Neuronal modulation of hepatic lipid accumulation induced by bingelike drinking


INTRODUCTION

Alcoholic binge drinking is prevalent. In 2015, about 1 in 6 adults in the United States admitted to binge drinking, with an average individual consumption of 7 drinks per episode and 470 binges per year (20). One of the major health consequences of excessive alcohol consumption is the development of acute and chronic liver disease. Hepatic steatosis, seen in more than 90% of heavy alcohol drinkers (1, 11, 45, 46), is an early consequence of excess alcohol consumption, including binge drinking, and can be induced by even a single dose of alcohol in rodents (21, 24, 48). Moreover, alcohol-induced hepatic steatosis predicts the pathogenesis of more chronic and severe diseases, including hepatitis, fibrosis, and ultimately cirrhosis (11, 26, 49).

However, the etiology of alcoholic fatty liver, although likely multifaceted, remains incompletely understood. One potential driver of this process is the direct effect of alcohol on the liver. Ethanol is oxidized by alcohol dehydrogenase (ADH) in the cytosol of hepatocytes to yield acetaldehyde, which is subsequently metabolized by aldehyde dehydrogenase (ALDH) to produce acetate (56). In addition, hepatocytes metabolize ethanol by the microsomal ethanol-oxidizing system, which is catalyzed by cytochrome P450 2E1 (CYP2E1) and catalase in peroxisomes (56). Ethanol also influences the activity of hepatic peroxisome proliferator-activated receptor-α (PPARα), sterol regulatory element binding protein 1c (SREBP-1c), and AMP-dependent protein kinase (AMPK), which in turn inhibit fatty acid oxidation and stimulate de novo lipogenesis, leading to fat accumulation (11, 26, 54). Alcohol may also exert indirect effects on the liver, by disrupting the metabolic function of adipocytes (41), promoting white adipose tissue lipolysis, and enhancing free fatty acid flux to liver (50, 57).

Notably, evidence also suggests that adenosine signaling plays an important role in the etiology of alcoholic liver diseases (13). All cells generate adenosine during the breakdown of ATP. Alcohol increases extracellular adenosine levels by at least two mechanisms (13). First, the liver oxidizes ethanol to yield acetate, which is metabolized to adenosine, thus increasing systemic adenosine levels when alcohol is consumed in excess (6, 40). Second, ethanol inhibits purine reuptake through the type I equilibrative nucleoside transporter (ENT1) in tissues including neurons, leading to the buildup of extracellular adenosine (31, 37). There are four adenosine receptors in vertebrates, termed A1, A2A, A2B, and A3 (16). Notably, mice with germline deletion of A1 or A2B, but not A2A, are resistant to alcohol-induced hepatic steatosis (43).
While the roles of the liver and white adipose tissue in alcohol-induced hepatic lipid accumulation have been extensively investigated, the role of the brain, by comparison, has not. Upon consumption, alcohol quickly enters the brain, altering neuronal functions and impairing neurological, cognitive, and social behaviors. Increasing evidence suggests that alcohol may also exert metabolic effects through the brain. For example, binge drinking rapidly induces systemic insulin resistance by impairing hypothalamic insulin signaling (27). Within the hypothalamus, agouti-related protein (AgRP) is a neuropeptide that is expressed by a group of neurons in the mediobasal hypothalamus. AgRP antagonizes the action of α-melanocyte-stimulating hormone (α-MSH), a product of the proopiomelanocortin (POMC)-expressing neurons. Stimulating insulin signaling in AgRP neurons directly inhibits their activity and leads to suppression of hepatic glucose production (23). AgRP neurons regulate peripheral tissues including liver and adipose tissues via modulation of the autonomic nervous system innervating these tissues (2). For example, we recently showed that AgRP inhibits hepatic sympathetic activity and that eliminating AgRP in leptin-deficient mice alters both hepatic lipid accumulation and distribution in an age-dependent manner (29).

Notably, direct alcohol exposure stimulates the electrical activity of AgRP neurons (5), and ethanol dose-dependently increases hypothalamic AgRP immunoreactivity within 2 h of systemic injection (12). Conversely, alcohol-associated over-eating is mediated via activation of AgRP neurons (5). Intracerebroventricular infusion of AgRP promotes alcohol drinking, whereas AgRP deficiency reduces ethanol-reinforced lever-pressing and bingelike alcohol drinking (33, 35, 36). Together, these results suggest that AgRP neurons are modulated by alcohol and that some of the metabolic effects of alcohol may depend on AgRP neurons.

