Lithocholic Acid Feeding Induces Segmental Bile Duct Obstruction and Destructive Cholangitis in Mice

We determined the mechanisms of hepatobiliary injury in the lithocholic acid (LCA)-fed mouse, an increasingly used model of cholestatic liver injury. Swiss albino mice received control diet or 1% (w/w) LCA diet (for 1, 2, and 4 days), followed by assessment of liver morphology and ultrastructure, tight junctions, markers of fibrosis and key proteins of hepatobiliary function, and bile flow and composition. As expected LCA feeding led to bile infarcts, which were followed by a destructive cholangitis with activation and proliferation of periductal myofibroblasts. At the ultrastructural level, small bile ducts were frequently obstructed by crystals. Biliary-excreted fluorescence-labeled ursodeoxycholic acid accumulated in bile infarcts, whereas most infarcts did not stain with India ink injected into the common bile duct; both findings are indicative of partial biliary obstruction. Expression of the main basolateral bile acid uptake proteins (sodium-taurocholate cotransporter and organic anion-transporting polypeptide 1) was reduced, the canalicular transporters bile salt export pump and multidrug-related protein 2 were preserved, and the basolateral transporter multidrug-related protein 3 and the detoxifying enzyme sulfotransferase 2a1 were induced. Thus, we demonstrate that LCA feeding in mice leads to segmental bile duct obstruction, destructive cholangitis, periductal fibrosis, and an adaptive transporter and metabolic enzyme response. (Am J Pathol 2006, 168:410–422; DOI: 10.2353/ajpath.2006.050404)

The effects of the monohydroxy bile acid lithocholic acid (LCA) have been studied in rodents (ie, mainly in the rat, hamster, and rabbit) since the early 1960s to understand the role of potentially hepatotoxic bile acids in the pathogenesis of cholestatic liver injury.1–4 The proposed concepts on the pathogenesis of LCA-induced cholestasis in rodents (ie, mainly rats) include biochemical alterations of the bile canalicular membrane,5,6 the development of crystalline plugs in bile canaliculi due to the poor solubility of LCA,3,7 and impaired trafficking or increased retrieval of canalicular export pumps to and from the canalicular membrane.8,9 In addition, LCA is increasingly being investigated in (knockout) mice to clarify the role of hepatic metabolic phase I and II detoxifying enzymes and their regulatory nuclear receptors in the hepatic defense against toxic bile acids.10–15 However, the cholestatic phenotype and the pathogenesis of hepatobiliary injury of LCA-treated mice are still poorly understood.

Potentially toxic bile acids may affect not only hepatocytes but also bile duct epithelial cells (cholangiocytes).16 Under physiological conditions, bile acids and biliary phospholipids are transported into bile via the bile salt export pump (Bsep/Abcb11) and the canalicular phospholipid flippase (multidrug resistance gene 2 [Mdr2]/Abcb4), respectively, and subsequently form mixed micelles, which protect cholangiocytes from bile acid toxicity.17 We have previously described the development of sclerosing cholangitis in mice with targeted disruption of the Mdr2 gene and subsequent absence of biliary phospholipid secretion.18,19 This results in an in-
Immunohistochemistry for Cytokeratin 19 (CK19)

Immunohistochemistry for CK19 was performed on cryosections fixed with a modified formalin/methanol/acetone fixation. In brief, sections were fixed in 4% buffered formaldehyde for 5 minutes and then fixed with methanol for 5 minutes at −20°C and with acetone for 3 minutes at −20°C. CK19 was detected using the monoclonal rat anti-Troma-III antibody (developed by Rolf Kemler, Max Planck Institute, Freiburg, Germany), which was obtained from the Developmental Studies Hybridoma Bank (developed under the auspices of the National Institute of Child Health and Human Development and maintained by the University of Iowa, Iowa City, IA). Binding of the antibody was detected using the ABC system (Dako) with AEC (Dako) as substrate.

