Ethanol Inhibition of M-Current and Ethanol-Induced Direct Excitation of Ventral Tegmental Area Dopamine Neurons

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INTRODUCTION

The ventral tegmental area dopamine (DA VTA) neurons are cells of origin of the mesolimbic dopamine pathway and provide their projections to the nucleus accumbens (NAcb) (Oades and Halliday 1987). Earlier behavioral and pharmacological studies showed that the mesolimbic dopamine pathway is critical for the reinforcing effects of ethanol (Appel et al. 2004; DiChiara and Imperato 1988; Weiss et al. 1993). Pharmacological study with dopamine agonists and antagonists also supports the involvement of dopamine in ethanol reinforcement (Samson et al. 1990). It was previously reported that rats self-administer ethanol directly into the VTA (Roddis-Henricks et al. 2000) and ethanol excites dopamine cells in the VTA (Gessa et al. 1985). Thus ethanol-induced excitation of DA VTA neurons would increase dopamine release in the NAcb and mediate the reinforcing effects of ethanol.

There are many studies on the ethanol modulation of ligand-gated ion channels in central neurons. In hippocampal neurons, N-methyl-d-aspartic acid (NMDA)-receptor–mediated current is inhibited by ethanol (Hendricson et al. 2004; Lovinger et al. 1989; Peoples and Weight 1995). γ-Aminobutyric acid type A (GABA_A)-receptor–mediated response is potentiated by ethanol in hippocampal neurons (Aguayo and Pancetti 1994; Procotor et al. 1992). Glycine-receptor–mediated response is potentiated by ethanol in midbrain dopamine neurons (Ye et al. 2001) and hypoglossal motoneurons (Eggers and Berger 2004). Adenosine (P2X) receptor–mediated current is inhibited by ethanol in hippocampal neurons (Li et al. 2000). Although ethanol modulates ligand-gated ion channels and affects the excitability of central neurons, the effect of ethanol on modulation of voltage-dependent ion channels in the CNS has not been well studied. DA VTA neurons have intrinsic pacemaker activity with the average firing frequency of 0.5–5 Hz and several voltage-dependent ionic currents contribute to the excitability of these neurons (Appel et al. 2003, 2006; Brodie and Appel 1998; Brodie et al. 1990, 1999; Koyama and Appel 2006a,b; Koyama et al. 2005; Mueller and Brodie 1989; Neuhoff et al. 2002; Okamoto et al. 2006). Therefore the effects of ethanol on voltage-dependent ion channels and the subsequent change in the excitability of these neurons are important cellular mechanisms involved in the reinforcing effects of ethanol.

We previously reported that ethanol increases the spontaneous firing frequency of acutely dissociated DA VTA neurons, in which the influence of synaptic activity on the excitability of these neurons was eliminated (Brodie et al. 1999). Our current-clamp studies suggest that ethanol excites DA VTA neurons by reducing a K^+ current, which contributes to the action potential (AP) afterhyperpolarization (AHP) (Appel et al. 2003; Brodie and Appel 1998). In addition, our voltage-clamp studies showed that ethanol reduces the sustained K^+ currents of DA VTA neurons (Appel et al. 2003; Brodie et al. 2000). Thus it is likely that ethanol excites DA VTA neurons directly by the reduction of K^+–channel conductance.

M-current (I_M) is a sustained K^+ current and activated at the subthreshold range of membrane potential, regulating AP generation (Aiken et al. 1995; Brown and Adams 1980). Five types of KCNQ (KCNQ1–5) channels were previously reported to mediate I_M (Lerche et al. 2000; Selyanko et al. 2000; Shapiro et al. 2000; Sogaard et al. 2001; Wang et al. 1998). Immunohistochemical co-localization of KCNQ2 and KCNQ4 channel subunit proteins was previously found in VTA neurons (Cooper et al. 2001; Kharkovets et al. 2000). We recently showed that I_M regulates action potential generation to reduce...
the excitability by underlying fast and slow AHP components without changing the middle component of AHP (Koyama and Appel 2006a). Consistent with this finding, it was earlier shown that the KCNQ4 channel subunit is the major KCNQ subunit in DA VTA neurons and regulates the excitability of these neurons (Hansen et al. 2006). Interestingly, Moore et al. (1990) showed that ethanol reduces \( I_M \) in CA1 hippocampal pyramidal neurons.