Intriguingly, recent evidence also suggests that presynaptic adenosine signaling via the A1 receptor inhibits AgRP neuronal activity (52). Adenosine is a potent neuromodulator, regulating neuronal activity and modulating signaling by other neurotransmitters (17). Alcohol exposure acutely increases central nervous system (CNS) adenosine levels, and elevated adenosine levels mediate the ataxic and sedative effects of ethanol through A1 receptor activation in the cerebellum, striatum, and cerebral cortex (44). In contrast to presynaptic A1 receptors, which are Gαi3-coupled G protein-coupled receptors (GPCRs), A2B is a Gαo-coupled postsynaptic GPCR. Of note, Gαi signaling potently regulates the function of AgRP neurons (32). This, along with the observations that alcohol increases AgRP neuronal activity (5) and expression (12), and that mice lacking A2B are resistant to alcohol-induced hepatic steatosis (43), led us to investigate if neuronal A2B signaling and AgRP contribute to the development of alcoholic fatty liver disease induced by bingelike drinking.

MATERIALS AND METHODS

Animals

Mice were housed in a barrier facility under a 12-h light cycle (lights on from 7 AM to 7 PM) with ad libitum access to water and standard mouse chow (21.6%, 23.2%, and 55.2% kcal from fat, protein, and carbohydrate, respectively; Purina mouse diet no. 5058). Agrp−/− mice were originally provided by Dr. Gregory Barsh at Stanford University. Transgenic mice expressing Cre recombinase under control of the Nestin promoter [B6.Cg-Tg(Nestin-Cre);1LtmJ/I] or the AgRP promoter [Agrpim1(cre);L Alloy/I] were originally obtained from the Jackson Laboratory. Adora2bflox/flox mice were provided by Dr. Holger Eltzschig at University of Colorado. To generate Agrp−/− mice, Agrp+/+ mice were bred with Agrp−/− mice to generate equal ratio of control (Agrp+/+) and mutant (Agrp−/−) mice. Littermates were compared whenever possible to minimize differences in genetic background and litter-specific effects. Adora2bflox/flox females were mated with Nestin-Cre+/+; Adora2b−/− males to generate mice lacking the A2B receptor in the nervous system (CNS-Adora2b−/−; Nestin-Cre+/+, Adora2bflox/flox) and their littermate controls (Nestin-Cre+/+; Adora2bflox/flox). Since mice carrying Nestin-Cre alone have a body weight phenotype (4), mice expressing Nestin-Cre were used as controls. To generate mice lacking the A2B receptor in AgRP neurons, Adora2bflox/flox females were mated with Adora2b−/−; Agrp-Cre males. This breeding resulted in control (Adora2bflox/flox or Adora2b−/−) and AgRP-specific A2B knockout (Agrp-Cre+/+; Adora2b−/−; Agrp-Cre−/−) mice (AgRP-Adora2b−/−). For the fasting experiment, 23-wk-old mice were fasted to 1000 and euthanized at the onset of the dark phase (1900). All experiments were approved by the University of California San Francisco Institutional Animal Care and Use Committee.

Alcohol Treatment

Preparation, administration, and dosing of alcohol in mice followed the protocol of Bertola et al. (3). Briefly, at 1900 on day 1, mice were given an oral gavage of ethanol (5 g/kg body wt), while control mice were given isocaloric dextrin (control, 9 g/kg body wt). At 800 on day 2, mice received a repeat oral gavage of either ethanol or vehicle and were euthanized at around 1700. The entire experiment was performed under thermoneutral ambient conditions (30°C), as alcohol administration can induce hypothermia in mice housed at room temperature. Blood alcohol content was measured as described previously (49).

RNA-Interference

RNA-interference was used to knockdown Agrp mRNA levels in 8-wk-old C57BL/6J mice (Jackson Laboratories) following a published protocol (47). The experiment was conducted at 30°C thermoneutrality. Briefly, at 1900 on day 1, half of the mice (n = 12, 2 mice per cage) were given an intraperitoneal injection of Agrp-DsiRNA (Integrated DNA Technologies, Inc., MEC-RNALON07427.12.1.5 μg per mouse) while an equal number received a control injection (Integrated DNA Technologies, Inc., Ds NC1, 5 μg per mouse). Subsequently, all mice received an initial ethanol gavage. At 0700 on day 2, all mice received another injection of either Agrp-DsiRNA or control, after which they received a second dose of ethanol. Tissues were then harvested at 1700 on day 2, alternating between control and Agrp-DsiRNA-treated mice.

Chemical Sympathectomy

At 15–16 wk of age, X male C57BL/6J mice underwent chemical sympathectomy by injection with a single dose of 6-hydroxydopamine (6-OHDA, 250 mg/kg in 0.9% NaCl and 10−7 M ascorbic acid, Sigma-Aldrich, St. Louis, MO) versus vehicle (0.9% NaCl and 10−7 M ascorbic acid). Body weights were monitored, and mice were allowed to recover for 10 days. The effectiveness of sympathectomy was validated by confirming the complete ablation of NPY-positive sympathetic fibers in the livers of the 6-OHDA-treated mice.

