperitoneal dose of 4 g/kg versus two successive doses of 2 g/kg suggested that peak blood alcohol concentration was not as important as duration of alcohol exposure (Khanna et al., 1996). Numerous studies suggest the existence of alcohol triggered molecular “switches” that may contribute to the formation of behavioral tolerance. For example, changes in biochemical pathways can outlast the presence of the initiating drug of abuse and thereby qualify as drug-activated switches (Pandey et al., 2001, 2003; Borlikova et al., 2006). Examples of potential drug-activated molecular switches include transcription factors such as δfosB and CREB (Kelz et al., 1999; Nestler et al., 2000), and secondary signaling molecules such as PKA (Coe et al., 1996; Pandey et al., 2001; Lin et al., 2006). In flies, CREB, which is activated by PKA, mediates upregulation of dslo during rapid tolerance to benzyl alcohol (Wang et al., 2007).

**Six hour time point**

Six hours of drug exposure is emerging as an important temporal threshold for a number of phenomena related to drug tolerance and addiction. For example, escalation of cocaine self-administration occurs after 6 h, but not 1 h of drug access (Ahmed and Koob, 1998). Ben-Shahar et al. (2006) found that 6 h, but not 1 h, of daily access to self-administered cocaine results in elevated levels of dopamine transporter (Ben-Shahar et al., 2006). Ethanol promotes transcription of the catalytic subunit of PKA from the Golgi area to the nucleus after 6 h, but not 1 h of drug exposure (Dohrman et al., 2002). To our knowledge, there have not yet been any studies showing that a single exposure...
to alcohol results in escalated drinking. Indeed, in light of our results, it would be interesting to examine whether the duration of a single drinking bout influences subsequent alcohol drinking pattern, as with cocaine usage.

Our data may implicate a recently described mechanism involving miR-9, a representative of the class of small modulatory RNAs known as microRNAs (Pietrzykowski et al., 2008). Within 15 min of exposure, alcohol increases miR-9 levels in hypothalamic and striatal neurons, resulting in the selective and rapid degradation of those BK transcripts containing a sequence with complementarity to miR-9 (Pietrzykowski et al., 2008). Those transcripts that were degraded encoded channel isoforms with the highest sensitivity to alcohol. For example, the transcript levels of ALCOREX, which form a channel highly sensitive to ethanol, were greatly degraded, whereas STREX transcripts, which form a channel insensitive to ethanol, were relatively enriched. Thus, although not tested here, our data are consistent with a mechanism whereby the emergent predominance of STREX mRNA represents a time-dependent transition involving miR-9. However, this scenario contains an issue requiring further exploration. Upregulation of miR-9 occurs rapidly after exposure to ethanol, evident within 15 min (Pietrzykowski et al., 2008), and well developed before a putative 6 h transition time point. One potential explanation consistent with our data would require that the duration of ethanol exposure influence either the duration or consequences of miRNA upregulation. These parameters are currently unknown. Additionally, the increase in STREX returns to baseline after 24 h, yet at that time, the increase in response to ethanol is still suppressed. Clearly, a fuller understanding of the mechanistic underpinnings of the time-dependent transition requires a greater understanding than we currently have of the temporal dynamics between mRNA changes, BK synthesis, and channel trafficking and insertion.

Here, we describe a mechanism underlying molecular tolerance to alcohol involving a switch in the α subunit isoform composition of BK channels. Indeed, our approach was designed to test this possibility. Moreover, there may be additional benefits inherent to this emerging picture. For example, the substitution of STREX for preexisting BK isoforms not only reduces the sensitivity to alcohol, but increases the sensitivity to Ca²⁺, important in light of the concurrent inhibition of Ca²⁺ channels produced by the drug (Wang et al., 1991). However, other mechanisms also play a role in BK acute tolerance. These include short-duration post-transcriptional changes such as phosphorylation/dephosphorylation, as well as more lasting alterations such as membrane lipid modification (Yuan et al., 2008; Treistman and Martin, 2009), and epigenetic phenomena such as chromatin remodeling (Wang et al., 2007, 2009). Why do multiple and seemingly redundant BK adaptive mechanisms, such as both desensitization and subsequent internalization exist? This is likely necessitated by the particularly large conductance of the channel, such that persistent potentiation would lead to serious nervous system dysfunction.

In a previous study (Pietrzykowski et al., 2008) we reported that ethanol induced a rapid reduction in BK message, including STREX, in hypothalamic neurons. While the absolute amount of mRNA for all BK α isoforms in hypothalamic neurons was reduced, the relative representation of STREX was enriched. Here, we show that over the course of hours, the absolute amount of STREX message in cultured striatal neurons is increased, in contrast to the results obtained in hypothalamic neurons. The most interesting explanation for this difference would be attributable to regional brain differences. Indeed, the actions of ethanol have been found to differ not only between neurons, but even between different subcellular compartments within a single neuron (Martin et al., 2004).
The time-dependent transition to PMT described here may have important implications for the relationship between an individual's drinking pattern and development of tolerance, dependency, and addiction. While the relationship between events at the level of molecules and the level of complex behavior is difficult to pin down, recent studies have attempted to bridge this gap by showing that BK subunit composition in genetically altered mice influences the response to alcohol at the levels of the single-channel protein, the action potential, and the behavioral response to the drug, including propensity to drink alcohol (Martin et al., 2008). Finally, we may question how such sophisticated responses to ethanol have arisen, and whether the evolutionary pressure was in response to ethanol in the environment, or whether ethanol has “hijacked” a mechanism designed to provide neural plasticity in response to other internal or external pressures.

References


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