Enzymatic Responses to Alcohol and Tobacco Nicotine-Derived Nitrosamine Ketone Exposures in Long Evans Rat Livers

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Abstract

Background: Chronic feeding plus binge administration of ethanol causes very high blood alcohol concentrations. However, its co-administration with tobacco Nicotine-Derived Nitrosamine Ketone (NNK) results in somewhat lower blood alcohol levels, suggesting that NNK and therefore smoking, alters alcohol metabolism in the liver. To explore this hypothesis, we examined effects of ethanol and/or NNK exposures on the expression and activity levels of enzymes that regulate their metabolism in liver.

Methods: This study utilized a 4-way model in which Long Evans rats were fed liquid diets containing 0% or 26% ethanol for 8 weeks, and respectively i.p. injected with saline or 2 g/kg of ethanol 3 times/week during Weeks 7 and 8. The control and ethanol-exposed groups were each sub-divided and further i.p. treated with 2 mg/kg of NNK or saline (3x/week) in Weeks 3–8. ADH, catalase and ALDH activities were measured using commercial kits. CYP450 mRNA levels (17 isoforms) were measured by qRT-PCR analysis.

Results: Ethanol significantly increased hepatic ADH but not catalase or ALDH activity. NNK had no effect on ADH, ALDH, or catalase, but when combined with ethanol, it increased ADH activity above the levels measured in all other groups. Ethanol increased CYP2C7, while NNK increased CYP2B1 and CYP4A1mRNA levels relative to control. In contrast, dual ethanol + NNK exposures inhibited CYP2B1 and CYP4A1 expression relative to NNK.

Conclusion: Dual exposures to ethanol and NNK increase hepatic ethanol metabolism, and ethanol and/or NNK exposures alter the expression of CYP450 isoforms that are utilized in NNK and fatty acid metabolism.

Keywords: Alcoholic liver disease; Ethanol metabolism; NNK; Alcohol dehydrogenase; Aldehyde dehydrogenase; Cytochrome P450

Introduction

Severity of alcohol induced liver injury correlates with dose and duration of alcohol exposure and the levels of its toxic metabolites [1,2]. Acute alcohol-mediated liver injury is characterized by reversible steatosis. High levels of chronic or binge alcohol exposures cause simple steatosis to progress through stages including steatohepatitis, fibrosis, and finally cirrhosis [3]. An important mediator of alcohol-induced liver injury is the accumulation of its highly toxic metabolite, acetaldehyde [1]. However, one relatively recent consideration regarding the pathogenesis of progressive alcohol-mediated liver injury is that a very high percentage of heavy drinkers also smoke tobacco products (mainly cigarettes) [4]. Of note is that chronic exposures to ethanol, tobacco smoke/products, and tobacco specific nitrosamine, 4-(methylnitrosamino)-1-(3-pyridyl)-1-butane (NNK) all cause steatohepatitis with oxidative damage [5-9]. In our efforts to examine the role of NNK as a co-factor in alcoholic liver disease, we made the unexpected observation that dual exposures to NNK and chronic plus binge ethanol administration results in lower blood alcohol concentrations compared with the same ethanol treatments alone [8]. These observations suggested that hepatic enzyme activities needed to metabolize alcohol could be modulated by NNK exposures and therefore probably smoking.

Ethanol is metabolized in the liver through three distinct oxidative pathways including Alcohol Dehydrogenase (ADH), Cytochrome P450 2E1 (CYP2E1), and catalase (Figure 1). ADH is main enzyme responsible for the metabolism of ethanol. ADH is an abundantly expressed NAD+ dependent cytosolic enzyme [10]. The CYP2E1 microsomal ethanol oxidizing pathway requires NADPH as a cofactor [11]. Catalase, localized in peroxisomes, utilizes hydrogen peroxide to catalyze alcohol metabolism [12]. Acetaldehyde, a highly toxic and carcinogenic metabolite is generated by all three
oxidative pathways; acetaldehyde can covalently bind and form adducts with DNA [13], phospholipids [14], hepatic microsomal proteins, hemoglobin [15], and erythrocyte membrane proteins [16,17]. Aldehyde Dehydrogenase (ALDH) detoxifies acetaldehyde to acetate in mitochondria [18]. Acetaldehyde accumulations caused by unbalanced increases in ADH or inhibition of ALDH activity lead to increased oxidative and ER stress and cellular injury [19,20]. ADH metabolism of ethanol also yields reduced NADH which promotes steatosis by stimulating fatty acid synthesis [20].