In our present study, we examined whether ethanol inhibition of \( I_M \) contributed to the excitation of DA VTA neurons. For this purpose, we used acutely dissociated VTA neurons with enzyme treatment, in which both excitatory and inhibitory synaptic influences on these neurons were eliminated (Koyama et al. 2005) and nystatin-perforated patch recording to minimize intracellular dialysis and prevent the rundown of \( I_M \) (Koyama and Appel 2006a). Part of this work previously appeared in abstract form (Koyama and Appel 2004).

**METHODS**

**Preparation of dissociated neurons**

Animals used in this study were treated in strict accordance with the U.S. National Institutes for Health Guide for the Care and Use of Laboratory Animals and all experimental methods were approved by the Animal Care Committee of the University of Illinois at Chicago. Each Fisher 344 rat (14–18 days old) was decapitated and the brain quickly removed. The brain was placed in the ice-cold cutting solution consisting of (in mM): 220 sucrose, 2.5 KCl, 2.4 CaCl\(_2\), 1.3 MgSO\(_4\), 1.24 NaH\(_2\)PO\(_4\), 26 NaHCO\(_3\), and 11 D-glucose, which was constantly bubbled with 95% O\(_2\)-5% CO\(_2\). Transverse brain slices, at a thickness of 400 \( \mu \)m, were made using a Vibratome. The brain slices were incubated for 30 min in artificial cerebrospinal fluid (ACSF), consisting of (in mM): 126 NaCl, 2.5 KCl, 2.4 CaCl\(_2\), 1.3 MgSO\(_4\), 1.24 NaH\(_2\)PO\(_4\), 26 NaHCO\(_3\), and 11 D-glucose (osmolality 300 mosM), which was constantly bubbled with 95% O\(_2\)-5% CO\(_2\) at room temperature (23–25°C). The brain slices were then incubated in a HEPES-buffered solution (in mM: 145 NaCl, 2.5 KCl, 2.4 CaCl\(_2\), 1.3 MgCl\(_2\), 10 HEPES, and 11 D-glucose (pH adjusted to 7.4 with NaOH; osmolality, 300 mosM) containing papain (15–18 U/ml) at 32°C for 20–25 min. After papain treatment, the brain slices were further incubated in the ACSF for 20–40 min. The VTA neurons were dissociated using a vibrating stylus apparatus dispersing cells from the brain slices as previously described (Koyama et al. 2005). Once the cell dissociation procedure was completed (4–7 min), the brain slice was removed from the culture dish and the dissociated neurons settled and adhered to the bottom of the dish within 20 min. The dissociation procedure with papain described above was earlier shown to yield acutely dissociated VTA neurons (Koyama et al. 2005) and no spontaneous synaptic potentials or synaptic currents were recorded from these neurons in the present study. DA VTA neurons were identified in current-clamp recording based on electrophysiological characteristics previously described (Koyama and Appel 2006a).