Bile Flow and Composition

Bile flow was determined as described previously. Biliary phospholipid concentrations were determined using a commercially available kit (Phospholipid B; Wako, Neuss, Germany) according to the manufacturer’s instructions. In detail, 5 μl of bile was mixed with 750 μl of color reagent and incubated for 10 minutes at 37°C and measured at 505 nm. Concentrations were determined using the standard solution of the kit. Biliary cholesterol concentrations were determined using a commercially available kit (cholesterol liquicolor; Human, Wiesbaden, Germany) according to the manufacturer’s instructions. For cholesterol extraction, 5 μl of bile and 95 μl of distilled water were mixed with 400 μl of chloroform:methanol (2:1) and centrifuged 5 minutes at 14,000 g. The lower phase was concentrated, and the obtained pellet was dissolved in 5 μl of methanol. After the addition of 500 μl of color reagent and incubation at 37°C for 5 minutes, probes were measured at 500 nm. Concentration was determined using the standard solution of the kit. For biliary glutathione, proteins in bile were precipitated with 5% metaphosphoric acid (2.5 μl of bile in 50 μl of metaphosphoric acid), and glutathione analysis was performed using the glutathione assay kit (Calbiochem, San Diego, CA), according to the manufacturer’s instructions. Biliary bile acid concentration required bile to be diluted 1:300 in physiological saline and analyzed using an Aα-hydroxysteroid dehydrogenase assay (Ecoline S”; Diasys, Holzheim, Germany) according to the manufacturer’s instructions. Equipment and conditions used for extraction, electrospray mass spectrometry, and gas chromatography-mass spectrometry were the same as previously described.

mRNA Analysis and Polymerase Chain Reaction of Transporters, Metabolic Enzymes, and Fibrosis Key Genes

RNA isolation, cDNA synthesis, and Taqman real-time polymerase chain reaction were performed as described.
previously. The following primers and 5'-FAM, 3'-TAMRA-labeled probes were used: multidrug resistance-associated protein (Mrp2); bile salt export pump (Bsep); 28S rRNA; Mrp3; cytochrome p450 (Cyp) 3a11 fwd, 5'-CCACCAGTAGACACTTTCC-3'; Cyp3a11 rev, 5'-TTCATCTCCATCAAGGTGTTGTACC-3'; Sult2a1 fwd, 5'-GGAAGGACCACGACTCATAAC-3'; Sult2a1 rev, 5'-GATTCTTCACAAGGTTTGTGTTACC-3'; Sult2a1 probe, 5'-CCCATCCATCTCTTCTCCAAGTCTTTCTTCAG-3'; sodium-taurocholate cotransport (Ntcp) fwd, 5'-CACCATGGAGTTCAGCAAGA-3'; Ntcp rev, 5'-AGCACTGAGGGGATGATAC-3'; Ntcp probe, 5'-AGGC-TCACTTCTGGAAGC-CCAAA-3'; collagen 1a1 (Col1a1) fwd, 5'-CAATGCAATGAAGAACTGGACTGT-3'; Col1a1 rev, 5'-TCCTACATCTTCTGAGTTTGGTGA-3'; Col1a1 probe, 5'-CAGAAAGCACAACCTCGCCCTCC-3'; tissue inhibitor of metalloproteinases 1 (TIMP-1) fwd, 5'-CATGGAAAGCCTCTGAGATATG-3'; TIMP-1 rev, 5'-AA-GCTGCAAGCCATTTGATG-3'; TIMP-1 probe, 5'-CTCA-TCACCGCGGGCAAAAGAA-3'; matrix metalloprotease-2 (MMP-2) fwd, 5'-CTTTGAGAAGGATGGCAAGTATGG-3'; MMP-2 rev, 5'-

![Figure 1](image1.png)

**Figure 1.** Macroscopic appearance of the liver in lithocholic acid-fed mice. Compared with the control liver on the left, the liver from a mouse fed a 1% (w/w) LCA-supplemented diet for 4 days is enlarged and shows numerous small necrotic areas (ie, bile infarcts) on its surface (indicated by arrows). Note also the enlarged gall bladder in the LCA-fed mouse liver (indicated by the asterisk).