NNK is metabolized by reduction of its carbonyl group [21], oxidation of its pyridine nitrogen [22], α-hydroxylation of the methylene carbon adjacent to the N-nitroso nitrogen [23], and α-hydroxylation of methyl carbons adjacent to the N-nitroso nitrogen [24]. Although several CYP450 isoforms can catalyze the α-hydroxylation reactions, CYP2B1 is the dominant isoform expressed in rat liver, whereas CYP1A2, CYP2A1, and CYP3A have minor roles [25,26]. Metabolism of NNK via α-methyl or α-methylene hydroxylation produces reactive metabolites that can methylate [27] or pyridyloxobutylate DNA [28]. In addition, in NNK-associated experimental steatohepatitis, the high hepatic levels of O6 methyl-Guanine adducts may indicate that liver injury is consequential to NNK metabolism [8].

This study was prompted by the finding that dual exposures to ethanol and NNK produced additive adverse histopathological and molecular effects on liver, despite lower blood alcohol levels relative to ethanol-only exposures. One potential explanation for this phenomenon is that NNK increases the rate of ethanol metabolism, yielding higher levels of toxic metabolites and attendant exacerbation of liver injury. Herein, we examined the effects of ethanol, NNK and ethanol + NNK exposures on the mRNA expression or activity levels of enzymes that mediate ethanol and NNK metabolism.

Methods

Experimental model

Liver tissue samples were obtained from a previously generated model [8]. In brief, 4 week old Long Evans male rats were fed isocaloric liquid diets containing 0% or 26% ethanol for 8 weeks (n=12/group). Ethanol-fed rats were binged with 2 g/kg of ethanol by intraperitoneal (i.p.) injection, 3 times per week during the last two weeks of the experiment. Control rats were treated in parallel with i.p. saline. Half the rats in each group were also i.p. injected with NNK (2 mg/kg) or saline 3 times per week from Week 3 through Week 8. Snap-frozen liver tissue harvested at sacrifice and stored at -80°C was used in these studies. The rat experiments were conducted in accordance with the Guide for the Care and Use of Laboratory Animals as adopted and promulgated by the U.S. National Institutes of Health, and approved by the Lifespan Institutional Animal Care and Use Committee (IACUC).

Enzyme activity assays

ADH, ALDH, and catalase activities were measured using commercial kits (BioVision, California, USA). Results normalized to sample protein concentration which was measured with the Pierce TM Thermo Scientific, Rockford, IL) bicinchoninic acid assay. The ADH colorimetric assay is based on alcohol-to-aldehyde interconversion via reduction of NAD+ to NADH and uses isopropanol. ALDH assay is based on oxidation of acetaldehyde, yielding NADH. Changes in absorbance for both the ADH and ALDH assays were measured over 90 minutes at 450 nm. The 10-minute time point was used for
inter-group comparisons of enzymatic activity since linear reaction rates were detected within 10 minutes (Figures 2b and 2d). Catalase activity levels were based on the rates in which H₂O₂ was decomposed to water and oxygen using OxiRed™ to probe unconverted H₂O₂. Absorbances (570 nm) were inversely proportional to catalase activity.

Quantitative Reverse Transcriptase Polymerase Chain Reaction (qRT-PCR) Assays

The RNeasy Mini Kit (Qiagen, Hilden, Germany) was used to extract total RNA from 100 mg samples of fresh frozen liver. RNA was reverse transcribed using the AMV 1st Strand cDNA Synthesis Kit (Roche, Indianapolis, IL). The resulting cDNA templates were used to measure expression levels of 17 cytochrome P450 (CYP) isoforms (Supplementary Table 1) by qPCR in a Roche Lightcycler 480 System as described [29]. Primer pairs were designed using Mac Vector 12 software (Cary, NC). Relative mRNA abundance was calculated using the 2^ΔΔCt method with results normalized to Hypoxanthine-guanine phosphoribosyltransferase (HPRT) and Beta-actin (Actβ) as internal controls.

Statistical analysis

Results are expressed as mean ± SD. Inter-group comparisons were made by two-way Analysis Of Variance (ANOVA) with Tukey or linear trend post hoc tests (GraphPad Prism 7, San Diego, CA, USA). All assays were performed with 6 independent samples per group since preliminary studies and power analysis determined that this group size was sufficient to achieve 80% power. P-values less than 0.05 were considered as statistically significant.