**Nystatin-perforated patch recording in dissociated neurons**

Electrophysiological measurement was performed using an Axopatch-1B patch-clamp amplifier (Axon Instruments, Union City, CA). The tip resistances of the electrodes ranged from 1 to 2.5 M\( \Omega \) when filled with pipette solution (in mM: 60 K-acetate, 60 KCl, 1 CaCl\(_2\), 2 MgCl\(_2\), and 40 HEPES; pH adjusted to 7.2 with KOH; final [K\(^+\)] = 131 mM; osmolality, 290 mosm). To minimize the rundown of \( I_M \), nystatin-perforated patch recording was used as previously described (Koyama and Appel 2006a). Nystatin was dissolved in methanol at a concentration of 10 mg/ml. This nystatin stock solution was diluted with pipette solution to a final concentration of 100–200 \( \mu \)g/ml and the electrodes were backfilled with this solution. After cell-attached configuration had been completed, access resistance was periodically monitored and capacitative transients were cancelled. When the access resistance had reached a steady level (8–14 M\( \Omega \)), the recording was started. Completion of perforated patch was distinguished from membrane rupture because the hallmark of membrane rupture was accompanied by the sudden appearance of steep capacitative transients with the reduction of access resistance to 3–7 M\( \Omega \); this can be compared with the access resistance in conventional whole cell configuration, described previously (Koyama and Appel 2006b). Pore formation with the nystatin-perforated patch technique is not accompanied by steep capacitance transients. Current- and voltage-clamp recordings were carried out with the HEPES-buffered solution constantly bubbled with 100% O\(_2\). The liquid junction potential between the pipette solution and the HEPES-buffered solution was estimated to be 5 mV (Neher 1992) and the results were corrected. Membrane currents and voltages were filtered at 1 kHz and acquired at a sampling frequency of 10 kHz. Data acquisition was performed by a DigiData 1322A interface and pClamp software version 9.0 (Axon Instruments). The dissociated VTA neurons were visualized under phase-contrast on an inverted microscope (Diaphot 300, Nikon, Tokyo, Japan). All experiments were performed at room temperature (23–25°C).

**Drug application for dissociated neurons**

Neurons were continuously bathed in control HEPES-buffered solution and drugs were dissolved at a final concentration in the same solution. Drug solutions were applied by multiple channel manifold (Becton Dickinson). Each channel of the manifold was connected to a gravity-fed reservoir with tubing (ID 860 \( \mu \)m). The output of the manifold was connected to an outflow tube (ID 580 \( \mu \)m), the tip of which was placed within 200 \( \mu \)m of the soma of the recorded neuron. Solution flowed continuously through one manifold channel. Application of drug solutions was controlled by opening or closing valves connected to the reservoirs. The behaviorally active range of blood ethanol concentrations in the rat extends from 40 mM (sedation) to 90 mM (loss of righting reflex) (Majchrowicz and Hunt 1976); the lethal blood ethanol concentration in rats is about 400 mM (loss of righting reflex) (Majchrowicz and Hunt 1976); the lethal blood ethanol concentration in rats is about 400 mM (loss of righting reflex) (Majchrowicz and Hunt 1976). Rats self-administer 44–55 mM ethanol directly into the VTA, indicating that this concentration of ethanol is reinforcing in the whole animal (Rodd-Henricks et al. 2000). In the present study, we examined ethanol concentrations in the range of 20 to 120 mM, which are pharmacologically relevant and sublethal concentrations in the rat.

**Preparation of brain slices**

After rapid removal of the brain, the tissue block containing the VTA was mounted in the Vibratome and submerged in the ice-cold cutting solution. Coronal sections (400 \( \mu \)m thick) were cut and the slice was placed on a mesh platform in the recording chamber. The slice was totally submerged in the ACSF maintained at a flow rate of 2 ml/min; the temperature in the recording chamber was kept at 35°C. The ACSF was saturated with 95% O\(_2\)-5% CO\(_2\) (pH = 7.4). Equilibration time of \( \pm 1 \) h was allowed after placement of the brain slice in the recording chamber before electrodes were placed in the tissue. Recording electrodes were placed in the VTA under visual control. Only those neurons that were anatomically located within the VTA and that conformed to the electrophysiological criteria for dopaminergic neurons (Mueller and Brodie 1989) were studied. These criteria include broad action potentials and regular spontaneous firing frequency at 0.5–5 Hz.
**Extracellular recording in brain slices**

Extracellular recording electrodes were fabricated from 1.5-mm-diameter glass tubing and were filled with 0.9% NaCl. Tip resistance of the microelectrodes ranged from 3 to 8 MΩ. The amplifier (Fintronics, Orange, CT) used in these recordings included a window discriminator, the output of which was fed to both a rectilinear pen recorder and a computer-based data acquisition system that was used for on-line and off-line analysis of the data. The multiplexed output of the amplifier was displayed on an analog storage oscilloscope, for accurate adjustment of the window levels used to monitor single units. An IBM PC-based data acquisition system was used to calculate, display, and store the frequency of firing over 5-s and 1-min intervals.

