Motivational Effects of Ethanol in DARPP-32 Knock-Out Mice

Fred O. Risinger, Pierre A. Freeman, Paul Greengard, and Allen A. Fienberg

DARPP-32 (dopamine and adenosine 3',5'-monophosphate-regulated phosphoprotein, 32 kDa) is an important component of dopaminergic function in brain areas thought to be important for drug and alcohol addiction. The present experiments characterized the acquisition of ethanol-induced conditioned taste aversion, ethanol-induced conditioned place preference, and ethanol self-administration in DARPP-32 knock-out (KO) mice compared to wild-type (WT) controls. For taste conditioning, KO and WT mice received access to 0.2 M NaCl solution followed immediately by intraperitoneal injection of 0–4 gm/kg ethanol. Ethanol produced dose-dependent conditioned taste aversion that was the same in both genotypes. For place conditioning, KO and WT mice received eight pairings of a tactile stimulus with ethanol (2 gm/kg, i.p.), and a different stimulus with saline. Ethanol produced increases in locomotor activity during conditioning, with KO mice showing higher activity levels after ethanol compared to WT mice. WT mice, but not KO mice, acquired conditioned preference for the ethanol-paired stimulus. In the self-administration procedure, KO and WT mice were trained to lever press for access to 10% v/v ethanol. Subsequently, the mice had 23 hr/d access to food, ethanol, and water. Response patterns were determined using 0–30% v/v ethanol concentrations. WT mice displayed concentration-dependent responding for ethanol. Responding on the ethanol lever by KO mice did not change as a function of ethanol concentrations. Saccharin (0.2% w/v) was subsequently added to the ethanol mixture, and responding was examined at 0, 5, 10, and 20% ethanol concentrations. Ethanol responding increased in both genotypes, although WT mice showed higher rates at all concentrations.

Key words: ethanol; conditioned taste aversion; conditioned place preference; self-administration; reward; reinforcement; DARPP-32 knock-out mice

Contemporary views of addiction emphasize behaviors related to the experience of hedonic consequences after drug exposure (Wise, 1998). For example, individual differences in sensitivity to the rewarding or aversive effects of ethanol are thought to contribute to the development of excessive drinking (Tabakoff and Hoffman, 1988). Although several neurotransmitter systems contribute to ethanol reward (Koob et al., 1998), dopaminergic mechanisms have achieved the most prominence in neural circuitry models of the positive motivational effects of abused drugs including ethanol (Koob, 1999). A dopaminergic pathway important for reward function appears to be mesocorticolimbic brain areas, which include the ventral tegmental area, nucleus accumbens, olfactory tubercle, amygdala, frontal cortex, and septal area (Koob, 1992). Currently, dopamine receptor subtypes are classified as D1-like (D1 and D5 receptors) and D2-like (D2, D3, and D4 receptors) (Civelli et al., 1993). Both D1-like and D2-like receptors are found in reward-related brain areas (Civelli et al., 1993).

An important molecular mechanism for the effects of dopamine acting through D1- and D2-like receptors is protein phosphorylation (Greengard et al., 1998). One phosphoprotein, DARPP-32 (dopamine and adenosine 3',5'-monophosphate-regulated phosphoprotein, 32 kDa), appears particularly important for regulation of striatal dopaminergic systems (Greengard et al., 1999). To date, specific pharmacological antagonists for DARPP-32 are unavailable. However, mice with targeted disruption of the DARPP-32 gene have been recently developed, enabling investigations into the role of this protein in mediating the actions of dopamine as well as drugs of abuse (Fienberg et al., 1998; Fienberg and Greengard, 2000). DARPP-32 null mutants, which lack detectable DARPP-32 protein in the brain, appear developmentally normal but show diminished sensitivity to dopamine in electrophysiological and biochemical preparations (Fienberg and Greengard, 2000). DARPP-32 knock-out (KO) mice have normal levels of dopamine D1 and D2 receptors, but reduced immediate early gene expression after treatment with a D1 receptor agonist SKF 82958 (Svenningsson et al., 2000). In addition, acute cocaine-stimulated activity is reduced in DARPP-32 KO mice, as is raclopride-induced catalepsy (Fienberg et al., 1998).

The experiments reported here were devoted to characterizing sensitivity to ethanol's motivational effects in DARPP-32 KO mice, using several different procedures including taste conditioning, place conditioning, and ethanol self-administration. Each of these procedures has been associated with indexing the motivational effects of ethanol. Most self-administered drugs produce conditioned taste aversion, and this response has been hypothesized to be positively correlated with sensitivity to drug reward (Hunt and Amit, 1987). Place conditioning procedures are frequently used to index the rewarding properties of self-administered drugs as indicated by the acquisition of conditioned place preference (Carr et al., 1989; Tzschentke, 1998). Oral

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ethanol self-administration has long been regarded as an index of the rewarding efficacy of ethanol (Myers and Veal, 1972). We hypothesized that DARPP-32 mutant mice would display reduced sensitivity to the rewarding and aversive effects of ethanol, given that pharmacologically or molecularly induced reductions in dopamine function are associated with reductions in ethanol drinking (Dyr et al., 1993; Phillips et al., 1998), conditioned taste aversion (Risinger et al., 1999), ethanol-stimulated activity (Risinger et al., 1992a), and oral ethanol self-administration (Risinger et al., 2000).