Results

Effects of ethanol and NNK on alcohol metabolizing enzyme activity levels in liver

Two-way ANOVA tests demonstrated significant ethanol and NNK effects on ADH activity, and significant ethanol effects on ALDH activity (Table 1). In contrast, no significant ethanol, NNK or ethanol x NNK interactive effects were detected for Catalase activity.

ADH activity

Mean (± S.D.) hepatic ADH activity in control (31.7 ± 4.6 nmol/min/mg protein) and NNK-only (38.3 ± 3.5 nmol/min/mg) were similar. Ethanol significantly increased ADH activity (46.5 ± 11.3 nmol/min/mg) by 1.5-fold relative to control (P=0.04). The combined ethanol + NNK treatments increased ADH activity (59.7 ± 12.2 nmol/min/mg) by 1.9-fold relative to control (P=0.001) and 1.6-fold relative to NNK exposure (P=0.002) as demonstrated with Tukey post hoc tests (Figure 2a, Table 1). These findings suggest that ethanol increases the metabolic rate for conversion of ethanol to acetaldehyde, and dual ethanol + NNK exposures have additive effects on hepatic ADH activity in rats.

ALDH activity

The mean levels of hepatic ALDH activity in the control (53.2 ± 5.5 nmol/min/mg) and NNK-exposed (56.8 ± 5.7 nmol/min/mg)
Multiple comparisons post-hoc significance test outcomes are graphed in Figure 2. mRNA was measured by qPCR analysis. Corresponding data with Tukey ethanol + NNK treated Long Evans rats (n=6 rats/group). Gene expression cytochrome P450 genes expressed in control, ethanol-fed, NNK-exposed and ethanol + NNK treated Long Evans rats (n=6 rats/group). Two-Way ANOVA results from comparing hepatic mean levels of Alcohol dehydrogenase (ADH), aldehyde dehydrogenase (ALDH), and catalase activities in control, ethanol-fed, NNK-exposed and ethanol + NNK treated Long Evans rats (n=6 rats/group). Significant differences are highlighted with bold font. Corresponding data with Tukey post-hoc significance test outcomes are graphed in Figure 2.

<table>
<thead>
<tr>
<th>Enzyme activity</th>
<th>Ethanol Factor</th>
<th>NNK Factor</th>
<th>Ethanol x NNK Interaction</th>
</tr>
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<tr>
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<td>F-Ratio</td>
<td>P-Value</td>
<td>F-Ratio</td>
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<td>ALDH</td>
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<td>Catalase</td>
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Table 2: Ethanol, NNK, and Their Interactive Effects on Cytochrome P450 mRNA Expression. Two-Way ANOVA results from comparing hepatic mean levels of 17 different cytochrome P450 genes expressed in control, ethanol-fed, NNK-exposed and ethanol + NNK treated Long Evans rats (n=6 rats/group). Gene expression (mRNA) was measured by qPCR analysis. Corresponding data with Tukey multiple comparisons post-hoc significance test outcomes are graphed in Figure 3. Significant differences are highlighted with bold font.

<table>
<thead>
<tr>
<th>Gene</th>
<th>Ethanol Factor</th>
<th>NNK Factor</th>
<th>Ethanol x NNK Interaction</th>
</tr>
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<tbody>
<tr>
<td></td>
<td>F-Ratio</td>
<td>P-Value</td>
<td>F-Ratio</td>
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<tr>
<td>CYP1A1</td>
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<td>CYP1A2</td>
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<td>CYP4A1</td>
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Groups were similar, whereas the levels in the ethanol (64.1 ± 14.1 n mol/min/mg) and ethanol + NNK (71.0 ± 22.5 n mol/min/mg) were higher. Although significant ethanol effects were observed by two-way ANOVA (Table 2), post-hoc Tukey tests failed to reach statistical significance (Figure 2e, Table 1). Nonetheless, the results demonstrate that ethanol exposure has stimulatory effects on hepatic ADH activity whereas NNK does not. However, the magnitude of ethanol-induced increase in ADH activity was modest compared with ADH, suggesting that acetaldehyde metabolism lagged relative to its generation in all 3 experimental groups.

Catalase activity

Hepatic catalase activity, irrespective of treatment, was approximately 1000 times higher than ADH activity. The mean level of catalase activity in control livers (10.1 ± 4.2 µmol/min/mg) was similar to those measured in the ethanol (9.6 ± 3.5 µmol/min/mg), NNK (11.4 ± 4.4 µmol/min/mg), and ethanol + NNK (9.8 ± 3.1 µmol/min/mg) groups, corresponding with results obtained by two-way ANOVA (Figure 2e, Table 1).