**Drug administration for brain slices**

Drugs were added to the ACSF by means of a calibrated infusion pump from stock solutions 100- to 1,000-fold the desired final concentrations. The addition of drug solutions to the ACSF was performed in such a way as to permit the drug solution to mix completely with the ACSF before this mixture reached the recording chamber. The use of a calibrated, variable-speed infusion pump permits the accurate addition of several concentrations of drugs from the same stock solution. Final concentrations were calculated from the ACSF flow rate, pump infusion speed, and the concentration of drug stock solution. The small volume chamber (about 300 μl) used in this study permitted the rapid application and washout of drug solutions. Drugs typically reached equilibrium in the tissue after 2 to 3 min of application. A stock solution of 95% ethanol was used in the pump and infusion of ethanol never exceeded 1% of the flow rate of the ACSF.

**Drugs and chemical agents**

Dimethyl sulfoxide (DSMO), ethanol, HEPES, and nystatin were purchased from Sigma–Aldrich (St. Louis, MO). Papain was purchased from Worthington (Lakewood, NJ). 10,10-Bis(4-pyridinylmethyl)-9(10H)-anthracenone (XE991) dihydrochloride was purchased from Tocris-Cookson (Ellisville, MO).

**Data analysis**

Action potentials were analyzed off-line by pClamp software (version 9.0, Axon Instruments). Interspike interval (ISI) histograms were created as previously described (Cocatre-Zilgien and Delcomyn 1992). The number of bins was equal to the square root of the number of ISIs. Bin width was obtained by dividing the ISI range (maximum ISI minus minimum ISI) by the number of bins. To assess the changes of spontaneous firing with drugs, drug effect was quantitated as the mean change in firing rate (normalized as the percentage of control) for a 60-s-long interval during the peak of the drug response as previously described (Brodie et al. 1990). The formula for this normalization is

\[
\text{Firing rate increase (\%)} = \left( \frac{\text{Firing rate with drug}}{\text{Firing rate in control}} \right) \times 100
\]

Data from cells with action potential amplitudes <50 mV were discarded. All average values are expressed as means ± SE. Statistical comparison to assess significant differences was done by one-way ANOVA as appropriate. When needed, the Student–Newman–Keuls post hoc test was used to test multiple comparisons. Correlation was evaluated by linear regression with P < 0.05 being considered significant.

**RESULTS**

**Ethanol inhibition of M-current (I_M) is concentration dependent**

As illustrated in Fig. 1, I_M of DA VTA neurons was recorded in nystatin-perforated patch voltage-clamp configuration, with which I_M is stably recorded without rundown. I_M was measured in the standard deactivation protocol (Brown and Adams 1980) with 1-s-long hyperpolarizing voltage step from a holding potential (V_H) of −25 to −40 mV. Ethanol at concentrations of 40, 80, and 120 mM reduced I_M amplitude as well as the negative shift of the baseline outward current in DA VTA neurons (Fig. 1, A1–A3). Figure 1B shows the measurement of the inward relaxation current caused by deactivation of I_M.
during the voltage step. $I_M$ amplitude was 32 pA in control and ethanol (120 mM) decreased $I_M$ amplitude to 20 pA.