MATERIALS AND METHODS

Animals. The present study used homozygous KO mice (−/−) in comparison to homozygous wild-type (WT) mice (+/+). Male congenic mice on a C57BL/6J background (N10) were produced at The Rockefeller University (Fienberg et al., 1998). Animals were between 4 and 6 months old at the beginning of training. Naïve mice were used for each experiment. Mice used in the taste conditioning study were housed individually in stainless-steel hanging cages (14 × 18 × 18 cm) with wire-mesh fronts and bottoms. Mice in the place conditioning study were housed individually in groups of four in polycarbonate cages (27.9 × 9.5 × 12.7 cm) with cob bedding. Mice in the operant self-administration study were initially housed in a single cage in polycarbonate cages. After training, mice in the operant self-administration study were housed individually in mouse operant chambers for 23 hr/d (see procedure below). For all studies, a 12 hr light/dark cycle was in effect (lights on at 7:00 A.M.), and the colony or testing rooms were maintained at an ambient temperature of 21 ± 1°C. The place conditioning and taste conditioning studies were conducted during the light cycle. For the ethanol self-administration study, lever response training was conducted during the light cycle. Lab chow was available ad libitum in the home cage. Animals in the taste conditioning and ethanol self-administration experiments had their access to fluids restricted as described herein.

Taste conditioning. The taste conditioning study was conducted in the home cages. Fluids were presented at room temperature in 25 ml graduated glass cylinders fitted with stainless-steel drinking spouts inserted through the front of the cage. Consumption was measured to the nearest 0.1 ml and was corrected for evaporation and spillage by subtracting the mean fluid loss measured in two drinking tubes placed on an empty cage for an equal amount of time. Mice of each genotype were randomly assigned to ethanol dose groups (n = 5–8/group). Subjects were adapted to a water restriction regimen (2 hr of water per day from 9:00–11:00 A.M.) for 48 hr (0.1 ml water was provided over 6 hr of water availability). At 48 hr intervals over the next 10 d, mice had access to a 0.2 M NaCl solution between 9:00 and 10:00 A.M. We have found NaCl to be an effective flavor stimulus in inbred and Swiss–Webster mice using this design (Risinger and Cunningham, 1992, 1995; Risinger, 1997; Risinger et al., 1999). For the first four trials (conditioning), immediately after access to the NaCl solution, each mouse received 10% v/v sucrose solution (i.e., fixed ratio 1 schedule of reinforcement). On the final test trial, mice were given access to NaCl, but did not receive ethanol. During conditioning, all mice also received 30 min access to tap water 5 hr after each NaCl access period, to prevent dehydration. On intervening days, subjects had 2 hr access to water (9:00–11:00 A.M.).

Place conditioning. The place conditioning apparatus consisted of eight identical acrylic and aluminum chambers (30 × 15 × 15 cm), each enclosed in a ventilated, light and sound-attenuating box (ENV-015M; MedAssociates, St. Albans, VT). Infrared light sources and detectors were positioned opposite each other at 5 cm intervals on the long walls of each chamber, 2.2 cm above the floor surface. Occlusion of the infrared light beams was used both as a measure of locomotor activity and to determine the animal’s position in the chamber. Data were recorded each minute by computer. The floor of each chamber consisted of interchangeable halves with one of two distinctive textures: “hole” floors were made from perforated stainless steel with 6.4 mm round holes on 9.5 mm staggered centers; “grid” floors were composed of 2.3 mm stainless-steel rods mounted 6.4 mm apart in Plexiglas rails. The place conditioning procedure was conducted daily Mondays through Fridays. The experimental sequence began with a 5 min habituation session, which was intended to reduce the novelty and stress associated with handling, injection, and exposure to the apparatus. All subjects received saline (10 ml/kg) and were immediately placed in the conditioning apparatus for 5 min on a smooth floor covered with paper. For conditioning, KO mice (n = 24) and WT mice (n = 24) were randomly assigned to one of two conditioning subgroups (n = 12/group) and exposed to an unbiased differential conditioning procedure. Conditioning was conducted using a between-group discrimination design (Cunningham, 1993), where conditioning trials consisted of pairings of a distinctive floor after ethanol exposure and pairings of a different floor with saline. Conditioning subgroups within each genotype were matched for exposure to ethanol and floor type and differed only in the specific floor-ethanol relationship (Cunningham, 1993). On alternate days mice received 2 gm/kg ethanol (CS− sessions) before placement on the grid floor (Grid+ subgroup) or the hole floor (Grid− subgroup). Mice received saline (CS− sessions) before placement on the other floor type. Presentation of CS+ and CS− sessions was counterbalanced for order of presentation. The 2 gm/kg ethanol dose was chosen because this dose produces reliable conditioned place preference in a variety of mouse genotypes (Risinger et al., 1994, 1996; Cunningham, 1995; Risinger and Oakes, 1996a). Eight conditioning sessions (four CS+, four CS−) were given before the first preference test. Four additional conditioning sessions were performed before the second preference test, and an additional four conditioning sessions performed before a third and final preference test. For each preference test, all subjects received saline injections before placement in the apparatus for a 60 min session with half grid floor and half hole floor (left–right position counterbalanced within groups).

