Alcohol-associated health problems, including alcoholic liver disease (ALD), are major global health problems. ALD progresses through the course of several pathologies, including steatosis, alcoholic hepatitis, cirrhosis, and potentially hepatocellular carcinoma. Alcoholic hepatitis occurs in approximately 10% to 35% of chronic heavy drinkers, and severe alcoholic hepatitis accounts for significant morbidity and mortality, approaching 35% to 45%.1 Approximately 10% to 20% of heavy drinkers will eventually develop cirrhosis.2,3 The specific mechanisms responsible for ALD development and progression are not fully understood, and there is no Federal Drug Administration—approved therapy for any stage of ALD.

Supported by NIH grants R21 AA020849-01A1 (L.A.K.), R01 AA024102-01A1 (L.A.K.), U01AA022489 (A.E.F. and C.J.M.), 1U01AA021901-01 (C.J.M.), 1U01AA021893-01 (C.J.M.), and R01AA023681 (C.J.M.); the Department of Veterans Affairs grant I01BX000350 (C.J.M.); the Department of Defense grant W81XWH-11-1-0595 (C.J.M.); the National Institute on Alcohol Abuse and Alcoholism and National Institute on Aging Intramural Programs (C.E.R.); the National Institute of General Medical Sciences of the NIH Institutional Development Award under grant P20GM113226 (C.J.M.), 1U01AA024337 (C.J.M.); and the National Center for Research Resources, NIH grant S10RR027926 (Wayne State Lipidomics Core Facility).

D.R.W. and H.L. contributed equally to this work.

Disclosures: None declared.
The link between diet, specifically dietary fat, and alcohol consumption in ALD pathogenesis has been increasingly appreciated. Findings from an epidemiological study that analyzed dietary fat intake in individuals from countries with similar per capita alcohol consumption revealed that intake of saturated fat (SF) was associated with lower mortality rates, whereas dietary intake of unsaturated fat (USF) was associated with a higher mortality from alcoholic cirrhosis. The beneficial effects of dietary SF (predominantly rich in medium- or long-chain saturated fatty acids) and the damaging effects of dietary USF [primarily enriched in linoleic acid (LA), an ω-6 polyunsaturated fatty acid (PUFA)] on alcohol-induced liver injury have been demonstrated in numerous preclinical studies using rodent models of ALD. LA is required for the development of experimental ALD, with the severity of liver pathology correlating to the amount of LA in the diet. On a mechanistic level, the deleterious effects of USF in comparison to the protective effects of SF are thought to result from alterations in hepatic lipid homeostasis, induction of hepatic lipid peroxidation and oxidative stress, changes in the gut microbiota, impaired intestinal barrier integrity, and elevated endotoxemia with subsequent hepatic macrophage activation and increased production of hepatic proinflammatory cytokines. LA is the most abundant ω-6 PUFA in human diets and in human plasma and membrane lipids. The consumption of LA, which has dramatically increased during the past several decades, is positively correlated with increasing prevalence of several pathological conditions, including obesity. LA can be enzymatically converted to bioactive oxidized LA metabolites (OXLAMs), primarily via the actions of cellular lipooxygenases [arachidonate 15-lipoxygenase (ALOX15) and ALOX15B in humans, and ALOX15 (12/15-LO) in rodents] or nonenzymatically via free radical–mediated oxidation in response to oxidative stress. OXLAMs are involved in various intracellular signaling pathways and may induce a proinflammatory response in different cell types. OXLAMs play a role in the development of different pathological conditions, such as inflammatory hyperalgesia and the metabolic syndrome. OXLAMs, specifically 9- and 13-hydroxyoctadecadienoic acids (9- and 13-HODEs, respectively), were elevated in patients with nonalcoholic fatty liver disease, and a decrease in plasma OXLAM levels was correlated with hepatic histological improvement in these patients. Increased levels of 9- and 13-HODEs were found in patients with alcoholic liver cirrhosis in parallel with the induction of hepatic ALOX15 and ALOX15B. Elevated 9- and 13-HODE levels were also observed in experimental rodent models of ALD and were associated with ethanol-induced liver injury, steatosis, and inflammation. Based on these observations, we hypothesized that OXLAMs may play a pathogenic role in ALD development and/or progression; however, the underlying mechanisms remain to be determined.

In the present study, we tested the hypothesis that the combination of ethanol and a diet with a high content of LA contributes to and exacerabtes ethanol-induced liver injury through increased OXLAM production and OXLAM-mediated induction of hepatic proinflammatory responses. We postulated that both increased substrate availability (LA-rich dietary fat) and activation of metabolic pathways of OXLAM production, in particular the 12/15-LO–mediated pathway, contribute to an ethanol-mediated increase in OXLAMs. Therefore, we further examined whether genetic ablation of Alox15 (the gene encoding 12/15-LO) would decrease OXLAM levels and attenuate liver injury caused by ethanol and an LA-enriched diet. For our studies, we used a chronic-binge ethanol exposure animal model of ALD (NIH National Institute on Alcohol Abuse and Alcoholism model). This model recapitulates the drinking pattern in humans and induces a moderately severe alcoholic liver injury with increased hepatic neutrophil infiltration, which is often observed in alcoholic hepatitis patients.