Body Composition and Metabolic Analysis

Body composition was measured using an EchoMRI-700 machine (Echo Medical Systems, LLC, Houston, TX). Indirect calorimetry, locomotor activity, and food intake were measured over 5 days at 30°C.
Measurement of Plasma and Tissue Lipid Levels

Several different direct and indirect methods were used to measure tissue triacylglycerol (TAG) levels and are specified in each figure.

Enzymatic TAG determination. Forty to fifty milligrams of liver were homogenized in 500 μl of buffer (250 mM sucrose, 50 mM Tris-HCl pH 7.4), and lipid levels were determined using an enzymatic TAG determination kit (TR0100, Sigma-Aldrich, St. Louis, MO) according to the manufacturer’s instructions. This approach measured total tissue glycerol content, which is the sum of both TAG-associated and free glycerol.

Thin layer chromatography. Forty to fifty milligrams of liver were homogenized in 500 μl of buffer (250 mM sucrose, 50 mM Tris-HCl pH 7.4). Lipids from homogenized samples were extracted in a 2:1 mixture of chloroform and methanol (15), adsorbed onto silica-coated glass slides, and then separated along with serial dilutions of similarly dissolved TAG standards by thin layer chromatography (TLC) using a solvent containing hexane-diethyl ether-acetic acid (80:20:2, vol/vol). This approach allows for direct assessment of tissue TAG levels.

Oil-red-O staining. Liver sections from mice treated with control and AgRP-DsIRNAs were prepared and stained with Oil-Red-O (Sigma-Aldrich, St. Louis, MO), which stains neutral lipids, including TAG. The protocol for visualizing and quantifying Oil-Red-O staining intensity in liver sections was based on that by Mihlem et al. (30). Tissue protein content, for normalization, was quantified with Bradford Reagent (B6916, Sigma-Aldrich, St. Louis, MO). Plasma TAG levels were measured using the same determination kit that was used for liver samples. Plasma free fatty acids were measured by the University of California-Davis Mouse Metabolic Phenotyping Center (MMPC).

Lipidomics

Liver samples were weighed and flash frozen. Lipidomic analyses were performed using mass spectrometry as previously described by Louie et al. (28). Specific lipid abundances were normalized to tissue weights and internal standards, i.e., dodecylglycerol for the positive mode and pentadecanoic acid for the negative mode.

Gene Expression Studies

Total RNA from mouse tissues was extracted using the RNeasy Lipid Tissue Mini Kit (Qiagen, Valencia, CA). Gene expression was assessed by quantitative PCR using Taqman gene expression assays (Thermo Fisher Scientific, Inc.). For hypotalamic tissues, relative mRNA levels of Adora2b (mm00839329_m1) and Adora1 (mm1308023_m1) were normalized to that encoding 18S ribosomal RNA (Rn18S; mm03928990_g1). For liver samples, mRNA levels of the gene encoding aldehyde dehydrogenase 1 (Adh1; mm00507711_m1) were normalized to that encoding beta-actin (Actb).

Western Blot Analysis

Approximately thirty milligrams of each liver sample were homogenized in 1 mL of RIPA buffer (sc-364162A), denatured by Laemml Sample buffer (Bio-Rad no. 1610737) and beta-mercaptoethanol, and then subjected to SDS-PAGE (Bio-Rad no. 4561104) for fractionation. Gels were then electroblotted onto polyvinylidene difluoride (PVDF) membranes (Bio-Rad no. 162-0177) at 120 mA for 100 min. Membranes were then blocked with dried nonfat milk in TBST (20 mM Tris, pH 7.6, 140 mM NaCl, and 0.1% Tween 20) and subsequently incubated with antibodies against either ADH1 (Cell Signaling no. 5295, 1:1,000) or GAPDH (Santa Cruz Biotechnology no. 25778, 1:10,000) in 5% BSA for 12 h at 4°C. After being washed three times, the membranes were incubated with an horseradish peroxidase (HRP)-conjugated anti-rabbit antibody diluted in nonfat milk (1:2,000) for 1 h. Membranes were then washed with TBST five times and developed with the SuperSignal West Pico Chemiluminescent Substrate (Thermoscientific no. 34079) per manufacturer’s protocol. Protein bands were analyzed and quantified using ImageJ.