Operant ethanol self-administration. Lever response training was conducted with four mouse operant chambers (modular mouse test chamber, ENV-307A; MedAssociates) each equipped with one ultra sensitive mouse lever (ENV-310; MedAssociates), liquid dipper with a 0.02 ml cup (ENV-303; MedAssociates), and 100 mL a house light. The house light was located on the opposite wall from the location of the lever and liquid dispenser, and was on when a lever was active. Each chamber was enclosed in a light–sound-attenuating cubicle (ENV-015M; MedAssociates). For 23 hr sessions, 16 mouse operant chambers (ENV-003; MedAssociates) enclosed in light/sound attenuating cubicles were used. Each chamber was equipped with two ultra-sensitive mouse levers, liquid dipper with a 0.02 ml cup, 20 mg pellet dispenser (ENV-203-20; MedAssociates), drinking tube, and house light. The access well for the liquid dipper was located in the center of the right side panel. The access well for the pellet dispenser was located in the center of the left panel. The levers were placed on the left side of the liquid dipper well and pellet dispenser. The drinking tube (25 ml glass graduated cylinder fitted with a stainless steel drinking spout) was located in the center of the front panel and connected to a contact lickometer (ENV-250A; MedAssociates). The house light was centered on the side panel 9.5 cm above the floor. Session parameters and data collection were controlled by computers adjacent to the chambers using MedAssociates interface modules.

During training, subjects received 2 hr access to water each day, 4 hr after training sessions. Subjects were first trained to lever press for 20% w/v sucrose solution. Initially, one lever press resulted in 10 sec access to 10% v/v sucrose solution (fixed ratio 1 schedule of reinforcement). During the course of a 10 d training phase, the schedule of reinforcement was gradually increased to FR4, and the dipper access period was reduced to 5 sec. When training was complete, the subjects entered a 15 d initiation phase during which an increasing concentration of ethanol was gradually introduced to the sucrose solution. The concentration of sucrose was gradually reduced such that at the end of this phase subjects were receiving access to 10% v/v ethanol in tap water. Eight mice of each genotype began and completed lever response training and the initiation phase.

After the initiation phase, subjects (n = 8 KO, 8 WT) were placed in operant chambers for 23 hr sessions. Initially, 10% v/v ethanol was available from the dipper (FR4), food from the pellet dispenser (20 mg Noxes formula A pellets; FR1), and water from the drinking tube. Each day, subjects were removed from the chamber for 1 hr to clean and resupply the chambers. A 12 hr light/dark cycle was maintained throughout the procedure.

The first session was used for acclimation to the chambers and procedure, and data from this session were not subjected to analysis of genotype differences. In addition, malfunction of the pellet dispenser in one chamber required the removal of one KO mouse from the study. Subjects completed phase 1 consisting of 4 weeks with 10% v/v ethanol available. At the end of phase 1, the concentration of ethanol was changed every four sessions (designated as phase 2). The following percentages of concentrations of ethanol were presented in the following order: 5, 10, 20, 30, and 0. For phase 3, 0.2% w/v saccharin was used.
as the ethanol vehicle. The addition of 0.2% saccharin was expected to increase overall ethanol consumption in both strains (cf. Risinger et al., 1998). The % v/v concentration of ethanol was changed every four sessions in the following order: 0, 5, 10, and 20.

Statistical analysis. ANOVA was used for all initial comparisons. The alpha level for all analyses was set at 0.05. For the floor preference tests in the place conditioning study, planned between-group comparisons examined conditioning subgroup for each genotype (Keppel, 1991). For the operant self-administration study, initial comparisons of genotype focused on daily session response rates for ethanol and food, and water intake. Further, a microanalysis of eating and drinking was conducted using a procedure that defined temporally related sequences of behavior as bouts (Samson et al., 1988). An ethanol bout consisted of four or more dipper presentations with ≤2 min between each dipper presentation. A food bout was defined as two or more pellet deliveries within ≤2 min. A water bout consisted of at least 25 consecutive licks with ≤2 min between each lick. The relationship between food intake and ethanol bouts was also determined (prandial bouts or nonprandial bouts). Prandial bouts occurred during or within 5 min of completion of a food bout. Nonprandial bouts occurred at least 5 min after the completion of a food bout.

RESULTS

Acquisition of ethanol-induced conditioned taste aversion

Figure 1 depicts mean (±SEM) NaCl intakes for each genotype over the course of the five NaCl access periods. Ethanol produced dose-dependent reductions in NaCl intake over trials, indicating the development of conditioned taste aversion. Both knock-out and wild-type mice showed similar levels of conditioned aversion, suggesting no influence of genotype in sensitivity to the aversive effects of ethanol measured in this design. Genotype x ethanol dose x trial analysis showed significant effects of ethanol dose ($F_{(2,37)} = 72.6; p < 0.001$), trial ($F_{(4,148)} = 22.7; p < 0.001$), and ethanol dose x trial ($F_{(8,148)} = 46.3; p < 0.001$). Effects of genotype, genotype x ethanol dose, genotype x trial, and genotype x ethanol dose x trial were nonsignificant (all $F$ values < 1.5).