**Materials and Methods**

**Animals and Treatments**

Eight-week-old genetically unaltered wild-type (WT; C57BL/6J) and Alox15 gene knockout (B6.129S2-Alox15tm11Fun/J, 11th backcross generation) male mice were obtained from the Jackson Laboratory (Bar Harbor, ME). Animals were housed in a specific pathogen-free barrier facility accredited by the Association for Assessment and Accreditation of Laboratory Animal Care. A 10-day ethanol feeding plus a single binge of ethanol administration was used as an experimental model of ALD (NIH National Institute on Alcohol Abuse and Alcoholism model). Briefly, the mice were provided free access to a Lieber-DeCarli control or ethanol (5% w/v)-containing diet for 10 days. On day 11, the mice were gavaged with a single dose of ethanol solution (20% w/v) prepared in ultrapure water; the gavage volume was calculated to deliver 5 g ethanol/kg body weight. The mice in control groups received an isocaloric/isovolumetric maltodextrin solution. The mice were euthanized 9 hours after the gavage. The experimental paradigm is outlined in Figure 1A. Two separate experiments were performed: WT mice were provided a USF- or SF-enriched diet with or without ethanol; and WT and Alox15 knockout mice were fed a USF-enriched diet with or without ethanol. The USF diet was supplemented with corn oil, a rich source of LA, and the SF diet was supplemented with beef tallow and medium-chain triglycerides (Research Diets, New Brunswick, NJ) (Table 1). The detailed composition of these diets has been described previously. In the control group, the levels of protein, carbohydrate, and fat were held constant at 17%, 43%, and 40% of total energy, respectively. In the alcohol-consuming diets, ethanol (35% of total calories) was substituted for carbohydrate energy. The diets were prepared fresh each day, and the food consumption was monitored daily. The control groups

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(the pair-fed mice) received the same amount of isocaloric food (maltodextrin-containing diets) that ethanol-fed animals consumed in the previous day (pair-feeding protocol). At the conclusion of the experiment, the mice were euthanized and portions of liver tissue were snap frozen in liquid nitrogen or were fixed in 10% neutral-buffered formalin. Blood was collected from the inferior vena cava via venipuncture using heparinized syringes. Plasma was obtained, aliquoted, and stored at −80°C for further analysis. The study protocol was approved by the University of Louisville Institutional Animal Care and Use Committee (Louisville, KY).

Biochemical Analysis and Enzyme-Linked Immunosorbent Assays

Plasma alanine transaminase (ALT; a marker of liver damage) activities were measured using the Piccolo Xpress chemistry analyzer (Abaxis, Union City, CA). Hepatic lipids were extracted with chloroform and methanol, and triglycerides were measured using Triglycerides Reagent (Thermo Fisher Scientific Inc., Middletown, CA). Liver thiobarbituric acid reactive substances (TBARSs) were measured in whole-liver homogenate using a commercially available kit (Cayman Chemical, Ann Arbor, MI). Plasma lipopolysaccharide (LPS) levels were measured with the Limulus Amoebocyte Lysate kit (Lonza, Walkersville, MD). Soluble CD14 (R&D Systems, Minneapolis, MN) and

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Nutritional Composition of the SF and USF Experimental Diets</th>
</tr>
</thead>
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<tr>
<td>Ingredients</td>
<td>SF diet</td>
</tr>
<tr>
<td>Protein, kcal%</td>
<td>17</td>
</tr>
<tr>
<td>Carbohydrate, kcal%</td>
<td>43</td>
</tr>
<tr>
<td>Fat, kcal%</td>
<td>40</td>
</tr>
<tr>
<td>Saturated FA, %</td>
<td>84.1</td>
</tr>
<tr>
<td>Monounsaturated FA, %</td>
<td>9.3</td>
</tr>
<tr>
<td>Polyunsaturated FA, %</td>
<td>6.6</td>
</tr>
</tbody>
</table>

FA, fatty acid; SF, saturated fat; USF, unsaturated fat.
lipopolysaccharide binding protein (Hyct Biotech, Plymouth Meeting, PA) were measured by enzyme-linked immunosorbent assays, according to the manufacturers’ instructions.

Liver Histological Analysis and Staining

Mouse liver tissues were collected, fixed in 10% neutral-buffered formalin solution, and embedded in paraffin. Hematoxylin and eosin staining was performed for liver tissue histological evaluation. Oil-Red-O staining on liver tissue cryostat sections was performed to evaluate hepatic fat accumulation. Apoptotic cells were identified by terminal deoxynucleotidyl transferase dUTP nick-end labeling assay using the ApopTag Peroxidase In Situ Apoptosis Detection kit (Millipore, Billerica, MA). Hepatic neutrophil accumulation was assessed by chloroacetate esterase staining using a commercially available kit (Sigma, St. Louis, MO). Hepatic 4-hydroxy-2-nonenal levels were detected by immunohistochemistry using antibodies from Alpha Diagnostic International (San Antonio, TX).