Immunofluorescence

AgRP-Cre: Adora2b^flox/flox mice were anesthetized with ketamine/xylazine and perfused with 4% (wt/vol) paraformaldehyde (PFA). Brains were excised and postfixed in 4% (wt/vol) PFA solution for 2 h at 4°C AND then transferred into a 30% sucrose solution for cryoprotection at 4°C overnight. Samples were then embedded in Shandon M-1 embedding matrix (Thermo Fisher Scientific, Inc., Carlsbad, CA) and stored at −80°C. Brain samples were sectioned using a cryostat to obtain 10-μm-thick coronal sections and double stained by simultaneous incubation at 4°C overnight with primary antibodies against c-Fos (1:4,000, Santa Cruz Biotechnology, Inc., Santa Cruz, CA) and AgRP (1:1,000, Neuronics, Minneapolis, MN). Sections were then washed and incubated at room temperature for 1 h with secondary donkey anti-rabbit Alexa 594 and donkey anti-goat Alexa 488 (1:200, Thermo Fisher Scientific, Inc.) antibodies, respectively. Nuclei were detected by counterstaining for 4′,6-diamidino-2-phenylindole (DAPI, 1:2,000) during incubation with the secondary antibody. Stained sections were prepared using Vectashield (Vector Laboratories, Burlingame, CA) mounting media. For assessing hepatic NPY expression, liver sections were immunostained with a polyclonal antibody against NPY (1:1,000, Peninsula Laboratories, San Carlos, CA) following the same experimental procedures as above.

Images were captured using an Olympus BX51WI microscope equipped with a QImaging Retiga 2000R digital camera. Images were taken with the same exposure, avoiding pixel saturation using QCapture Pro 6 (Qimaging, Surrey, BC, Canada). Only sections processed from the same experiments were compared. Image J (NIH, Bethesda, MD) was used to count c-Fos-positive cells. Investigators were blinded during specific cell counting from the images. Sections were matched by anatomical landmarks according to Bregma (anterior, Bregma: −1.84 mm; medial, Bregma −1.94 mm; posterior, Bregma −2.06 mm) using the Mouse Brain Atlas. At least two sections per region per mouse were quantified and then averaged. Only overlapping DAPI and c-Fos signals were used to count c-Fos-positive cell numbers.

Statistical Analysis

Statistical analyses were performed using Prism software (GraphPad Software, Inc., La Jolla, CA), and specific statistical tests for different experiments are described in the figure legends. Briefly, the two-tailed Student’s t test was used to compare two independent groups of mice, and a two-way ANOVA with multiple comparisons was used when two genotypes and two treatments were compared. Repeated-measures two-way ANOVA was used to analyze data captured from sets of mice monitored repetitively over time.

RESULTS

AgRP-Deficient Mice Have Reduced Hepatic Lipid Accumulation in Response to Bingelike Alcohol Consumption

Given that alcohol consumption impacts AgRP neuronal activity and neuropeptide levels (5, 12) and that AgRP neurons are implicated in regulating hepatic lipid metabolism (29), we sought to determine if AgRP plays a role in the development of hepatic steatosis induced by acute bingelike drinking by examining mice that lack the AgRP neuropeptide (Agrp^−/− mice). Agrp^−/− and wild-type mice had similar body weights and terminal levels of blood glucose, free fatty acids and...
plasma TAG following receipt of either alcohol or maltose dextrin by oral gavage (Fig. 1, A–H). Food intake was also similar between genotypes (Fig. 1I). These data suggest that under our experimental conditions acute bingelike alcohol consumption or AgRP deficiency does not have significant impact on either energy balance or circulating indicators of metabolic function.

However, this was notably not the case within the liver, as Agrp<sup>−/−</sup> mice had reduced hepatic total glycerol content (both TAG-associated and free glycerol) following alcohol gavage than did control mice, whereas there were no differences between the two genotypes following gavage with isocaloric maltose dextrin (Fig. 2, A and B). There was no correlation between rates of food intake and hepatic glycerol content (Fig. 2C). Moreover, this protection was not likely due to gross alterations in systemic ethanol metabolism, as both mutant and control mice had similar levels of blood alcohol and hepatic alcohol dehydrogenase (ADH1), assessed by both mRNA and protein expression (Supplemental Fig. S1, A–D; all Supplemental material is available at https://doi.org/10.6084/m9.figshare.11786766).

To identify specific hepatic lipid species that are affected by acute alcohol consumption in an AgRP-dependent manner, we performed lipidomic analysis on the livers of Agrp<sup>−/−</sup> and control mice that were given oral gavage of either maltose dextrin or alcohol (complete data set shown in Supplemental Table S1). Many of the 117 hepatic lipid species we examined were different in abundance following alcohol versus maltose dextrin treatment. To this end, we applied a two-tier criterion to identify lipid species that are regulated by alcohol consumption more so in control than in mutant mice. First, we identified lipid species that were significantly up- or downregulated in alcohol-treated, versus maltose dextrin-treated, control mice. Next, we asked whether the levels of any of these lipid species were significantly different between alcohol-treated control and Agrp<sup>−/−</sup> mice. Three lipid species, namely C16:0/C20:4/C16:0 TAG (1 of the 6 different TAGs), C18:0/C20:4 diacylglycerol (DAG), and cholesterol (red dots in Fig. 2D) met this criterion. Upon reanalyzing these three lipid species by two-way ANOVA with Sidak’s multiple comparisons test, we found that whereas both hepatic cholesterol and C18:0/C20:4