Ethanol place preference conditioning

Locomotor activity levels measured during the place conditioning procedure indicated that each genotype had similar levels of activity in the absence of drug treatment. However, KO mice showed greater sensitivity to ethanol-stimulated activity than WT mice. Activity during the habituation session (activity counts per minute: KO mice, 107.3 ± 4.6; WT mice, 109 ± 5.1) was the same in both genotypes ($F_{(1,46)} = 0.11; p < 0.7$). Mean (±SEM) activity counts per minute during conditioning are shown in Figure 2. For each CS+ session, when subjects received 2 gm/kg ethanol, mean activity levels were higher than those seen on CS− sessions when subjects received saline. Genotype x session type (CS+/CS−) comparisons for each conditioning trial yielded significant effects of session type on each conditioning trial (all $F$ values$_{(1,46)} > 45.4; p$ values < 0.001). Significant genotype x session type effects were seen on conditioning trials 1–5 (all $F$ values$_{(1,46)} > 7.2; p$ values < 0.01), but not on conditioning trials 6–8 (all $F$ values$_{(1,46)} < 3.9; p < 0.053$). Comparisons of session type in each genotype separately yielded significant session type effects on all conditioning trials (all $F$ values$_{(1,23)} > 5.3; p$ values < 0.03). For CS+ sessions, significant genotype effects were noted on conditioning trials 1–5 (all $F$ values$_{(1,46)} > 7.2; p$ values < 0.01), but not conditioning trials 6–8 (all $F$ values$_{(1,46)} < 3.9; p$ values < 0.053). Genotype effects were not seen on any CS− session (all $F$ values$_{(1,46)} < 2.8, p$ values < 0.1). Overall, activity levels declined over the course of conditioning. Both genotypes showed reduced levels of activity under CS+ and CS− conditions after conditioning trial 1, indicating habituation to the apparatus and procedure. Analysis of activity levels over CS− sessions yielded a significant effect of session ($F_{(7,322)} = 68.3; p < 0.001$), but not genotype ($F_{(1,46)} = 0.7; p < 0.4$) or genotype x session ($F_{(7,322)} = 1.4; p < 0.2$). Analysis of activity levels over CS+ sessions yielded significant effects of genotype ($F_{(1,46)} = 14.0; p < 0.001$), session
Food was presented on an FR1 schedule of reinforcement. Each subject’s data was averaged into four session blocks. KO mice did not show evidence of place conditioning on any test (all \( F \) values_{(1,22)} < 0.2; \( p \) values < 0.7). In addition, analysis of conditioning group over test sessions for the WT mice yielded significant effects of conditioning group (\( F_{(1,22)} = 4.5; p < 0.05 \)) and test \( \times \) conditioning group (\( F_{(2,44)} = 5.9; p < 0.005 \)).

An alternative analysis of genotype differences in the preference tests used difference scores calculated for each subject of the total seconds spent on the ethanol-paired floor minus the seconds spent on the saline-paired floor. Mean (±SEM) difference scores for the KO mice were \( -2.2 \pm 218.4 \) on test 1, \( 15.5 \pm 392.4 \) on test 2, and \( -36.4 \pm 383.1 \) on test 3. Mean (±SEM) difference scores for the WT mice were \( 203.3 \pm 176.9 \) on test 1, \( 532 \pm 245.4 \) on test 2, and \( 686.5 \pm 270.7 \) on test 3. Although WT mice spent more time on the ethanol-paired floor compared to KO mice, significant genotype effects were not seen on any test (all \( F \) values_{(1,46)} < 2.4; \( p \) < 0.1).

Operant ethanol self-administration

Figures 4–6 depict ethanol level responding, food lever responding, and water intake for phases 1–3, respectively. Table 1 gives ethanol bouts, water bouts, and food bouts for each phase. Each subject’s data from phase 1 was averaged into four-session blocks. For phases 2 and 3, each subject’s data was averaged over each ethanol concentration. During phase 1, WT mice responded more overall on the ethanol lever than KO mice. Genotype \( \times \) trial block analysis showed a significant effect of genotype (\( F_{(1,13)} = 4.7; p < 0.05 \)), but not trial block (\( F_{(4,52)} = 2.0; p < 0.1 \)) or genotype \( \times \) trial block (\( F_{(4,52)} = 0.6; p < 0.7 \)). A similar analysis showed that food lever responding did not differ between KO and WT mice (genotype, \( F_{(1,13)} = 0.8; p < 0.3 \); trial block, \( F_{(4,52)} = 2.0; p < 0.1 \); genotype \( \times \) trial block, \( F_{(4,52)} = 0.4; p < 0.8 \)). Water intakes were similar for both genotypes and declined slightly over sessions. Analysis indicated a significant effect of trial block (\( F_{(4,52)} = 4.6; p < 0.005 \)), but not genotype (\( F_{(1,13)} = 1.2; p < 0.3 \)) or genotype \( \times \) trial block (\( F_{(4,52)} = 0.3; p < 0.8 \)). Both KO and WT mice displayed ethanol responding in the form of bouts, and ethanol bout frequency increased over the course of phase 1 (trial block: \( F_{(4,52)} = 4.1; p < 0.03 \)). Although WT mice showed higher bout frequencies per session, this difference did not reach statistical significance (genotype: \( F_{(1,13)} = 3.9; p < 0.07 \)). Analysis indicated that both KO and WT mice had overall similar frequencies of food lever bouts (genotype: \( F_{(1,13)} = 1.6; p < 0.2 \)). However, significant fluctuations in food bout frequency were noted (trial block, \( F_{(4,52)} = 7.9; p < 0.001 \); genotype \( \times \) trial block, \( F_{(4,52)} = 4.4; p < 0.004 \)). Follow-up analyses indicated WT mice had higher food bout frequencies compared to KO mice.
during the last trial block ($F_{(1,13)} = 10.9; p < 0.006$). Analysis of water bout frequency showed no significant effect of genotype ($F_{(1,13)} = 0.7; p < 0.4$), trial block ($F_{(4,52)} = 2.3; p < 0.09$), or genotype $\times$ trial block ($F_{(4,52)} = 0.4; p < 0.7$).