Effects of ethanol and tobacco nitrosamine NNK on hepatic cytochrome P450 expression

Cytochrome P450 (CYP) genes that were measured regulate metabolism of hepatic steroid hormones (CYP1A1, CYP1A2, CYP2A1, CYP2A2, CYP2A11, CYP2C12, CYP2C13, CYP2D1, CYP2E1, CYP3A2, and CYP3A23), fatty acids (CYP1A1, CYP1A2, CYP2B1, CYP2E1, and CYP4A1), and retinoic acid (CYP2C7) and xenobiotics (CYP1A1, CYP1A2, CYP2A1, CYP2A2, CYP2C6, CYP2D2, and CYP3A2) or bioactivation of carcinogens (CYP2B1, CYP2B2, and CYP2E1) (Supplementary Table 1). CYP mRNA levels were measured by qRT-PCR analysis.

Two-way ANOVA tests demonstrated significant ethanol effects on the expression levels of CYP2B1, CYP2C6v1, CYP2C7, CYP2C11, CYP2E1, and CYP3A23; no significant ethanol x NNK effects on the expression levels of CYP2B1, CYP2C7, CYP2D1, CYP2D2, and CYP4A1; and no significant NNK effects (Table 2). Post hoc tests demonstrated that ethanol significantly increased expression of CYP2C7 (1.8 fold; P < 0.03) and modestly, but not significantly decreased expression of CYP2C11, CYP2E1, and CYP3A23 relative to control.

NNK increased CYP2B1 expression relative to control (1.9-fold; P = 0.06), ethanol (2.4-fold; P < 0.03), and ethanol + NNK (4.8-fold; P = 0.005) (Figure 3e). In addition, NNK significantly increased CYP4A1 expression relative to control (2.5-fold; P < 0.03) and ethanol + NNK effects (2.5-fold; P = 0.04), and had a trend increase relative to ethanol-only effects (1.9-fold; P = 0.08) (Figure 3q). Another notable point was that the mean levels of CYP2B1 (Figure 3n) and CYP3A23 (Figure 3p) were similar in control and NNK livers, whereas ethanol and ethanol + NNK had relative inhibitory effects on the expression of those genes (both P = 0.08). Like ethanol, ethanol + NNK exposures inhibited expression of CYP2E1 and CYP3A23 (1.6-fold; P = 0.04) relative to control (Figure 2p). In addition, CYP3A23 was also reduced by ethanol + NNK relative to NNK only exposures (1.8-fold; P = 0.01).

Discussion

This study examined the effects of chronic plus binge ethanol, NNK, and ethanol + NNK exposures on the activity or mRNA levels of major enzymes that regulate alcohol and NNK metabolism in liver using a 4-way experimental paradigm generated in adult male Long Evans rats. The research was driven by the earlier observation that dual ethanol + NNK exposures resulted in lower steady-state and post-binge blood alcohol concentrations compared with the
same ethanol-only exposures [8]. This finding led to the hypothesis that alcohol metabolizing enzymes were modulated by ethanol and/or NNK exposures. This concept is consistent with previously published data demonstrating that experimental chronic ethanol administration increases mRNA, protein, and activity levels of ADH in rat livers [19,30]. Similarly, in other studies, in rats that were infused continuously with ethanol-containing diets, blood and urine ethanol concentrations cycled between 0 and 500 mg/dL [31], but after the urine concentrations reached 300 mg/dL, ADH mRNA expression and activity were induced [32], consistent with adaptive responses to alcohol load. Our studies extend these earlier finding by measuring activity and expression of a broader range of ethanol-metabolizing enzymes, and also consider the effects of independent and co-exposures to NNK. The importance of the NNK parallel studies is that consideration is given to the co-factor role of tobacco exposures/smoking in the pathogenesis of alcohol-mediated liver injury.

In this study, we demonstrated that ethanol, ethanol + NNK significantly increase ADH but just modestly increase ALDH relative to control, with larger effect sizes produced by the dual exposures. Although NNK alone had no detectable effect on ADH or ALDH activity relative to control, the finding that ethanol + NNK had somewhat additive responses compared with ethanol-alone suggests that NNK, and therefore probably also smoking, has modifying effects on ethanol-stimulated ADH and ALDH activities. The greater increase in ADH activity associated with ethanol + NNK versus ethanol-only exposures accounts for the lower blood alcohol concentrations measured in the dual-exposure group. On the other hand, the disproportionately greater increases in ADH compared with ALDH activity effected by both ethanol and ethanol + NNK favored excess formation and accumulation of acetaldehyde in liver. In addition, the larger effects of ethanol + NNK versus ethanol suggest that hepatocacetaldehyde levels were higher in the dual exposure group, an effect that could account for the greater severity of liver injury compared with the ethanol-only group.