![Figure 2](image2.png)

**Figure 2.** Lithocholic acid (LCA) feeding leads to bile infarcts and destructive cholangitis in mouse liver. Livers of mice fed 1% (w/w) LCA-supplemented diet for 2 days (A) and 4 days (B–F). A: Liver histology of a 2-day LCA-fed mouse reveals characteristic bile infarcts (asterisks) and disseminated single-cell necroses (arrows) in hepatic acinar zone 1 and 2. B: Small bile duct in a 4-day LCA-fed mouse liver with pronounced cholangitis with numerous neutrophils surrounding and invading the bile duct. C and D: Larger bile duct with pronounced periductal edema, pericholangitis, periductal fibrosis, and subepithelial fluid accumulation leading to detachment of biliary epithelium (asterisk). Arrowheads mark pseudopapillary proliferation of BECs. E and F: Bile duct with pericholangitis, periductal fibrosis, and ulceration of the bile duct epithelium (asterisk). Note also neutrophils invading the ulcer (F). bd, bile duct. hematoxylin and eosin staining; original magnification for A, ×10; for B, ×63; for C and E, ×20; for D and F, ×40.

![Figure 3](image3.png)

**Figure 3.** LCA feeding leads to obstruction of intralobular bile ducts by crystals, bile infarcts, and destructive cholangitis in mouse liver—ultrastructural characteristics. A–F: Electron microscopy from the liver of a 4-day LCA-fed mouse. A: Totally obstructed interlobular bile duct containing crystalline material. B: Larger magnification of the highlighted area of A shows crystals spearing BECs. C: Large crystal (indicated by the arrowheads) spearing BECs of an interlobular bile duct. D: Bile infarct (asterisk) containing crystalline material surrounded by severely altered hepatocytes and invading neutrophils (arrowhead). E: Interlobular bile duct with altered bile duct epithelium and ulceration (indicated by the asterisk). Arrowheads mark pseudopapillary proliferation of BECs. F: Higher magnification of the ulcer showing invading neutrophilic granulocytes (NG). bd, bile duct.
TTTAGGAGGTGCCCTGG-AA-3' and MMP-2 probe, 5'-CAGATGGACAGCCCTGCAAGTTCCC-3'.

Preparation of Liver Membranes and Analysis of Transporter Protein Levels by Western Blotting

Liver membranes were prepared as described previously.21 Transporter protein levels were determined using polyclonal antibodies against Bsep (dilution, 1: 7500; kindly provided by Dr. Renxue Wang, Vancouver, Canada), Mrp2 (dilution, 1: 1000; kindly provided by Dr. Bruno Stieger, Zurich, Switzerland), Mrp3 (dilution, 1: 1000; kindly provided by Dr. Dietrich Keppler, Heidelberg, Germany), Ntcp (dilution, 1: 2500; kindly provided by Dr. Bruno Stieger), and organic anion transporting protein (Oatp1) (dilution, 1: 1000; kindly provided by Dr. Bruno Stieger). Blots were reprobed with an anti-actin antibody (dilution, 1:5000; Sigma, Steinheim, Germany) to confirm the specificity of changes in transporter protein levels.21,22,24

Preparation of Total Liver Protein and Analysis of αSMA Expression by Western Blotting

Protein was isolated by sonicating liver tissue in a homogenization buffer (0.25 mol/L sucrose, 10 mmol/L HEPES, pH 7.5, and 1 mmol/L EDTA, pH 8.0, containing protease inhibitors phenylmethylsulfonyl fluoride, aprotinin, leupeptin, and pepstatin). Protein (30 mg) was run on a 10% sodium dodecyl sulfate-polyacrylamide gel, transferred to nitrocellulose, and blotted with a monoclonal mouse antibody against α-SMA (dilution, 1:2000; Dako). Binding was detected using peroxidase-conjugated rabbit anti-mouse immunoglobulins (Dako). Determination of protein concentration, protein loading, and protein expression levels was performed as described previously.21,22 Blots were reprobed with an anti-α-SMA antibody (dilution, 1:5000; Sigma, Steinheim, Germany) to confirm the specificity of changes in α-SMA expression.