During phase 2, WT continued to show higher levels of responding on the ethanol level compared to KO mice at certain ethanol concentrations. Overall genotype $\times$ ethanol concentration analysis yielded significant effects of genotype ($F_{(1,13)} = 4.7; p < 0.05$) and ethanol concentration ($F_{(4,52)} = 3.7; p < 0.01$). Further analyses at each ethanol concentration indicated WT mice responded more on the ethanol lever than KO mice at the 20 and 30% concentrations (both $F$ values$(_{(1,13)} > 6.5; p$ values $< 0.02$, but not at the 0, 5, or 10% concentrations (all $F$ values$(_{(1,13)} < 3.2; p$ values $< 0.08$). WT mice showed concentration-dependent changes in responding ($F_{(4,28)} = 3.2; p < 0.03$), whereas KO mice did not ($F_{(4,24)} = 1.2; p < 0.5$). Paired $t$ comparisons indicated responding in WT mice was greater for 20 and 30% ethanol than for plain water ($t$ values$(_{(7)} > 3.5; p$ values $< 0.01$). Food lever responding was the same in both KO and WT mice and did not change when ethanol concentration was altered (Genotype: $F_{(1,13)} = 0.1; p < 0.7$; ethanol concentration: $F_{(4,52)} = 2.0; p < 0.1$; genotype $\times$ ethanol concentration, $F_{(4,52)} = 0.9; p < 0.5$). Water intakes were the same in both genotypes ($F_{(1,13)} = 2.1; p < 0.1$). WT mice showed a higher frequency of ethanol lever bouts compared to KO mice with analysis yielding a significant effect of genotype ($F_{(1,13)} = 4.6; p < 0.05$) and ethanol concentration ($F_{(4,52)} = 3.5; p < 0.01$). WT mice had higher numbers of food bouts compared to KO mice. Analysis yielded a significant effect of genotype ($F_{(1,13)} = 8.5; p < 0.01$) but not ethanol concentration ($F_{(4,52)} = 0.5; p < 0.7$) or genotype $\times$ ethanol concentration ($F_{(4,52)} = 0.4; p < 0.8$). Both KO and WT mice had similar frequencies of water bouts during phase 2 (Genotype: $F_{(1,13)} = 0.04; p < 0.8$).

The addition of saccharin in phase 3 increased overall responding in both genotypes. Compared to KO mice, WT mice showed higher response rates on the ethanol lever overall, even when ethanol was not present (i.e., 0.2% saccharin alone). Overall analysis yielded a significant effect of genotype ($F_{(1,13)} = 5.4; p < 0.04$) and ethanol concentration ($F_{(3,39)} = 5.6; p < 0.003$). Although analysis of ethanol concentration in WT mice indicated concentration-dependent changes in responding ($F_{(3,21)} = 3.6; p < 0.03$), follow-up paired $t$ comparisons indicated no significant differences across concentrations ($t$ values $< 1.9; p$ values $< 0.1$). However, KO mice showed significant changes in responding across ethanol concentration ($F_{(3,18)} = 6.8; p < 0.003$). Specifically, response rates were higher for 5% ethanol ($T_{(6)} = 2.9; p < 0.03$) and 10% ethanol ($T_{(6)} = 4.0; p < 0.007$) compared to response rates for saccharin alone. KO and WT mice did not differ in food lever responding ($F_{(1,13)} = 0.1; p < 0.8$), although food lever responding decreased with the addition of ethanol ($F_{(3,39)} = 24.6; p < 0.001$). KO mice consumed more water than WT mice (Genotype: $F_{(1,13)} = 5.2; p < 0.04$). Analysis also yielded a significant effect of ethanol concentration ($F_{(3,39)} = 22.7; p < 0.001$) and genotype $\times$ ethanol concentration ($F_{(3,39)} = 4.1; p < 0.01$). Follow-up analyses indicated both WT and KO mice differ from KO mice in ethanol lever responding but not water intake.

![Figure 5](image-url)  Figure 5. Mean ($\pm$ SEM) number of ethanol lever responses per session (left panel), food lever responses (middle panel), and milliliters of water intake (right panel) during phase 2 for DARPP-32 KO mice ($n = 7$) and WT mice ($n = 8$). Ethanol was presented on an FR4 schedule of reinforcement. Food was presented on an FR1 schedule of reinforcement. Subjects were presented with each ethanol concentration (% v/v) for four consecutive 23 hr sessions.