Figure 2A shows the time course of ethanol effect on normalized $I_M$ amplitude in DA VTA neurons. Ethanol inhibition of $I_M$ was reversible and concentration dependent. Figure 2B shows the concentration–response relationship for ethanol inhibition of $I_M$ amplitude. Average maximal ethanol inhibition of $I_M$ amplitude obtained with each concentration was $6.0 \pm 3.5\%$ with 20 mM ethanol ($n = 8$), $7.2 \pm 2.9\%$ with 40 mM ethanol ($n = 8$), $15.3 \pm 2.6\%$ with 80 mM ethanol ($n = 10$), and $29.2 \pm 2.7\%$ with 120 mM ethanol ($n = 9$). From the graph in Fig. 2B, 20% inhibition of $I_M$ by ethanol was estimated to be 90 mM. Although the concentration of nystatin in the stock solution was relatively low and the concentration of methanol in the pipette solution to dissolve nystatin was only $6.0\mu M$, 20% inhibition of $I_M$ by methanol was reversible and concentration dependent. From the $7.1\%$ at $25$ to $40$ mV. This hyperpolarizing voltage step was given in every 20 s. All $I_M$ amplitudes were normalized to the average $I_M$ amplitude obtained from the 5 events just before the application of ethanol. Note that ethanol inhibition of $I_M$ was concentration dependent; 40 mM (filled circles), 80 mM (filled triangles), and 120 mM (filled squares).

**Ethanol inhibition of $I_M$ is not voltage dependent**

We then examined whether ethanol inhibition of $I_M$ was voltage dependent. Figure 3A shows $I_M$ induced by a series of hyperpolarizing voltage steps before, during, and after treatment with 120 mM ethanol. Ethanol inhibited $I_M$ measured with all four hyperpolarizing voltage steps in seven DA VTA neurons. Each $I_M$ in Fig. 3A was obtained by averaging the currents from the seven DA VTA neurons. Figure 3B shows the relationship between membrane voltage and normalized $I_M$ amplitude before (open circles), during (filled circles), and after (open triangles) treatment with 120 mM ethanol in the seven DA VTA neurons. The average normalized $I_M$ amplitude at $-45$ mV was $108.1 \pm 2.1\%$ in control, $74.2 \pm 5.7\%$ in the presence of 120 mM ethanol, and $92.3 \pm 9.3\%$ after washout of ethanol. Ethanol significantly reduced normalized $I_M$ amplitude (one-way ANOVA, $F = 6.9$, $P < 0.01$) but there was no significant difference in normalized $I_M$ amplitude between control and washout values (Student–Newman–Keuls post hoc test, $P > 0.05$). Figure 3C shows the relationship between membrane voltage and mean percentage inhibition of $I_M$ by ethanol. The average ethanol inhibition of $I_M$ obtained with each voltage was $27.3 \pm 3.0\%$ at $-35$ mV, $31.4 \pm 4.9\%$ at $-45$ mV, $31.9 \pm 7.1\%$ at $-55$ mV, and $31.3 \pm 4.4\%$ at $-65$ mV. There was no correlation between membrane voltage and $I_M$ inhibition rate by ethanol ($r = -0.76$, $P > 0.2$).

**Ethanol increases the firing frequency of DA VTA neurons**

In nystatin-perforated patch current-clamp recording, we examined the effect of ethanol on the spontaneous firing of DA VTA neurons. Ethanol (40 mM) increased firing frequency and this ethanol effect was reversible (Fig. 4A1). Figure 4A2 shows the time course of the effect of 40 mM ethanol on spontaneous firing. The effect of 40 mM ethanol on firing frequency for 60-s-long continuous recording was analyzed by an interspike interval (ISI) histogram (see METHODS) (Fig. 4A3); the distribution of ISI is shifted slightly leftward. Ethanol (80 mM) increased firing frequency and this ethanol effect was also reversible (Fig. 4B1). Figure 4B2 shows the time course of the effect of 80 mM ethanol on spontaneous firing. The effect of 80 mM ethanol on firing frequency was analyzed by the ISI histogram (Fig. 4B2); the distribution of ISI is shifted leftward. As shown in Fig. 4C, ethanol increased the firing frequency of DA VTA neurons in a concentration-dependent manner. Average ethanol-induced increase in firing frequency at each concentration was 6.5 ± 1.0% by 20 mM ethanol ($n = 5$), 11.4 ± 3.1% by 40 mM ethanol ($n = 5$), and 26.8 ± 4.9% by