Immunofluorescence Microscopy

Hepatocellular localization of the key basolateral (Ntcp and Mrp3) and canalicular (Mrp2 and Bsep) transporters, staining for cytokeratin 8/18 (CK8/18), CK19, laminin, and activated caspase-3 was studied by immunofluorescence microscopy as described previously.18,21

Lobular Distribution of India Ink and Fluorescent-Labeled Ursodeoxycholic Acid (UDCA)

India ink was injected into the common bile duct and visualized as previously described18 to determine its lobular distribution in LCA-fed mice and for comparison in 3-day common bile duct ligated (CBDL)-mice.18 To study a potential relationship between the cholestatic phenotype observed in LCA-fed mice and the lobular distribution of bile acids, fluorescent UDCA (UDC-lysyl-nitrobenzoxadiazolyl; kindly provided by Dr. Alan Hofmann, University of California, San Diego, La Jolla, CA) was injected into the inferior vena cava, and excreted fluorescent UDCA was viewed by fluorescence microscopy and compared with CBDL mice as described previously.19

Table 1. Serum Biochemistry of 1-, 2-, and 4-Day 1% LCA-Fed Mice

<table>
<thead>
<tr>
<th></th>
<th>Control (n = 6)</th>
<th>1-day LCA (n = 5)</th>
<th>2-day LCA (n = 5)</th>
<th>4-day LCA (n = 8)</th>
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<tr>
<td>ALT (U/L)</td>
<td>83 ± 42</td>
<td>102 ± 47</td>
<td>2398 ± 2278</td>
<td>4837 ± 1884*</td>
</tr>
<tr>
<td>AP (U/L)</td>
<td>220 ± 65</td>
<td>165 ± 58</td>
<td>536 ± 323</td>
<td>1334 ± 555*</td>
</tr>
<tr>
<td>Bili tot (mg/dl)</td>
<td>0.15 ± 0.05</td>
<td>0.32 ± 0.13</td>
<td>0.62 ± 0.33</td>
<td>2.33 ± 1.02*</td>
</tr>
<tr>
<td>SBA (μmol/L)</td>
<td>21 ± 25</td>
<td>16 ± 4</td>
<td>70 ± 67</td>
<td>280 ± 118*</td>
</tr>
</tbody>
</table>

*P < 0.05 compared with standard diet-fed controls (ANOVA with Bonferroni post testing).

Values are expressed as mean ± SD. ALT, alanine aminotransferase; Bili tot, total bilirubin; SBA, serum bile acids.
Figure 6. Tight junctions are altered in LCA-fed mouse liver. Double-immunofluorescence labeling for CK8/18 (in green) and the tight junction protein ZO-1 (in red) in control liver (A–C and G–I) and the liver from a 4-day LCA-fed mouse (D–F and J–L). B and C: ZO-1 staining of bile ducts in control diet-fed mice shows a distinct pattern outlining the cell contacts between BECs. E and F: In contrast, the liver of a 4-day LCA-fed mouse shows strikingly altered tight junctions morphologically characterized by partially disrupted and missing ZO-1 staining (arrowheads). Note also that BECs still form an intact epithelial monolayer (D and F), suggesting that the observed missing ZO-1 staining is not related to BEC necrosis. G–I: Staining of the intermediate filament network and tight junctions of hepatocytes in control diet-fed mice. J: The liver of a 4-day LCA-fed mouse shows a significantly increased density of the cytokeratin intermediate filament (CK-IF) network with a pronounced pericanalicular sheath (arrowheads). K: Strikingly altered tight junctions between hepatocytes in 4-day LCA-fed mouse liver morphologically characterized by elongation and distortions of the ZO-1 staining pattern (arrowheads). bd, bile duct. Bar = 10 μm.
Bacterial Counts of Liver and Spleen