Fig. 1. Acute bingelike alcohol consumption does not alter energy balance in control or AgRP-deficient mice. Agouti-related protein (AgRP)-deficient mice [Agrp<sup>−/−</sup> knockout (KO); n = 10] and control mice (Agrp<sup>+/+</sup>, Ctrl; n = 12) were given oral gavage of alcohol (5 g ethanol per kg body wt). Another cohort of control and mutant mice was given oral gavage of isocaloric maltose dextrin (n = 6 each). Body weight (A and B), terminal blood glucose (C and D), plasma free fatty acids (E and F), plasma triglycerides (G and H), and food intake (I) were determined. Data are means ± SD. *P < 0.05; ns, nonsignificant. Student’s t test was used to compare 2 independent groups.
DAG levels were reduced by alcohol treatment more profoundly in control, versus mutant mice, the absolute hepatic levels of these species were not different between the two genotypes in the context of alcohol treatment (Fig. 2, E and F). By contrast, hepatic C16:0/C20:4/C16:0 TAG content was found to be upregulated by bingelike alcohol consumption in control, but not in Agrp−/− mice and also to have a reduced absolute abundance in alcohol-treated Agrp−/− versus control mice (Fig. 2G). These data together suggest that AgRP mediates the ability of alcohol to alter hepatic levels of C16:0/C20:4/C16:0 TAG, which is a highly abundant hepatic TAG.

Downregulation of AgRP by RNA-Interference Reproduces the Protection against Alcohol-Induced Hepatic Steatosis Seen in Agrp−/− Mice

One caveat to using Agrp−/− mice is the potential for developmental compensation that could confound phenotypic

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Fig. 2. Agouti-related protein (AgRP)-deficient mice have reduced hepatic steatosis in response to bingelike alcohol consumption. A and B: liver total glycerol content [triacylglycerol (TAG)-derived and free glycerol] of control and AgRP-knockout (KO) mice in response to oral gavage of maltose dextrin and alcohol. C: food intake did not show significant correlation with liver total glycerol. D–G: lipidomics analysis of liver tissues from maltose dextrin (MD)- or alcohol (Alc)-treated Agrp−/− control (Ctrl; n = 5) and Agrp−/− (KO; n = 5) mice. Designation of color-coded lipid species are indicated on the graph. With the use of Student’s t-test, a 2-tier criterion was applied to identify lipid species that are regulated by alcohol more so in control than in KO mice. E and G: analysis of 3 significant lipid species from D by two-way ANOVA with Sidak’s multiple comparisons test. Lipid abundance represents relative values that were normalized to internal standard dodecylglycerol (positive mode), pentadecanoic acid (negative mode), and tissue weight. Data are means ± SD. *P < 0.05, **P < 0.01, ****P < 0.0001; ns, nonsignificant.
outcomes. Therefore, we sought to knockdown AgRP function in adult animals and examine its effects on alcohol-induced hepatic steatosis. Small interfering RNAs (siRNA) are an effective tool for this purpose (51). Dicer-substrate RNAs (DsiRNAs) are chemically synthesized 27-mer RNA duplexes with a potency up to 100-fold greater than that of traditional siRNAs (22). Notably, peripheral injection of DsiRNAs against AgRP led to specific knockdown of hypothalamic Agrp but not Npy expression (47). This is likely due to the fact that the majority of AgRP neurons are uniquely situated outside the blood-brain barrier, allowing them to readily take up circulating substances (39, 55). To further evaluate the role of AgRP in alcohol-induced hepatic steatosis, we therefore used this DsiRNA approach to acutely knockdown AgRP expression in C57BL/6 mice, an inbred strain sensitive to alcohol.

Eight-week-old male C57BL/6J mice were divided into two weight-matched groups and housed under thermonutral conditions (30°C) to minimize any potential confounding effects of AgRP knockdown on thermogenesis and consequent energy balance. The subsequent treatment protocol took place over two successive days. At both 7 PM on day 1 and 7 AM on day 2, the mice were intraperitoneally injected with either Agrp-DsiRNA or negative control (NC DsiRNA) and then a dose of alcohol by oral gavage. Food intake and blood glucose levels were measured, and tissues were harvested 10 h after the second treatment. Liver lipid content was analyzed using a colorimetric TAG assay from homogenized liver lysates. When compared with NC-DsiRNA, Agrp-DsiRNA treatment did not impact body weight, food intake, blood glucose, or plasma TAG levels (Fig. 3, A–D). However, it reduced both the concentration of plasma free fatty acids and hepatic total glycerol content (TAG-derived and free glycerol) in the setting of alcohol treatment (Fig. 3, E and F). The findings obtained by analyzing total hepatic glycerol content were corroborated by histological data, as the intensity of Oil-Red-O staining, which analyzing total hepatic glycerol content were corroborated by glycerol content (TAG-derived and free glycerol) in the setting of alcohol-induced fatty liver seen in AgRP (Supplemental Fig. S2). Thus acutely knocking down reduced acute alcohol-induced hepatic steatosis in the absence of AgRP (Supplemental Fig. S2). Thus acutely knocking down AgRP in adult mice reproduces the protection against alcohol-induced fatty liver seen in Agrp−/− mice.