Mechanistically, acetaldehyde’s highly toxic and carcinogenic [17] effects are due to its strong electrophilic structure and ability to form adducts with nucleophilic cellular components (DNA, RNA, and protein), leading to diverse pathophysiological responses including enzyme inactivation, DNA damage, and increased cell death [33]. Impairment of cellular enzyme functions leads to increased formation of reactive metabolites and free radicals that promote oxidative stress, lipid peroxidation, and membrane damage, exacerbating tissue injury and organ dysfunction. Therefore, detoxification of acetaldehyde via ALDH activation is a vital cellular protective mechanism that has therapeutic implications [33,34]. In regard to the present work, it is likely that acetaldehyde accumulations mediated by the minimal or absent activation of ALDH vis-à-vis significant increases in ADH activity significantly contributed to liver injury following ethanol and/or NNK exposures.

Among the 17 CYP isoforms included in the qPCR array, the expression levels of 9 (53%) were significantly modulated by ethanol (n=6) and/or ethanol + NNK (n=5) exposures. Just two of those CYP genes were significantly altered by both ethanol and ethanol + NNK, and NNK-only effects were not observed. These findings suggest that NNK and therefore smoking, differentially modifies hepatic profiles of CYP gene expression in the context of chronic binge ethanol...
NNK is primarily oxidized by CYP2B1 [35], and less commonly by CYP1A2 and CYP2A1 in rat livers [25,26]. Induction of CYP2B1 activates NNK via α-methyl or α-methylene hydroxylation pathways. Hydroxylated NNK metabolites can form methyl or pyridyloxobutyl adducts with DNA [27,28]. Therefore, the NNK associated increases in CYP2B1 expression may have contributed to liver injury due to accumulation of DNA adducts and attendant DNA damage.

Previous studies showed that ethanol induces hepatic CYP2E1 protein levels and enzyme activity [36], and it impairs mitochondrial β-oxidation but stimulates ω-oxidation of free fatty acids in reactions catalyzed by microsomal CYPs, including mainly by CYP2A1 and CYP2E1 [37]. However, in the present study, ethanol and ethanol + NNK exposures resulted in modest reductions in CYP2E1, and no change in CYP4A1 mRNA levels. Discrepancies between mRNA and protein expression or enzyme activity occur under various conditions. One potential explanation for the reduced mRNA levels is adduct formation mediated by metabolites such as acetaldehyde. In contrast, NNK exposed rat livers had significant or trend elevations in CYP4A1 expression relative to the control, ethanol and ethanol + NNK groups. Increased levels of ω-oxidation of long chain fatty acids produce toxic Dicarboxylic Acids (DCAs) and ω-hydroxylated fatty acids, which impair mitochondrial oxidative phosphorylation and promote steatohepatitis [37,38]. Therefore, NNK’s selective stimulatory effects on CYP4A1 may have been an important mediator of steatohepatitis in that group.

CYP2C7, a constitutive enzyme involved in the oxidative metabolism of vitamin A (retinol) and retinoic acid [39,40], was previously shown to be stimulated by ethanol in rat liver microsomes [36]. Correspondingly, hepatic CYP2C7 expression was also significantly increased by ethanol and exhibited statistical trend elevations in the ethanol + NNK group. A potential consequence of CYP2C7 induction by ethanol is that with the increased catabolism of retinoic acid and attendant elimination of retinol (Vitamin A) from the body, hepatic stores of vitamin A could be rendered deficient, which occurs commonly in alcoholicics [41,42].

In conclusion, the findings of this study suggest that chronic alcohol and NNK exposures significantly alter hepatic expression, activation and profiles of enzymes used to metabolize ethanol and/or NNK. These effects of ethanol and/or NNK may contribute to the pathogenesis of steatohepatitis by increasing hepatic accumulation of ethanol and NNK mediated DNA adducts. These data provide a better understanding of the mechanisms of alcohol-related liver injury and the contribution of tobacco NNK or smoking as a cofactor in the pathogenesis of alcoholic liver disease.

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