![Graph showing relationship between membrane voltage and $I_M$ amplitude before, during, and after treatment with 120 mM ethanol in DA VTA neurons.](http://jn.physiology.org/)

**FIG. 2.** Ethanol effect is concentration dependent. A: time course of ethanol effect on $I_M$ in DA VTA neurons. $I_M$ was measured with 1-s-long hyperpolarizing voltage step from a $V_H$ of $-25$ to $-40$ mV. This hyperpolarizing voltage step was given in every 20 s. All $I_M$ amplitudes were normalized to the average $I_M$ amplitude obtained from the 5 events just before the application of ethanol. Note that ethanol inhibition of $I_M$ was concentration dependent; 40 mM (filled circles), 80 mM (filled triangles), and 120 mM (filled squares). B: concentration–response curve for ethanol inhibition of $I_M$. Average maximal ethanol inhibition was obtained with each concentration.
80 mM ethanol (n = 10). From the graph in Fig. 4C, 20% increase in spontaneous firing frequency by ethanol was estimated to be produced by 60 mM. Among 14 DA VTA neurons, 120 mM ethanol increased firing frequency by 13.9 ± 2.6% in seven neurons (50%) and prevented spontaneous firing accompanied with membrane depolarization (depolarization inhibition) in seven neurons (50%). Therefore we did not include the results by 120 mM ethanol in this study.

Inhibition of $I_M$ increases the firing frequency of DA VTA neurons

We next examined how $I_M$ inhibition contributes to the spontaneous firing of DA VTA neurons. For this purpose, we used a specific KCNQ channel inhibitor, XE991 (Wang et al. 1998). XE991 (1 μM) increased spontaneous firing frequency (Fig. 5A). Figure 5A, shows the time course of the effect of 1 μM XE991 on spontaneous firing. The effect of 1 μM XE991 on firing frequency was analyzed by the ISI histogram (Fig. 5A); the distribution of ISI is shifted leftward. XE991 (10 μM) increased spontaneous firing frequency, an effect that was reversible (Fig. 5B). Figure 5B, shows the time course of the effect of 10 μM XE991 on spontaneous firing. The effect of 10 μM XE991 on firing frequency was analyzed by the ISI histogram (Fig. 5B); the distribution of ISI is dramatically shifted leftward. As shown in Fig. 5C, XE991 increased the firing frequency of DA VTA neurons in a concentration-dependent manner. Average XE991-induced increase in firing frequency at each concentration was 8.7 ± 2.7% by 0.3 μM XE991 (n = 7), 23.6 ± 6.7% by 1 μM XE991 (n = 7), 42.0 ± 8.2% by 3 μM XE991 (n = 6), and 47.4 ± 7.4% by 10 μM XE991 (n = 8). Among 11 DA VTA neurons, 30 μM XE991 increased firing frequency by 62.1 ± 15.8% in seven neurons (64%) and completely prevented action potential generation accompanied with membrane depolarization in four neurons (36%). Therefore we did not include the results using 30 μM XE991 in this study.