**Adenosine Signaling in the CNS via the A2B Receptor Limits Hepatic Steatosis Induced by Bingelike Drinking**

Mice with deletion of the A2B gene (Adora2b) are also protected against the development of alcoholic fatty liver (43). To determine if this protection is mediated within the CNS, we generated mice lacking Adora2b specifically in the CNS by crossing Adora2b flox/flox mice with Tg.nestin-Cre mice (CNS-Adora2b−/−). Since mice expressing the Tg.nestin-Cre transgene are known to have a metabolic phenotype (4), we used control mice that also expressed this transgene (Tg.nestin-Cre, Adora2b flox/flox mice).

We confirmed the desired specificity of the knockout by observing that hypothalamic Adora2b mRNA expression was abolished in CNS-Adora2b−/− mice, both at baseline and after the alcohol gavage protocol, whereas Adora1 mRNA expression was not affected (Fig. 4, A and B).

**CNS-Adora2b−/− mice showed no alteration in body weight, lean mass, fat mass, respiratory exchange ratio (RER), or energy expenditure (Fig. 4C and Supplemental Fig. 3) nor any changes in ad libitum food intake, as measured by CLAMS (Fig. 4, D and E), when compared with control mice. Although CNS-Adora2b−/− mice did exhibit a nonsignificant trend toward reduced food intake during the initial hours of refeeding (Fig. 4D), their food intake was not different from that of control mice upon exposure to alcohol (Fig. 4F).** By contrast, whereas total hepatic glycerol content (both TAG-associated and free glycerol) was similar between CNS-Adora2b−/− and control mice at baseline (Fig. 4G), CNS-Adora2b−/− mice had significantly higher total hepatic glycerol content after acute alcohol consumption (Fig. 4H). These results suggest that, independent of caloric or macronutrient consumption, A2B signaling in the CNS acts to limit the ability of bingelike alcohol consumption to induce acute hepatic steatosis.

**Deletion of Adora2b in AgRP Neurons Is Not Sufficient to Alter Alcohol-Induced Hepatic Steatosis**

We next explored the possibility that A2B signaling plays a role in the mechanism by which AgRP neurons impact alcohol-induced hepatic steatosis. To this end, we generated mice that lack Adora2b specifically in AgRP neurons (AgRP-Adora2b−/−). To assess the effects of A2B deficiency on neuronal functions, male control and AgRP-Adora2b−/− mice were euthanized in the late afternoon after a 9-h daytime fast. The mice were then perfused, hypothalamic sections were prepared, and c-Fos expression was examined by immunofluorescence analysis. Fasted AgRP-Adora2b−/− mice had reduced c-Fos expression specifically in the mediobasal hypothalamus (DMH) (Supplemental Fig. S4, A–C) Despite this, however, AgRP-Adora2b−/− and control mice had no differences in body weight, body composition (lean and fat mass), food intake, energy expenditure, RER, or locomotor activity (Supplemental Fig. S4D and Supplemental Fig. S5).

To assess the impact of A2B signaling specifically in AgRP neurons on alcohol-induced hepatic steatosis, adiposity-matched AgRP-Adora2b−/− and control mice were given oral gavages of alcohol per the protocol described above. There were no differences in hepatic total glycerol content (TAG-associated and free glycerol) between the two genotypes, either at baseline (untreated) or in response to alcohol gavage (Supplemental Fig. S4, E and F) suggesting that abrogating A2B signaling in AgRP neurons is not sufficient to alter the acute effects of alcohol on hepatic steatosis. Given that we also found that brain-specific removal of Adora2b (CNS-Adora2b−/−) increases the sensitivity to alcohol-induced fatty liver, our data with AgRP-Adora2b−/− mice open up the possibility that A2B signaling may influence alcohol-induced liver fat accumulation either by acting directly on other components of AgRP neuronal circuits that do not include AgRP neurons.

**Chemical Sympathectomy Attenuates the Extent of Hepatic Steatosis Induced by Bingelike Alcohol Consumption**

Our results, probing both AgRP and A2B, support the concept that the brain plays an integral role to the development of...
bping alcohol-induced fatty liver. The brain regulates peripheral lipid metabolism by modulating the function of the autonomic nervous system innervating peripheral metabolic tissues. In particular, the AgRP neuropeptide inhibits sympathetic nervous system (SNS) activity in both the white adipose tissue and liver (29, 38). To this end, we sought to determine if intact SNS function is required for the development of hepatic steatosis induced by binge alcohol consumption.