![Figure 6](image-url)  Figure 6. Mean ($\pm$ SEM) number of ethanol lever responses per session (left panel), food lever responses (middle panel), and milliliters of water intake (right panel) during phase 3 for DARPP-32 KO mice ($n = 7$) and WT mice ($n = 8$). Ethanol was presented on an FR4 schedule of reinforcement. Food was presented on an FR1 schedule of reinforcement. Subjects were presented with each ethanol concentration (% v/v) for four consecutive 23 hr sessions.
mice had significant declines in water intake over ethanol concentration (KO mice: $F_{(3,18)} = 24.9; p < 0.001$; WT mice: $F_{(3,21)} = 4.1; p < 0.02$). KO mice drank more water compared to WT mice at the 0% ethanol concentration ($F_{(1,13)} = 10.9; p < 0.006$) but not at the 5, 10, or 20% concentrations (all $F$ values $< 4.0; p < 0.06$). The number of ethanol lever bouts was increased in both genotypes compared to levels seen without saccharin (i.e., phase 1 and 2). However WT mice showed higher ethanol bout frequencies than KO mice with analysis yielding a significant effect of genotype ($F_{(1,13)} = 6.3; p < 0.03$) and ethanol concentration ($F_{(3,39)} = 6.6; p < 0.001$). WT mice also had higher food bout frequencies compared to KO mice ($F_{(1,13)} = 19.2; p < 0.001$). KO mice showed higher numbers of water bouts with the analysis yielding a significant genotype $\times$ ethanol concentration interaction ($F_{(3,39)} = 30.6; p < 0.001$). Both genotypes showed significant effects of ethanol concentration (KO: $F_{(3,18)} = 42.9$; WT: $F_{(3,21)} = 7.3; p < 0.002$). Significant genotype effects were only seen at the 0% concentration ($F_{(1,13)} = 5.2; p < 0.04$).

Table 2 gives mean (±SEM) gram per kilogram ethanol intakes based on the number of dippers presented during a session. As would be expected based on ethanol response rates, WT mice had higher gram per kilogram ethanol doses per session than KO mice during each experimental phase. A significant genotype effect was noted in phase 1 ($F_{(1,13)} = 5.4; p < 0.04$), but no effect of trial baseline phase.
Table 4. Mean (±SEM) number of prandial and nonprandial fluid lever bouts during each experimental phase

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<th>Prandial KO</th>
<th>Prandial WT</th>
<th>Nonprandial KO</th>
<th>Nonprandial WT</th>
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</tr>
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<td>Phase 3 Ethanol (% v/v)</td>
<td></td>
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</tr>
<tr>
<td>0</td>
<td>14.1 (4.1)</td>
<td>13.9 (2.2)</td>
<td>6.3 (0.8)</td>
<td>14.4 (3.2)</td>
</tr>
<tr>
<td>5</td>
<td>11.3 (1.0)</td>
<td>16.2 (1.9)</td>
<td>8.8 (1.2)</td>
<td>15.7 (2.1)</td>
</tr>
<tr>
<td>10</td>
<td>11.4 (0.9)</td>
<td>14.8 (2.1)</td>
<td>8.5 (0.8)</td>
<td>15.6 (2.5)</td>
</tr>
<tr>
<td>20</td>
<td>9.6 (1.3)</td>
<td>13.4 (1.6)</td>
<td>5.9 (1.2)</td>
<td>10.3 (2.4)</td>
</tr>
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block or genotype × trial block (both F values4,52 < 1.7; p < 0.1). Analysis of phase 2 doses yielded a significant effect of genotype (F values1,13 = 7.8; p < 0.02), ethanol concentration (F(3,39) = 42.4; p < 0.001), and genotype × ethanol concentration (F(3,39) = 4.7; p < 0.007). Both genotypes showed significant increases in ethanol dose with increasing ethanol concentration (KO mice: F(3,18) = 28.9; p < 0.001; WT mice: F(3,21) = 25.2; p < 0.001). Significant genotype effects were seen at the 20% concentration (F(1,13) = 8.5; p < 0.01) and the 30% concentration (F(1,13) = 6.3; p < 0.03). During phase 3, overall analysis yielded a significant effect of ethanol concentration (F(2,26) = 50.9; p < 0.001) but not genotype (F(1,13) = 3.8; p < 0.07) or genotype × ethanol concentration (F(2,26) = 1.7; p < 0.2).

Table 3 gives mean (±SEM) ethanol bout size for each phase for both genotypes. No effects of genotype were seen in analyses of each phase (all F values1,13 < 2.4; p values, 0.1). Bout size remained constant over trial block during phase 1 (F4,52 = 0.7; p < 0.6) and over ethanol concentration during phase 3 (F3,39 = 1.2; p < 0.3) with no interactions of genotype with trial block in phase 1 (F4,52 = 0.1; p < 1.0) or ethanol concentration in phase 3 (F3,39 = 1.2; p < 0.3). However, analysis of phase 2 ethanol bout size yielded significant effects of ethanol concentration (F4,52 = 7.1; p < 0.002) and genotype × ethanol concentration (F4,52 = 2.6; p < 0.05). Follow-up analyses indicated ethanol bout size in KO mice did not differ from WT mice at any ethanol concentration (all F values1,13 < 2.4; p values < 0.08).