RNA Isolation and Real-Time RT-PCR Assay

Total RNA was purified using Trizol reagent (ThermoFisher, Waltham, MA), as described by the manufacturer. To ensure there was no genomic DNA carryover, samples were digested with DNase I (ThermoFisher). cDNA was synthesized using qScript cDNA Supermix (Quanta Biosciences, Beverly, MA). The cDNA equivalent of 10 ng RNA was used in each real-time quantitative PCR (with the exception of 18S ribosomal RNA, where 0.1 ng was used). RT-PCR assays were performed with PerfeCTa SYBR Green Fast Mix (Quanta Biosciences) on the Applied Biosystems 7900HT platform (Foster City, CA) using primers listed in Table 2. All data were normalized to the expression of the 18S ribosomal RNA gene, and the data were calculated as described previously.35

Western Blotting

Western blotting was performed to evaluate the levels of cytochrome p450 2E1 (CYP2E1) using a commercially available antibody (Abcam, Cambridge, MA). Liver tissue lysates were separated by SDS-PAGE and subsequently transferred to a polyvinylidene difluoride membrane. Blots were visualized using Amersham enhanced chemiluminescence Western blot detection (GE Health Care, Hercules, CA), and quantified using ImageJ software version 1.50i (NIH, Bethesda, MD; http://imagej.nih.gov/ij). Glyceraldehyde-3-phosphate dehydrogenase (Santa Cruz Biotechnology, Dallas, TX) was used as a loading control.

Liver Lipidomic Analysis

Lipid extraction from liver tissue and quantification of lipid metabolite levels were performed, as previously described,36-39 by the Wayne State University Lipidomic Core Facility (Detroit, MI). Briefly, tissue samples were homogenized in phosphate buffer, pH 7.2. Samples were spiked with internal standards and extracted for PUFA metabolites on C18 extraction columns, as described earlier.36-39 High-performance liquid chromatography was performed on a Prominence XR system (Shimadzu Corp., Kyoto, Japan) using a Luna C18 (3 μm, 2.1 × 150 mm) column. Mass spectra for each detected lipid mediator were recorded using the Enhanced Product Ion feature to verify the identity of the detected peak in addition to multiple reaction monitoring transition and retention time match with the standard. The data were collected using Analyst 1.6.2 software (SCIEX, Framingham, MA), and the multiple reaction monitoring transition chromatograms were quantitated by MultiQuant software version 3.0.2 (SCIEX). The internal standard signals in each chromatogram were used for normalization for recovery as well as relative quantitation of each analyte.

Plasma Oxidized Linoleic Acid Metabolites (OXLAMs) Measurement

Lipid extraction from plasma and quantification of 9- and 13-HODE and LA levels were performed as previously described.26,40 Briefly, plasma samples with antioxidant solution, internal standards [synthetic 9(S)-HODE-d4 or 13(S)-HODE-d4], and potassium hydroxide were added to glass test tubes and overlaid with argon. After hydrolysis under argon atmosphere, the released fatty acids were extracted twice into the hexane layer by liquid/liquid extraction. The combined hexane extracts were dried under nitrogen gas and resuspended in 85% methanol.

Table 2 Murine Primer Sequences for RT-PCR

<table>
<thead>
<tr>
<th>Gene</th>
<th>Forward primer</th>
<th>Reverse primer</th>
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<tbody>
<tr>
<td>18s</td>
<td>5’-CTCAACAGGGAAACCTGCAC-3’</td>
<td>5’-CGCTCCACCAAATAGAAGC-3’</td>
</tr>
<tr>
<td>Tnfa</td>
<td>5’-CTGATCGTGTCCCTCAAAGC-3’</td>
<td>5’-GGTGCTTTTGCTACAGTGC-3’</td>
</tr>
<tr>
<td>Mip2a</td>
<td>5’-GCCGCCCAGCAGAAGTGCATA-3’</td>
<td>5’-TCCAGTGTCAGTTACCTAGC-3’</td>
</tr>
<tr>
<td>Mcp1</td>
<td>5’-GGCTCGACGAGATGCAGT-3’</td>
<td>5’-TGAGCGGCTGAGAAAAAATAGCAG-3’</td>
</tr>
<tr>
<td>Il1b</td>
<td>5’-TTATCCTTGAAGAAAGGCCCCAT-3’</td>
<td>5’-TCCGAGCCTGTAAGCAGTTP-3’</td>
</tr>
<tr>
<td>Nos2</td>
<td>5’-GAGCAGAGGAGGATCCAGGAC-3’</td>
<td>5’-CCACGAGTAGTTGCTCTTTC-3’</td>
</tr>
<tr>
<td>Arg1</td>
<td>5’-CATGGGGCTGCGACGAGGAG-3’</td>
<td>5’-GCTGAAATGCTCTTCCATCA-3’</td>
</tr>
<tr>
<td>Tgfb1</td>
<td>5’-TGATACGCCTGAGTTGCTCTCT-3’</td>
<td>5’-CAAGAGCAGTGGACGCTAA-3’</td>
</tr>
</tbody>
</table>
methanol/water. Reconstituted lipid extracts were analyzed by high-performance liquid chromatography. Oxidized fatty acids were quantified on a triple quadrupole mass spectrometer (API 365; Applied Biosystems, Foster City, CA) with Ionics EP 10þ upgrade (Concord, ON, Canada) using the stable isotope dilution method and multiple reaction monitoring with characteristic parent-to-daughter ion transitions.