To determine the possible role of bacterial translocation in the pathogenesis of LCA-induced liver injury in mice, bacterial counts of liver and spleen were performed as previously described.\(^{19}\)

Measurement of Hepatic Hydroxyproline Content

To quantify liver fibrosis, hepatic hydroxyproline content was determined. The right liver lobe was homogenized in 6 N HCl (200 mg liver tissue/4 ml HCl) and hydrolyzed at 110°C for 16 hours. After filtration, 50 \(\mu\)l was added to 450 \(\mu\)l of 2.2% NaOH dissolved in citrate-acetate buffer (50 g of citric acid \(\times\) \(\text{H}_2\text{O}\), 12 ml of acetic acid, 120 g of sodium acetate \(\times\) 3 \(\text{H}_2\text{O}\), 34 g of NaOH, and 1 L of distilled water; pH 6.0). After neutralization, 250 \(\mu\)l of chloramin T solution was added and incubated for 20 minutes. After adding 250 \(\mu\)l of perchloric acid and 12 minutes of incubation at room temperature, 250 \(\mu\)l of \(p\)-dimethylaminobenzaldehyde solution was added and incubated at 60°C for 20 minutes. Hydroxyproline content was measured at 565 nm using a hydroxyproline standard curve.

Statistical Analysis

Data are reported as arithmetic means ± SD of five to eight animals in each group. Statistical analysis was performed using Student’s t-test when appropriate or analysis of variance with Bonferroni post test when three or more groups were compared using the Sigmastat statistics program (Jandel Scientific, San Rafael, CA). A \(P\) value <0.05 was considered significant.

Results

LCA Feeding Leads to Bile Infarcts, Destructive Cholangitis, and Periductal Fibrosis

Beginning at day 2, livers of LCA-fed mice contained numerous bile infarcts (Figures 1 and 2A) that were morphologically indistinguishable from those seen in CBDL mice.\(^{18,25}\) Small interlobular bile ducts were frequently obstructed with crystals, as revealed by electron microscopy (Figure 3, A and B). In addition, crystals penetrating into bile duct epithelial cells (BECs) were also observed (Figure 3C), suggesting that direct physicochemical LCA-induced injury of BECs may be important. The lobular lesions (ie, bile infarcts and single cell necroses) displayed light microscopic and ultrastructural characteristics of oncotic hepatocytes with hydropic swelling and disruption of cell and organelle membranes (not shown) as defined recently.\(^{26}\) In addition, crystals were also deposited within bile infarcts (Figure 3D). These findings suggest that bile infarcts and bile ducts are faced with the highest LCA concentration within the liver. Around days 3 and 4, animals developed a dense neutrophilic-granulocytic infiltrate around small interlobular (Figure 2B) and larger bile ducts (Figure 2, C–F). This observation was further confirmed by the finding of numerous CD11b-positive periductal cells in LCA-fed mice as shown in Figure 4B. BECs were irregular with enlarged nuclei and focal pseudopapillary proliferations (Figure 2, C–E). In addition, subepithelial fluid accumulation lifting the epithelial cell layer (Figure 2, C and D) followed by ulceration of the bile duct epithelium (Figures 2, E and F, and 3, E and F) and pronounced periductal edema was frequently observed in larger ducts. There was no evidence for apoptosis of BECs because we found neither BECs with activated caspase-3 nor BECs with characteristic CK intermediate filament alterations typical for apoptotic cell death\(^{18}\) (not shown), suggesting that necrosis may represent the primary mode of BEC death in our
model. These morphological findings were accompanied by a continuous increase in serum alanine aminotransferase levels followed by elevations of alkaline phosphatase and bilirubin (Table 1). Prolonged feeding of 1% LCA-supplemented diet beyond 7 days was associated with substantial mortality, indicating that continuous long-term treatment of this dose is not feasible (Figure 5). The acute cholangitis with neutrophils raised the possibility of a bacterial etiology related to increased bacterial translocation from the gut into the liver and spleen. However, we found no differences in bacterial counts in livers and spleens between LCA-fed mice and controls (data not shown). The development of pericholangitis in LCA-fed mice led to the hypothesis that this could be due to leakage of the tight junctions of BECs. As determined by immunofluorescence microscopy with staining for tight junction protein ZO-1 (Figure 6, E and F) and by electron microscopy (Figure 3, E and F), tight junctions were irregular and focally disrupted, suggesting that bile ducts were leaky under these experimental conditions. Time course studies revealed that tight junction alterations preceded lifting and ulceration of the epithelial cell layer, suggesting that tight junction alterations may be of primary importance in LCA-induced cholangiopathy. Tight junctions between hepatocytes also showed alterations of the ZO-1 pattern with distortion and widening indicative of dilated canaliculi (Figure 6, K and L).