Table 4 gives mean (±SEM) number of prandial and nonprandial bouts for each genotype during each phase. Each genotype generated both types of bouts, of which prandial bouts constituted ~50–60% of the total number. During phase 1, the frequency of prandial bouts increased over trials (F4,52 = 5.5; p < 0.001). Also, WT mice showed higher frequencies of prandial bouts compared to KO mice (F1,13 = 4.6; p < 0.05). Nonprandial bout frequency remained constant (F4,52 = 1.3; p < 0.3) and was similar in both genotypes (F(1,13) = 2.6; p < 0.1). Prandial bouts also increased during phase 2 (F4,52 = 4.6; p < 0.003) with WT mice showing higher prandial bout frequencies than KO mice (F(1,13) = 5.1; p < 0.04). As in phase 1, nonprandial bout frequency remained stable in phase 2 (F4,52 = 2.3; p < 0.07), and KO mice and WT mice had similar nonprandial bout frequencies (F(1,13) = 3.5; p < 0.08). The frequency of each bout type was higher in phase 3 for both genotypes. Prandial bout frequency was similar for KO and WT mice (F(1,13) = 1.4; p < 0.3) and remained stable over ethanol concentration (F3,39 = 0.7; p < 0.5). However, nonprandial bout frequency was higher in WT mice compared to KO mice (F1,13 = 6.1; p < 0.03). Also, nonprandial bout frequency decreased over ethanol concentration (F3,39 = 7.4; p < 0.001).

**DISCUSSION**

DARPP-32 KO mice self-administered less ethanol than WT mice across a variety of conditions, indicating a decrement in oral ethanol reinforcement. However, DARPP-32 KO mice and WT mice did not differ in the acquisition of ethanol-induced conditioned taste aversion, suggesting that sensitivity to the aversive effects of ethanol were equivalent. DARPP-32 KO mice showed enhanced sensitivity to ethanol-stimulated activity, indicating the KO mice did not have a general lack of response to ethanol. However, mice lacking DARPP-32 failed to acquire ethanol-induced conditioned place preference. Overall, these findings indicate DARPP-32 is an important component for sensitivity to ethanol reinforcement but not ethanol aversion.

Pairing a distinctive flavor with subsequent ethanol exposure generally results in a conditioned aversion to the flavor (Sherman et al., 1988). In the present study, both KO and WT mice displayed similar levels of conditioned taste aversion, showing that elimination of DARPP-32 had no influence on the aversive effects of ethanol measured in this design. However, pharmacological studies have indicated dopaminergic systems contribute to ethanol aversion. Specifically, dopamine D2 receptor blockade reduces the acquisition of ethanol-induced conditioned taste aversion (Sklar and Amit, 1977; Rissing et al., 1999). Dopamine D1 receptor or D4 receptor blockade only marginally reduce ethanol-conditioned taste aversion (Risinger et al., 1999; Thrasher et al., 1999). The present results do not show a positive relationship between ethanol reinforcement and sensitivity to ethanol-conditioned taste aversion. Furthermore, the present results are not consistent with the notion of dopaminergic mediation of ethanol aversion. However, ethanol appears to produce aversion by two mechanisms (Hunt and Amit, 1987). One mechanism is central in origin and resembles the mechanism of other reinforcing drugs. A second emotive mechanism relies on the accumulation of acetaldehyde (Aragon et al., 1991). Thus, the doses of ethanol used in the present study may have produced conditioned taste aversion by peripheral mechanisms in which central changes in dopaminergic systems would not be expected to be influential. For example, dopamine D2 receptor blockade does not prevent conditioned taste aversion produced by the emetic agent lithium chloride (Hunt et al., 1985). It should also be noted that reductions in NaCl intake may be related to conditioned behavioral changes other than those directly related to aversive motivational effects. For example, conditioned activation in the presence of the ethanol-associated flavor could have interfered with consumption.

In place conditioning using various mouse strains, pairing distinctive environmental cues with ethanol exposure results in subsequent conditioned place preference (Cunningham et al., 1992;
Risinger et al., 1994; Risinger and Oakes, 1996a). Place conditioning procedures have been used for examination of neuropharmacological mechanisms related to ethanol reward (Risinger et al., 1992a,b, 1996; Risinger and Oakes, 1996b; Thrasher et al., 1999). Neither dopaminergic nor opioid mechanisms, using pharmacological agents, have been implicated in the acquisition of ethanol-induced conditioned place preference (Risinger et al., 1992a; Cunningham et al., 1995; Thrasher et al., 1999). However, specific serotonergic receptor mechanisms appear particularly important for the acquisition of this task (Risinger and Oakes, 1996b; Risinger et al., 1996) as well as GABAergic receptor mechanisms (Chester and Cunningham, 1999a; although see Chester and Cunningham, 1999b; Risinger et al., 1992b). The results of the present study suggest a role for DARPP-32 in mediating ethanol reward. Specifically, lack of DARPP-32 is associated with lowered sensitivity to the rewarding effects of ethanol as illustrated by the failure of KO mice to acquire ethanol-induced conditioned place preference. However, the present results should be viewed with some caution. Although the comparisons of conditioning subgroup in the WT mice are consistent with the acquisition of conditioned preference, the overall magnitude of preference was modest compared to that seen in other mouse strains examined in a similar multiple conditioning–testing procedure (Risinger et al., 1996; Risinger and Oakes 1996a). This, in part, is likely attributable to the C57BL/6 background. Although C57BL/6 mice acquire place preference to ethanol paired cues (Kelly et al., 1997; Nocjar et al., 1999), this strain shows lower magnitudes of conditioned preference compared to other inbred mouse strains (e.g., DBA/2J; Cunningham et al., 1992). The lack of conditioning in KO mice and the modest conditioning noted in WT mice could be interpreted as attributable to a reduction in learning ability. However, results from the taste-conditioning study indicate KO and WT mice are able to learn Pavlovian associations. Furthermore, DARPP-32 KO mice are able to learn a food-reinforced operant task (Heyser et al., 2000). Thus, the place conditioning results are not likely attributable to a non-specific learning decrement or general insensitivity to the motivational effects of ethanol.