Ethanol inhibition of $I_M$ and ethanol-induced increase in firing frequency

The question remained whether ethanol-induced reduction of $I_M$ could fully account for ethanol-induced excitation of DA VTA neurons. Our previous study showed that 0.3, 1, 3, and 10 μM XE991 reduced $I_M$ amplitude by 27.2 ± 3.8, 51.4 ± 2.9, 81.4 ± 1.7, and 86.6 ± 2.5% in DA VTA neurons (Koyama and Appel 2006a). By comparing the effect of XE991 on $I_M$ inhibition in the previous study and the effect of XE991 on firing frequency shown in Fig. 5C, we found a positive correlation between $I_M$ inhibition and increase in the firing frequency of DA VTA neurons produced by XE991 (Fig. 6A, open circles). The slope of the linear relationship was 0.5. In addition, we compared $I_M$ inhibition by ethanol shown in Fig. 2B and the increase in firing frequency produced by ethanol shown in Fig. 4C. In the same manner, we obtained a positive correlation between $I_M$ inhibition by ethanol and ethanol-induced increase in the firing frequency of DA VTA neurons (Fig. 6A, filled circles). The slope of this linear relationship for ethanol effects was 1.3.

Ethanol effect on spontaneous firing under the block of $I_M$

Extracellular single-unit recording was used to measure the effect of ethanol (40, 80, and 120 mM) on the firing frequency of DA VTA neurons in brain slices. After washout of the last concentration of ethanol, 10 μM XE991, a concentration that blocked $I_M$ by 87% (Koyama and Appel 2006a), was applied and the same concentrations of ethanol were tested again. The percentage increase in firing frequency produced by each ethanol concentration was calculated with the formula shown.
in METHODS. Figure 7 shows the concentration–response relationship between ethanol and the percentage increase in firing frequency before and after treatment with 10 μM XE991 from 15 DA VTA neurons. The ethanol excitation was concentration dependent ($P < 0.001$). Before XE991 treatment, average ethanol-induced increase in firing frequency at each concentration was 10.4 ± 1.6% by 40 mM ethanol, 17.7 ± 1.9% by 80 mM ethanol, and 27.9 ± 4.8% by 120 mM ethanol (Fig. 7, filled circles). Although we anticipated the partial reduction of ethanol-induced excitation in the presence of 10 μM XE991, there was no statistically significant effect of XE991 on the ethanol concentration–response curve ($P > 0.05$). For the 15 DA VTA neurons, 10 μM XE991 produced no significant change of firing frequency (1.3 ± 1.5%).
neurons. However, ethanol reduced  
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shows that 120 mM ethanol reduces  
I  
M  
that 44 mM ethanol reduces  
I  
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study than in the present report. Moore et al. (1990) reported (Hansen et al. 2006). We also found that ethanol inhibition of  
VTA neurons and regulates the excitability of these neurons  
KCNQ4 channel subunit is the major KCNQ subunit in DA  
neurons (at room temperature). In our previous study per-  
formed at 34°C, depolarization blockade was not observed in dissociated DA VTA neurons (Brodie et al. 1999). We reported that higher concentrations of ethanol (>300 mM) produce initial increases in firing frequency followed by the complete block of firing in DA VTA neurons (depolarization inhibition) (Appel et al. 2006). There is a possibility that the ethanol-induced depolarization inhibition of spontaneous firing in DA VTA neurons may depend on the temperature of bathing solutions. However, the cellular mechanism of the depolarization inhibition by ethanol in these studies is unclear and additional studies will be needed to elucidate how ethanol produces depolarization inhibition.

Treatment with XE991 increased firing frequency in a concentration-dependent manner in DA VTA neurons. The increase in the spontaneous firing frequency of DA VTA neurons produced by XE991 correlated positively with  
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inhibition. This result suggests that  
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plays a role in maintaining the spontaneous firing frequency of DA VTA neurons. From the linear relationship obtained with XE991, the contribution of  
I  
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inhibition by 80 mM ethanol is estimated to cause a 6% increase in the spontaneous firing frequency, given that 80 mM ethanol inhibited  
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by 15%. The ethanol-induced increase in the spontaneous firing frequency of DA VTA neurons also correlated positively with  
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inhibition by ethanol. However, we found a difference in the slope values for the linear relationship of  
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inhibition versus increase in spontaneous firing frequency between ethanol (1.3) and XE991 (0.5). In addition, the 20% increase in spontaneous firing frequency by ethanol was obtained at 60 mM, which was smaller than the ethanol concentration that produced 20% inhibition of  
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amplitude (90 mM). Furthermore, in the pharmacological oc- clusion experiment using the submaximal concentration of XE991, ethanol still increased the spontaneous firing frequency of DA VTA neurons in a concentration-dependent manner. These observations suggest that if ethanol inhibition of  
I  
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underlies excitation of DA VTA neurons, other additional ethanol effects on these neurons increase the magnitude of the effect of  
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reduction on firing rate. Although we anticipated the partial reduction of ethanol-induced excitation in the pres- ence of XE991, there was no statistically significant effect of XE991 on the ethanol concentration–response curve. Because 10 mM XE991 inhibits  
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amplitude by 87% (Koyama and