Fifteen- to sixteen-week-old male C57BL/6J mice underwent chemical sympathectomy by injection with a single dose of 6-hydroxydopamine (6-OHDA, 250 mg/kg) or vehicle control (ascorbic acid). The effectiveness of this treatment was validated by confirming the complete ablation of NPY-positive sympathetic fibers in the liver of 6-OHDA-treated mice in pilot experiments (Supplemental Fig. S6). Ten days after 6-OHDA or vehicle injection, weight-matched mice were given a single oral gavage of alcohol. To minimize the likelihood of potential confounds resulting from any effects of SNS activity on carbohydrate metabolism, we opted to use gavage with water in a separate cohort of mice as a second control in addition to isocaloric maltose dextrin in this experiment. After alcohol or control treatment, food was subsequently removed from the cages housing the mice, and their livers were harvested 8 h later. 6-OHDA treatment affected body weight and blood glucose similarly when mice were treated with water, alcohol, or isocaloric maltose dextrin (Fig. 5, A–F). However, whereas chemical sympathectomy did not affect hepatic TAG content in water- or maltose dextrin-treated mice, it reduced hepatic TAG...
content in alcohol-treated mice, indicating that an intact SNS is needed for the full ability of acute alcohol consumption to induce hepatic steatosis (Fig. 5, G–K, and Supplemental Figs. S7 and S8).

**DISCUSSION**

Binge drinking is prevalent, is costly, and has serious health consequences (20, 53). Hepatic steatosis develops quickly after acute binge drinking (21, 24, 48), and marks an early stage of liver injury. Alcohol-induced hepatic steatosis is currently understood to be mediated by peripheral mechanisms, such as the direct toxic effects of alcohol in the liver and adipose tissues. In this study, we provide multiple lines of evidence to support the concept that the brain and SNS play important modulatory roles in the full development of hepatic steatosis after bingelike alcohol consumption. Specifically, we show that hypothalamic AgRP, CNS adenosine signaling, and an intact SNS are integral components of a regulatory program impacting hepatic TAG accumulation in the context of acute alcohol consumption.

AgRP neurons have previously been reported to mediate behavioral consequences of alcohol consumption. For example, AgRP infusion into the brain promotes alcohol drinking, whereas AgRP deficiency reduces alcohol-reinforced lever-pressing and bingelike alcohol drinking (33, 35, 36). Here we show that AgRP is required for the full development of acute
alcohol-induced hepatic steatosis, an indicator of the role of AgRP in mediating key metabolic consequences of alcohol consumption. Importantly, given that oral alcohol was provided by gavage to both the control and mutant mice, we were able to ensure that any genotype-specific effects we observed were not due to differences in alcohol consumption between groups. To this end, we also found that AgRP deficiency not only diminished the effects of alcohol on hepatic TAG content but also on hepatic cholesterol, although potentially to a lesser extent. In this regard, our findings align with other data suggesting that alcohol drinking impacts both TAG and cholesterol metabolism (8, 14, 18) and that fatty acid and cholesterol synthesis are often coordinately regulated in the liver (19).

It is worth noting that RNAi-mediated knockdown of AgRP expression in adult mice led to a more pronounced reduction of hepatic TAG levels in the context of acute alcohol administration when compared with that seen in mice with germline AgRP-deficiency. This could in part be due to potential compensatory changes induced in Agrp−/− mice due to the absence of AgRP during early development. It could also be attributed to differences in genetic background, as RNAi-treated mice were on a purer C57BL/6J background, which is susceptible to alcohol-induced hepatic steatosis, whereas the Agrp−/− mice were only partially (5–6 generations) backcrossed onto the C57BL/6J background.

One potential caveat of examining the role of AgRP neurons in alcohol-induced liver fat accumulation involves our use of maltose dextrin as an “isocaloric” control agent for comparison against alcohol. It has been recently shown that activation of hypothalamic AgRP neurons increases carbohydrate utilization while decreasing fat utilization (7). As such, maltose dextrin, a carbohydrate, could potentially have exerted independent metabolic effects on hepatic lipid through CNS mechanisms, including those potentially involving AgRP neurons. Although this was mitigated in our studies using 6-OHDA by examining a third, water-treated group of mice alongside those treated

Fig. 5. Chemical sympathectomy partially blocks the development of hepatic steatosis induced by bingelike alcohol consumption. Fifteen- to sixteen-week-old male C57BL/6J mice underwent chemical sympathectomy by injection with a single dose of 6-hydroxydopamine (6-OHDA; 250 mg/kg in 0.9% NaCl and 10⁻⁷ M ascorbic acid, Sigma-Aldrich, St. Louis, MO) or vehicle control (0.9% NaCl and 10⁻⁷ M ascorbic acid). After recovery, weight-matched mice were given a single oral gavage of water, maltose dextrin (9 g/kg body wt), or alcohol (5 g/kg body wt). Food was subsequently removed, and liver tissues were harvested 8 h later. 6-OHDA treatment affected body weight and blood glucose equally in water, maltose dextrin, or alcohol-gavaged mice (A–F). Sympathectomy by 6-OHDA did not affect total liver glycerol content [triacylglycerol (TAG)–derived and free glycerol] in water- or maltose dextrin-gavaged mice but caused a reduction in alcohol-gavaged mice (G–I). Similar results were seen with extracted and thin layer chromatography (TLC)-resolved TAG from livers (J and K).