Cell Culture and Treatments

RAW264.7 cells (a mouse monocyte/macrophage cell line) were obtained from ATCC (Manassas, VA) and cultured in Dulbecco’s modified Eagle’s medium supplemented with 10% fetal bovine serum and 100 U/mL of penicillin and streptomycin at 37°C and 5% CO2. The 9- and 13-HODEs (supplied in 100% ethanol) were purchased from Cayman Chemical. Cells were seeded at a density of 2×105 cells/cm² and incubated with 5 µmol/L 9- or 13-HODE (in the presence of 25 mmol/L ethanol) or vehicle/control (25 mmol/L ethanol) for 24 hours. LPS (from Escherichia coli 0111:B4; Sigma Chemical Co, St. Louis, MO) was then added to a final concentration of 100 ng/mL, and cultures were continued for an additional 3 hours. Cells were washed twice with sterile phosphate-buffered saline and scraped in Trizol (Thermo-Fisher), and total RNA was purified as described by the manufacturer. The HODE concentration was chosen on the basis of the observation that the circulating HODE levels could reach low micromolar range in humans with ALD.28

Five micromolar 9- or 13-HODE is the maximum concentration that did not cause significant cell death in RAW264.7 cells, as determined in our preliminary experiments.

Statistical Analysis

The data are expressed as the means ± SEM. For animal studies, statistical significance was determined using either the two-tailed t-test or two-way analysis of variance, followed by the Bonferroni multiple comparison test, where appropriate. For cell culture studies, the data were analyzed by one-way analysis of variance. P < 0.05 was considered statistically significant. Statistical analysis was performed using GraphPad Prism version 5.01 for Windows (GraphPad Software, Inc., La Jolla, CA).

Results

Liver Injury Caused by a Diet Rich in Linoleic Acid and Chronic-Binge—Ethanol Administration Is Associated with Elevated OXLAM Levels

A mouse model of chronic-binge—ethanol-induced liver injury was used in this study. The animals were fed a Lieber-deCarli control or ethanol-containing diet enriched in SF or USF. We first evaluated the effects of different types of dietary fat in combination with ethanol exposure on body/organ weights (Table 3). Body weights were significantly reduced in response to ethanol administration, regardless of the type of lipids in the diet. When compared to pair-fed mice, those receiving SF+ethanol had a significant increase in liver weights, whereas USF+ethanol-fed mice had no appreciable differences in liver weights. When expressed in terms of body weight, the liver/body weight ratios were similarly increased in both the SF+ethanol and USF-ethanol groups. Ethanol administration significantly reduced epididymal white adipose tissue as well as epididymal white adipose tissue/body weight ratios regardless of the type of dietary fat. There were no differences in body/organ weights between SF and USF or between SF+ethanol- and USF+ethanol-fed mice.

As a measure of liver injury, we assayed plasma ALT activity and found that it was significantly elevated in the USF+ethanol group when compared to pair-fed controls and to mice fed SF+ethanol (Figure 1B), suggesting a
Effect of Saturated or Unsaturated Dietary Fat on Chronic-Binge—Ethanol-Induced Hepatic Oxidative Stress

Oxidative stress, one of the factors contributing to OXLAM production, is a common feature of ethanol-induced liver damage and plays an important role in ALD pathogenesis. To determine whether the type of dietary fat plays a role in modulating the degree of chronic-binge—ethanol-induced oxidative stress, hepatic levels of CYP2E1, 4-hydroxy-2-nonenal, and TBARS were evaluated as surrogate markers of oxidative stress. Consistent with previous reports in different animal models of ALD, hepatic levels of CYP2E1 (the major enzyme for ethanol metabolism) were similarly elevated in SF- and USF-fed mice in response to chronic-binge ethanol administration (Figure 2, A and B). A trend toward elevated intensity of 4-hydroxy-2-nonenal—positive staining was observed in the USF and ethanol-fed mice, and these mice exhibited more liver injury compared to SF + ethanol-fed animals.

In the present study, we observed significant elevations in both liver (Figure 1F) and plasma (Figure 1G) OXLAM levels in USF + ethanol-fed mice, and these mice exhibited more liver injury compared to SF + ethanol-fed animals.
A Linoleic Acid—Enriched Diet Enhances Hepatic Inflammation in Response to Chronic-Binge Ethanol Exposure

The current chronic-binge ethanol feeding model is known to cause robust hepatic neutrophil infiltration. In our study, we observed significantly increased numbers of neutrophils in the livers of mice fed USF+ethanol compared to pair-fed controls and to mice fed SF+ethanol (Figure 3, A–C). There were no differences in liver neutrophil infiltration between mice fed SF and USF diets without alcohol. Expression of key macrophage M1 type (proinflammatory) response genes, including tumor necrosis factor-α (TNF-α) and IL-1β, was significantly increased in response to the USF+ethanol diet compared to pair-fed controls, an effect not observed in SF+ethanol-fed mice (Figure 3, D and E). Of note, hepatic mRNA levels of monocyte chemotactic protein 1 (MCP-1), a chemotactic factor for monocytes/macrophages, were significantly increased in response to ethanol regardless of the type of dietary fat (Figure 3F). Expression of inducible nitric oxide synthase was elevated in the livers of mice fed the USF+ethanol diet compared to pair-fed animals (Figure 3G). There were no changes in inducible nitric oxide synthase expression in SF+ethanol-fed mice compared to controls. No differences were observed in the expression of any liver proinflammatory cytokines between SF and USF in animals fed the control diets. Expression of the macrophage M2 type (anti-inflammatory) response gene, arginase 1, was not altered by the ethanol administration or type of dietary fat, whereas
transforming growth factor-β1 was up-regulated in USF+ethanol-fed mice and was unchanged in SF+ethanol-fed mice (Figure 3, H and I). Taken together, a USF compared to SF diet enhanced ethanol-mediated hepatic neutrophil infiltration and increased the macrophage M1 proinflammatory response (with the exception of MCP-1 mRNA, which was up-regulated in both SF+ethanol and USF+ethanol groups).