FIG. 6. Ethanol inhibition of  
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and ethanol-induced increase in firing frequency. Linear relationship between  
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inhibition rate and increase in the firing frequency of DA VTA neurons; XE991 (open circles) and ethanol (filled circles).

FIG. 7. Effect of ethanol on the firing frequency of DA VTA neurons in brain slices under the block of  
I  
M. Pooled concentration–response curves for ethanol (40, 80, and 120 mM) effects on spontaneous firing rate measured in DA VTA neurons with extracellular single-unit recording in the absence and presence of 10 mM XE991 (n = 15).
Appel 2006a) and this XE991 effect could increase the membrane input resistance, additional ethanol-induced excitatory effects on DA VTA neurons may be amplified through the increase in the membrane input resistance and compensating for the reduction of the contribution of ethanol-induced \( I_M \) inhibition to spontaneous firing frequency. It is noted that dissociated DA VTA neurons were more excited by ethanol and the block of \( I_M \) than DA VTA neurons in brain slices. For example, 80 mM ethanol-induced increase in spontaneous firing frequency was 27% in dissociated neurons and 21% in neurons from brain slices. The XE991 (10 \( \mu \)M)–induced increase in spontaneous firing frequency was 47% in dissociated neurons and 1.3% in neurons from brain slices. Because dissociated DA VTA neurons were truncated from distal dendrites, the elimination of \( K^+ \) conductance in distal dendrites presumably causes the dissociated neurons to be more excitable than the neurons in brain slices.

It is possible that ethanol may inhibit the other types of sustained \( K^+ \) currents rather than \( I_M \) in DA VTA neurons. We previously showed that 10 mM tetraethylammonium (TEA) blocks \( I_M \) amplitude by 98% in DA VTA neurons (Koyama and Appel 2006a) and ethanol can still increase the spontaneous firing frequency of these neurons in the presence of 10 mM TEA (Appel et al. 2003). We also showed that ethanol-induced excitation of DA VTA neurons is blocked by a low concentration of quinidine (20–40 \( \mu \)M) (Appel et al. 2003). Thus ethanol may inhibit a quinidine-sensitive and relatively TEA-insensitive sustained \( K^+ \) current in addition to its inhibition of \( I_M \) in DA VTA neurons. The candidates for the sustained \( K^+ \) current are three different \( K^+ \) channel families: Kv1 channels (Grismer et al. 1994; Yeola et al. 1996), ether-a-go-go (eag) channels (Ludwig et al. 1994; Schonherr et al. 2002), and TASK channels (Leonoudakis et al. 1998; Meadows and Randall 2001). It is also possible that other ion channels such as \( I_h \) (Okamoto et al. 2006) could also play a role in ethanol excitation (but see Appel et al. 2003). The data of the present study taken together indicate that, although ethanol causes inhibition of \( I_M \) and this results in some increase in firing frequency, another effect of ethanol is primarily responsible for the ethanol-induced increase in firing rate. Physiological reduction of this other predominantly ethanol-sensitive current (by neurotransmitter-mediated second-messenger events for example) could increase the relative importance of \( I_M \) in firing of DA VTA neurons.

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Ethanol inhibition of M-current