Data are means ± SD. *P < 0.05; **P < 0.01; ns, nonsignificant, by two-tailed Student’s t test.
with maltose dextrin and alcohol, respectively, future studies may want to utilize isocaloric controls that employ either a noncarbohydrate macronutrient as an additional control or multiple controls employing different carbohydrates. It is technically challenging, however, to develop a variety of controls with similar caloric density to alcohol, to ensure that the volumes of the various gavage agents would be comparable.

The mechanisms by which alcohol regulates AgRP neuronal function remain to be determined. Alcohol exposure rapidly increases hypothalamic AgRP immunoactivity in C57BL/6J mice (12). Alcohol also increases the activity of AgRP neurons in hypothalamic brain slices (5), suggesting that alcohol may act directly in the brain to regulate AgRP neurons, rather than only doing so indirectly through a peripheral mechanism. Acute alcohol exposure inhibits the activity of equilibrative nucleoside transporter 1 (ENT1) in the brain, leading to an elevation of extracellular adenosine levels (10). As adenosine is a potent neuromodulator (44) and mice lacking the adenosine receptors A₁ or A₂B are resistant to alcohol-induced hepatic steatosis (43), we evaluated if A₂B in the CNS is involved in development of alcohol-induced hepatic steatosis. Deleting Adora2b in the CNS abolished hypothalamic Adora2b mRNA expression, confirming that Adora2b is normally expressed in the hypothalamus. Interestingly, we show that abolishing A₂B signaling in the brain potentiates the ability of alcohol consumption to acutely increase liver fat in mice, which could be attributed to overcompensation by other adenosine receptors such as A₁ receptor. It should also be acknowledged that Tg.Nestin-Cre:A2B<sup>−/−</sup> mice would have been better controls, as the use of Tg.Nestin-Cre, A2B<sup>+/−</sup> as controls would presumably lead to an underestimate of the phenotypes.

By contrast, our results show that removing A₂B only in AgRP neurons does not alter alcohol-induced hepatic steatosis, suggesting that A₂B signaling in non-AgRP neurons are responsible for alcohol’s regulatory effects in this context. For example, alcohol also targets POMC neurons (9, 25, 34, 42). Adolescent bingelike alcohol consumption or chronic consumption of alcohol-containing diet reduces hypothalamic α-MSH immunoactivity (25, 34). The opposing regulatory effects of alcohol on POMC and AgRP neurons support the notion that these neurons are both CNS targets of alcohol, perhaps contributing to alcohol’s metabolic effects in a reciprocal or otherwise integrated manner. Indeed, it is possible that adenosine signaling in POMC neurons, and potentially others, may compensate for the loss of A₂B signaling solely in AgRP neurons.

One possible mechanism by which AgRP regulates alcohol-induced hepatic steatosis is via modulating the autonomic nervous system innervating the liver and the adipose tissues. AgRP inhibits sympathetic activity in the white adipose tissue to limit lipolysis (38). We have also shown that intracerebroventricular infusion of AgRP suppresses sympathetic activity in the liver (29). Moreover, AgRP neurons mediate the stimulatory effects of leptin on sympathetic activity in the adipose tissue and liver (2). Thus diminished AgRP function in the brain likely disrupts autonomic function in liver and white adipose tissues, lessening their response to alcohol-induced adipose tissue lipolysis as well as hepatic lipid synthesis. Although our findings suggest that chemical sympathectomy abrogated the ability of alcohol consumption to increase liver fat content by disrupting the mechanism by which AgRP neurons act to stimulate hepatic TAG accumulation, we cannot rule out the possibility that ablatting the SNS could also have reduced liver fat content in the context of acute alcohol consumption by other indirect mechanisms.

Taken together, our study indicates that the brain not only underlies the effects of alcohol on cognitive, psychological, and locomotor functions but also mediates the metabolic effects of alcohol, specifically the acute development of hepatic steatosis induced by bingeing drinking. Our results further suggest that alcohol exerts its acute pathological effects, at least in part, by impinging on the central melanocortin system, which in addition to functioning as a key regulator of energy balance, is emerging as an important regulator of peripheral lipid metabolism.

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**DISCLOSURES**

No conflicts of interest, financial or otherwise, are declared by the authors.

**AUTHOR CONTRIBUTIONS**


**REFERENCES**


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