Oxidized Metabolites of Linoleic Acid Induce Proinflammatory Response in Macrophages

To directly determine the effect of OXLAMs on cytokine expression, we assayed their effect on RAW264.7 cells, a mouse monocyte/macrophage cell line. RAW264.7 cells were treated overnight with 5 μmol/L 9- or 13-HODE or vehicle (25 mmol/L ethanol) and then with or without 100 ng/mL LPS for 3 hours. The expression of several cytokines was then determined by RT-PCR. Stimulation of RAW264.7 cells by 9-HODE alone, but not 13-HODE alone, led to a significant increase in TNF-α expression (Figure 4A). A similar pattern was observed when the expression of macrophage inflammatory protein-2α and MCP-1 was assayed (Figure 4, B and C), where 9-HODE alone and in combination with LPS enhanced their expression, but 13-HODE alone did not; rather, it slightly attenuated LPS induction of these two cytokines. In contrast, 13-HODE, but not 9-HODE, potentiated LPS induction of inducible nitric oxide synthase expression (Figure 4D). These results demonstrate that 9- and 13-HODEs have different, and sometimes opposing, roles in regulating cytokine gene expression in RAW264.7 cells, with 9-HODE promoting a proinflammatory response (M1 response) to a greater extent than 13-HODE. To determine the effect on the resolution phase of the inflammatory response (M2 response), the expression of arginase 1 and transforming growth factor-β1 was determined. There were no statistically significant effects of either 9- or 13-HODE on the expression of arginase 1 or transforming growth factor-β1 mRNA levels (Figure 4, E and F).

Loss of Alox15 Expression Modestly Attenuates Chronic—Binge-Ethanol-Induced Liver Injury with No Effect on Hepatic Steatosis

12/15-LO is the main enzyme involved in LA oxidation and OXLAM production and may represent a therapeutic target in ALD. Therefore, we questioned whether deficiency of

**Figure 4** The 9- and 13-HODE--induced cytokine expression in RAW264.7 macrophages. A--D: RT-PCR analysis of the expression of M1 macrophage markers: A: Tumor necrosis factor-α (TNF-α). B: Macrophage inflammatory protein-2α (MIP-2α). C: Monocyte chemoattractant protein-1 (MCP-1). D: Inducible nitric oxide synthase (iNOS). E and F: M2 macrophage markers: arginase-1 (ARG-1) (E) and transforming growth factor-β (TGF-β1) (F). mRNA levels were measured by RT-PCR. Genes were normalized to 18S rRNA as an internal control. Results are presented as fold change relative to vehicle/control (25 mmol/L ethanol). Data are expressed as means ± SEM. n = 3 (A--F). *P < 0.05. HODE, hydroxyoctadecadienoic acid.
Alox15, the gene encoding 12/15-LO, would reduce OXLAM levels and attenuate liver damage caused by the combination of ethanol and a LA-rich diet. We used a chronic-binge ethanol-induced model of ALD, where Alox15/C0/C0 and WT mice were fed USF (LA-enriched) control- or ethanol-containing diets. Body composition measurements revealed that the body weights of Alox15/C0/C0 mice were slightly lower than WT mice, but neither group had any significant weight loss because of ethanol (Table 4). However, the liver weight and liver/body weight ratio was significantly increased in both Alox15/C0/C0 and WT mice in response to ethanol feeding. There was a reduction in the amount of epididymal white adipose tissue in both groups, although epididymal white adipose tissue/body weight ratios were similar in the pair-fed and ethanol groups. Control diet–fed Alox15/C0/C0 mice had modestly, but not significantly, lower levels of liver OXLAMs, including 9- and 13-HODEs, compared to WT animals (Figure 5, A and B). However, OXLAM levels were similarly elevated in livers in both WT and Alox15/C0/C0 mice in response to chronic-binge ethanol exposure. In addition to OXLAMs, we measured hydroxyeicosatetraenoic acids (HETEs), which are the 12/15-LO oxidation products of arachidonic acid. There were no noticeable differences in the levels of 15-HETE between Alox15/C0/C0 and WT mice, whereas 12-HETE levels trended lower in Alox15/C0/C0 compared to WT mice in both control- and ethanol-fed mice (Figure 5, C and D). Notably, the degree of hepatic steatosis was similar in WT and Alox15/C0/C0 mice fed ethanol-containing diets (Figure 5, E and F), whereas chronic-binge–ethanol challenge in WT resulted in greater liver injury (compared to controls) than in Alox15/C0/C0 mice (Figure 5G). The

Table 4  Body Composition of WT and Alox 15−/− Mice in an Experimental Animal Model of Chronic-Binge—Induced Liver Injury