In mice, ethanol produces locomotor stimulation at doses ≤2 gm/kg (Risinger and Oakes, 1996a). Moreover, a positive relationship between ethanol-induced locomotor stimulation and ethanol reward has been proposed (Wise and Bozarth, 1987), and ethanol-stimulated activity is thought to reflect the rewarding euphoric effects of ethanol (Phillips and Shen, 1996). During conditioning, a 2 gm/kg ethanol dose produced locomotor stimulation, a response seen in a number of other mouse strains, including C57BL/6 (Cunningham, 1995). KO mice showed higher levels of activity after ethanol compared to WT mice. Both genotypes showed similar levels of activity after saline. Therefore, the notion of a positive relationship between ethanol-stimulated activity and ethanol reward was not supported, given that KO mice showed ethanol-induced stimulation of activity but not conditioned place preference. Dissociation of ethanol-stimulated activity and acquisition of ethanol-induced conditioned place preference has also been noted in studies using pharmacological manipulations. For example, haloperidol eliminates the locomotor stimulant effects of ethanol without influencing the magnitude of conditioned preference in a design similar to that used here (Risinger et al., 1992a). However, it has been noted that the relationship between ethanol-stimulated activity and the acquisition of ethanol-induced conditioned place preference depends on a variety of factors including gender, locomotor activity levels after initial exposure to ethanol or saline, and changes over multiple place conditioning trials in sensitivity to ethanol-stimulated activity (Nocjar et al., 1999). Thus, conclusions about differences in sensitivity to ethanol-stimulated activity and ethanol-conditioned place preference noted in the present genotypes await additional studies (e.g., ethanol dose manipulations).

The present study characterized oral ethanol self-administration using a relatively long-term procedure, with results indicating DARPP-32 exerts a modulating influence on ethanol-reinforced behavior. Responding for food and water intakes were similar in both genotypes. However, KO mice responded less on the ethanol-associated lever across a variety of ethanol-concentration conditions compared to WT mice. As expected, access to unsweetened oral ethanol was an effective reinforcing stimulus in WT mice with C57BL/6 lineage (cf. Risinger et al., 1998). For example, during phase 2, WT mice showed ethanol concentration-dependent responding, indicating ethanol is an effective reinforcer in this genotype. KO mice did not show concentration-dependent changes in responding for unsweetened ethanol. Although overall ethanol responding was reduced under all conditions in KO mice compared to WT mice, KO mice showed increases in ethanol-reaver responding when ethanol was added to a saccharin solution. KO mice also responded less for plain saccharin. Thus, KO mice display reductions in responding for more than one reinforcer, which is consistent with the notion of a general role for dopaminergic systems in reward processes (Wise et al., 1978) or in the production of instrumental responding (Salamone et al., 1997). WT mice generated a bout pattern of responding for ethanol similar to that seen in C57BL/6 mice. KO mice also produced ethanol bouts, although at lower levels than those seen in WT mice. Bout size remained relatively constant in both genotypes, with differences in ethanol intake determined by the number of bouts rather than bout size. Similar patterns have been noted in C57BL/6 mice (Risinger et al., 1998) and ethanol preferring P rats (Files et al., 1993). In addition, both genotypes showed prandial and nonprandial bouts. Higher ethanol intakes seen in WT mice compared to KO mice were based on greater frequencies of food-related drinking (i.e., prandial bouts) when ethanol was presented unsweetened. However, with sweetened ethanol available, higher ethanol intakes in WT mice were based on an increase in nonprandial drinking.

In summary, DARPP-32 KO mice showed specific decrements in ethanol reinforcement, but not in ethanol aversion. Food and water consumption were not altered in the KO mice. In contrast to the reduced sensitivity to ethanol reinforcement, DARPP-32 KO mice showed heightened sensitivity to ethanol-stimulated activity, but did not acquire ethanol-induced conditioned place preference. These results are generally consistent with pharmacological studies supporting a role for dopamine D1 and D2 receptor systems in the control of ethanol intake (Pfeffer and Samson, 1988; Dyr et al., 1993). Furthermore, mice lacking dopamine D1 receptors or dopamine D2 receptors consume less ethanol (El-Ghundi et al., 1998; Phillips et al., 1998). However, dopamine D2 receptor KO mice display a pronounced reduction in operant responding for a variety of reinforcers, including ethanol (Risinger et al., 2000). Dopamine system involvement in the production of ethanol-induced conditioned place preference appears less clear in studies showing that dopamine D2 and D4 receptor antagonists do not influence the acquisition of this response (Risinger et al., 1992a; Thrasher et al., 1999). However, DARPP-32 appears particularly important in the regulation of dopamine D1 receptor function (Fienberg et al., 1998). Results from D1 receptor-specific manipulations have not been reported, which may be an important mechanism for the production of ethanol conditioned place preference. Failure to
note differences between KO and WT mice in ethanol-conditioned
taste aversion is also consistent with a primary role for D1 mech-


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