<table>
<thead>
<tr>
<th>Parameters</th>
<th>WT</th>
<th>Pair fed</th>
<th>Ethanol</th>
<th>Alox 15−/−</th>
<th>Pair fed</th>
<th>Ethanol</th>
</tr>
</thead>
<tbody>
<tr>
<td>Body/tissue weights</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Body weight, g</td>
<td>23.8 ± 0.72</td>
<td>20.81 ± 0.56</td>
<td>23.8 ± 0.72</td>
<td>20.4 ± 0.63</td>
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<tr>
<td>Liver weight, g</td>
<td>0.77 ± 0.04</td>
<td>0.72 ± 0.04</td>
<td>0.77 ± 0.04</td>
<td>0.97 ± 0.01</td>
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</tr>
<tr>
<td>Liver/body weight</td>
<td>0.032 ± 0.0008</td>
<td>0.044 ± 0.001*</td>
<td>0.035 ± 0.001</td>
<td>0.047 ± 0.001*</td>
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<tr>
<td>eWAT weight, g</td>
<td>0.37 ± 0.03</td>
<td>0.27 ± 0.02</td>
<td>0.37 ± 0.03</td>
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<tr>
<td>eWAT/body weight</td>
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<td>0.015 ± 0.001</td>
<td>0.007 ± 0.001</td>
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</tbody>
</table>

Data are presented as the means ± SEM, n = 7 to 10 animals per group.
*P < 0.05 pair-fed versus ethanol-fed mice.
eWAT, epididymal white adipose tissue; WT, wild-type.

Figure 5  Genetic deletion of Alox15 modestly attenuates chronic-binge—ethanol (EtOH)–induced liver injury with no effect of hepatic steatosis. A and B: Liver levels of 9- and 13-HODEs. C and D: Liver 15- and 12-HETE levels. E: Representative images of hepatic hematoxylin and eosin (H&E) staining. F: Biochemical assessment of liver triglycerides (TGs). G: Serum ALT levels (data are from two independent experiments). Data are expressed as means ± SEM. Each experiment is a representative or the average of 5–8 mice per group. *P < 0.05. Original magnification, ×400 (E). WT, wild-type. HETE, hydroxyeicosatetraenoic acid; HODE, hydroxyoctadecadienoic acid.
expression of CYP2E1 was similarly increased by ethanol in both WT and Alox15−/− mice (Figure 6, A and B). Furthermore, in comparison to pair-fed control groups, the expression of TNF-α and macrophage inflammatory protein-2α (proinflammatory cytokines) was elevated in the livers of both WT and Alox15−/− ethanol-fed mice (Figure 6, C and D). Analysis of chloroacetate esterase staining revealed lower levels of hepatic neutrophil infiltration in Alox15−/− compared to WT animals in response to ethanol exposure (Figure 6, E and F). Collectively, these data demonstrate that disruption of the 12/15-LO pathway was not sufficient to decrease ethanol-induced OXLAM levels; nevertheless, Alox15 gene knockout attenuated liver injury caused by LA-enriched diet and chronic-binge ethanol exposure.

**Discussion**

In the present study, we report that the combination of ethanol and a diet rich in LA, an ω-6 PUFA, exacerbated chronic-binge ethanol-induced liver damage in mice. This is in agreement with previously published studies in various murine models of chronic-ethanol–induced ALD, confirming the harmful effects of dietary LA in experimental ALD regardless of the nature of ethanol administration. The results from this study suggest that an increase in OXLAMs (specifically 9-HODE), which can potently induce a proinflammatory response in macrophages, is likely one of the mechanisms triggering the progression from alcohol-induced steatosis to alcoholic steatohepatitis.

Liver steatosis is an early response to ethanol exposure. In line with previous reports, the degree of chronic-binge ethanol-induced hepatic steatosis in our study was greater in mice fed a USF (corn oil/LA-enriched) compared to animals fed an SF (medium-chain triglyceride oil-enriched) diet. One of the critical points in ALD pathogenesis is progression from simple steatosis to alcoholic steatohepatitis, which is characterized by the combination of hepatic fat accumulation and inflammation. An increase in neutrophil infiltration and proinflammatory cytokine production is a hallmark of ethanol-induced liver inflammation. Neutrophils (polymorphonuclear leukocytes, a major innate immune cell subset) play a central role in the host defense against infection- and sterile-based types of inflammation. After alcohol consumption, neutrophils might be recruited to the

**Figure 6** Effect of Alox15 deficiency on cytochrome p50 2E1 (CYP2E1) expression and markers of liver inflammation. A: CYP2E1 protein expression analyzed by Western blotting with glyceraldehyde-3-phosphate dehydrogenase (GAPDH) as loading control (Ctrl). B: The intensity of protein bands (from A) was quantified by densitometry using the ImageJ software. C: Tumor necrosis factor-α (TNF-α) mRNA. D: Macrophage inflammatory protein 2-α (MIP-2α) mRNA. C and D: Hepatic mRNA levels were measured by RT-PCR. Genes were normalized to 18S rRNA as an internal control. E: Representative images of chloroacetate esterase staining (CAE; arrows indicate CAE-positive neutrophils). F: Quantification of CAE staining performed by counting CAE-positive neutrophils in a random series of five digital images per animal. Results are presented as fold changes relative to the WT pair-fed group. Data are expressed as means ± SEM. Each experiment is a representative or the average of 5–8 mice per group. *P < 0.05. Original magnification, ×400 (E). EtOH, ethanol; WT, wild-type.
liver by chemoattractant cytokines and through their recognition of damage-associated molecular patterns (eg, high mobility group box-1) that are released from the damaged hepatocytes.\textsuperscript{31,42} The enhanced accumulation of neutrophils in hepatic tissue may lead to elevated production of proinflammatory mediators, reactive oxygen species, and production/release of proteases and hypochlorous acid,\textsuperscript{43} thus exacerbating liver injury.\textsuperscript{44} Our observation of increased neutrophil numbers in the livers of ethanol-fed mice is consistent with previous reports using this animal model of ALD,\textsuperscript{45} and suggests that elevated hepatic neutrophil infiltration in response to ethanol and USF compared to SF diet might be one of the contributing factors to more severe liver injury observed in these animals. In our study, USF+ethanol compared to SF+ethanol administration resulted in enhanced hepatic M1 type macrophage proinflammatory responses with significant elevation of TNF-\textit{z} and IL-1\textit{f} gene expression. Dysfunction of intestinal tight junction integrity, elevated endotoxemia, and associated Toll-like receptor\textsuperscript{4}–mediated increase in production of proinflammatory cytokines in liver macrophages have been implicated in the harmful effects of dietary USF in comparison to the protective effects of dietary SF in experimental animal models of ALD.\textsuperscript{7,14,46,47} Previous studies have reported either elevated or unchanged serum endotoxin levels in this particular (chronic hepatic ethanol exposure) experimental model of ALD in mice.\textsuperscript{30,48,49} In the present study, we observed no differences in LPS levels between SF and USF control- or ethanol-fed animals, suggesting that other factors (eg, diet-derived mediators) might contribute to the liver injury caused by USF and chronic binge ethanol exposure. In this regard, elevated levels of endogenous hepatic and circulating OXLAMs associated with USF and ethanol-induced liver inflammation and injury are one of the most important findings of the current study. Moreover, the increase in 9- and 13-HODE levels observed in the current study, in the chronic-binge ethanol model of ALD, as well as in chronic ethanol-induced ALD in mice (I.A.K., unpublished data), likely represents one of the common diet-dependent pathways/mechanisms contributing to ethanol-induced liver damage. The 9- and 13-HODEs can be formed enzymatically from LA, primarily via the actions of cellular lipoxigenases (eg, 12/15-LO), or nonenzymatically via free radical–mediated oxidation in response to oxidative stress. Elevated levels of hepatic CYP2E1, a marker of oxidative stress, were found in our study (in response to chronic-binge ethanol exposure) as well as in previous studies (in chronic ethanol feeding models\textsuperscript{6,7}), regardless of the type of dietary fat. The level of TBARS was elevated to a greater extent in USF+ethanol- compared to SF+ethanol-fed animals. This could be because of a high content of LA in a USF diet, which is prone to oxidative degradation, leading to production of a variety of oxidative metabolites. Therefore, oxidative stress and availability of the substrate (LA-enriched diet versus medium-chain triglyceride–enriched diet) are, most likely, contributing factors to the elevated levels of OXLAMs observed in mice fed USF+ethanol compared to SF+ethanol-fed animals. Elevated plasma 9- and 13-HODE levels have been found in rodents in parallel with alcohol-induced hepatic steatosis, oxidative stress, and hepatocyte damage,\textsuperscript{29,30} and in patients with alcoholic cirrhosis.\textsuperscript{28} In the study by Raszeja-Wyszomirska et al,\textsuperscript{28} the authors demonstrated that the concentrations of HODEs and HETEs were significantly different between healthy individuals, nonalcoholic fatty liver disease patients, and ALD patients. Specifically, patients with ALD had 46 times higher HODE levels than nonalcoholic fatty liver disease patients. Furthermore, HODE levels were higher in men with ALD compared to women with ALD. In addition, the expression of ALOX15 and ALOX15B was increased in patients with ALD compared to healthy controls.

The central hypothesis of the current study was that a diet with a high content of LA exacerbates ethanol-induced injury, in part via increased OXLAM production and OXLAM-mediated induction of hepatic proinflammatory responses. By using RAW264.7 cells, we directly tested the effect of each HODE on the production of inflammatory cytokines in a monocyte/macrophage cell type. In general, we found that 9-HODE increased proinflammatory cytokines, either alone or in combination with LPS. We also found that 13-HODE was largely ineffective, and even opposed to the activity of 9-HODE with respect to LPS-induced cytokine expression (Figure 4). There was no effect of either 9- or 13-HODE on the expression of M2 macrophage markers, suggesting that 9-HODE was primarily proinflammatory and that 13-HODE was mainly neutral or had some anti-inflammatory activity. These data are consistent with previous reports of differential activities of 9- and 13-HODE. In studies using human keratinocytes, Hattori et al\textsuperscript{22} demonstrated that 9-HODE, acting through G-protein coupled receptor (GPR)132, stimulated the expression of IL-6. However, they found no effect by 13-HODE. There are three known receptors for HODEs: GPR132, transient receptor potential vanilloid 1 (TRPV1), and peroxisome proliferator-activated receptor \(\gamma\) (PPAR\(\gamma\)). GPR132 binds to 9-HODE but binds only weakly to 13-HODE.\textsuperscript{18} TRPV1 and PPAR\(\gamma\) can bind to both 9- and 13-HODE\textsuperscript{25,50} RAW264.7 cells do not express TRPV1\textsuperscript{51} and express only low levels of PPAR\(\gamma\).\textsuperscript{52} Therefore, the primary response to HODEs in RAW264.7 cells is likely mediated by GPR132 and explains the relative inactivity of 13-HODE in these cells. Although GPR132 is the primary HODE receptor expressed in macrophages,\textsuperscript{53} the expression of TRPV1 and PPAR\(\gamma\) is much more widespread. PPAR\(\gamma\) is found in adipose tissue, the small intestine, muscle, brain, and liver.\textsuperscript{54,55} and TRPV1 has been found to be expressed in the liver and brain.\textsuperscript{57} Therefore, HODEs may affect multiple organ systems, including the liver, by acting through other receptors. Indeed, we have previously shown that \(Trpv1^{−/−}\) mice have much improved indexes of liver
injury and inflammation, such as decreased plasma ALT and decreased expression of several markers of inflammation (eg, TNF-α, IL-1β, IL-12p40, IL-6, MCP-1, and macrophage inflammatory protein-2α) after ethanol feeding.60

12/15-LO is the major enzymatic pathway of OXLAM production, and 12/15-LO deficiency protects against high-fat diet–induced insulin resistance, adipose tissue inflammation, and nonalcoholic fatty liver disease.58–61 Thus, 12/15-LO knockout mice were protected from high-fat diet–induced liver steatosis, liver inflammation, and lymphocyte infiltration in the liver.61 Disruption of Alox15 in hyperlipidemic ApoE–/– mice resulted in reduced serum ALT levels, decreased hepatic steatosis, inflammation with significant reductions in the expression of TNF-α, IL-18, and MCP-1, and macrophage infiltration, all of which were associated with lower hepatic 12-HETE levels.60 In addition, the absence of Alox15 reduced hepatocyte apoptosis, as revealed by reduced caspase-3 levels.60 In our study, Alox15 deficiency in mice resulted in the attenuation of ethanol-induced liver injury; however, there was no effect on hepatic steatosis. This might be because of the decreased levels of ALOX15-generated metabolites, some of which could be hepatotoxic. Notably, the absence of Alox15 in mice did not appreciably alter chronic-binge–ethanol-induced OXLAM levels. This observation suggests that nonenzymatic LA oxidation via free radical–mediated oxidation in response to oxidative stress, rather than through the 12/15-LO pathway of LA oxidation, is likely the main mechanism contributing to ethanol-mediated increase in OXLAM production. Another potential source of OXLAM production is that mediated by the cytochrome P450 system62,63; however, the significance of this pathway in ALD at this point is unclear. Although the absence of Alox15 conferred partial protection to the liver, that protection cannot be attributed to lower OXLAM levels, as liver 9- and 13-HODEs were similarly elevated in response to ethanol in WT and Alox15–/– mice. This suggests that other 12/15-LO–derived metabolites (eg, HETEs, the products of arachidonic acid oxidation) may play a role. Among several HETEs, 12-HETE (a major product of 12/15-LO in rodents) was reduced in the liver tissue from ethanol-fed Alox15–/– compared to WT mice. 12-HETE is a known proinflammatory lipid mediator through the induction of TNF-α, MCP-1, and IL-6 expression in macrophages.64,65 An increase in 12-HETE was observed in patients with ALD28 and was associated with liver inflammation and injury in hyperlipidemia-prone ApoE–/– mice; genetic disruption of Alox15 protected these mice against liver injury.60 In addition, the lower levels of other proinflammatory lipid mediators from arachidonic acid (eg, leukotriene B4 previously implicated in neutrophil recruitment66–68) in Alox15–/– compared to WT mice (I.A.K., unpublished data) might confer a decrease of neutrophil infiltration observed in ethanol-fed Alox15–/– compared to WT animals. Notably, the 12/15-LO pathway plays an important role in both inflammation and production of factors that promote resolution of inflammation (eg, lipoxin A4).69,70 Therefore, further studies examining the role of various lipid mediators involved in the development, progression, or resolution of ethanol-mediated inflammation are warranted.

In summary, the results of the current study support the concept that dietary LA, a ω-6-PUFA, exacerbates ethanol-induced liver injury and provides evidence that the increase in OXLAM production and promotion of an OXLAM-mediated proinflammatory response might be one of the underlying mechanisms. Strategies to decrease LA consumption through dietary modifications may be an effective approach to reduce ethanol-induced OXLAMs and to attenuate ALD.

Acknowledgments

We thank Jingwen Zhang and Ying Song for assistance with tissue staining, and Marion McClain for proofreading the manuscript.

I.A.K., C.E.R., A.E.F., and C.J.M. designed the study and interpreted the data; A.E.F. and S.S. performed OXLAM analyses; B.G. provided critical discussion; H.L. and M.E.M. performed animal studies and acquired data; D.R.W. acquired and analyzed data; I.A.K. and D.R.W. wrote the manuscript.

References


70. Vangaveti V, Baune BT, Kennedy RL: Hydroxyoctadecadienoic acids; novel regulators of macrophage differentiation and atherogenesis. Ther Adv Endocrinol Metab 2010, 1:51—60