Moreover, we frequently observed ruptured canals of Hering in the proximity to bile infarcts (Figure 7), parallel to previous observations in CBDL mice.18 The evolving cholangitis in LCA-fed mouse liver was accompanied by a significantly increased number of α-SMA-positive periductal myofibroblasts (Figure 8C) responsible for an astonishingly rapidly evolution and pronounced periductal fibrosis (Figure 8D) comparable with previous observations in CBDL mice.18 In addition, α-SMA protein levels were significantly increased in LCA-fed mice compared with controls (Figure 8E). Increased proliferation of periductal myofibroblasts was also demonstrated by a significantly increased number of Ki-67-positive (CK19-negative) periductal myofibroblasts (Figure 9). Liver fibrosis was confirmed biochemically by significantly higher hepatic hydroxyproline content in LCA-fed mice after 4 days (ie, 194 ± 24 vs. 104 ± 4 mg/g liver in controls; P > 0.05). This was accompanied by a significantly enhanced expression of collagen 1α1 mRNA (52-fold) compared with controls (Table 2). Of interest, the absolute increase of TIMP-1 mRNA expression (424-fold) was greater than for MMP-2 mRNA (10-fold), suggesting that liver fibrosis in LCA-fed mice may, at least in part, result from increased matrix production in the presence of decreased matrix degradation (Table 2). Taken together these findings support the concept that LCA feeding in mice results in partial biliary obstruction and subsequently bile infarcts. In addition, these findings suggest that LCA-induced alterations of tight junctions caused destructive cholangitis and activation of BECs, initiating proliferation and activation of periductal myofibroblasts and leading to periductal fibrosis.

Figure 8. Periductal fibrosis in LCA-fed mouse liver. Immunohistochemistry (A and C) for α-SMA-positive cells and Sirius red staining (B and D) for collagen in control liver (A and B) and in the liver from a 4-day LCA-fed mouse (C and D). A: In the control liver, only smooth muscle cells of the hepatic artery branches (ha) and the portal vein (pv) are α-SMA positive. B: In contrast, there are numerous α-SMA-positive myofibroblasts surrounding the bile duct (bd) of a 4-day LCA-fed mouse. Note that hepatic arteries and portal veins of the control liver and 4-day LCA-fed mice are of the same size, indicating that both situations are cut at the same level of the biliary tree. C and D: Sirius red staining outlines the pronounced periductal fibrosis in the LCA-fed mouse liver. Original magnification, ×20. E: Protein quantification for α-SMA and β-actin by Western blotting in control livers (CO) and livers from 4-day LCA-fed mice (4-day 1% LCA). Note the significant increased α-SMA and unchanged β-actin expression in LCA-fed mouse liver. Densitometry data are expressed as the fold change relative to control diet-fed animals. Values are the means from five animals in each group. *P < 0.05 compared with standard diet-fed controls (Student's t-test).

Comparison of the Lobular Distribution of Fluorescent UDCA and India Ink in LCA-Fed Mice and CBDL Mice: Clues to the Pathogenesis of the Cholestatic Phenotype

Because bile infarcts were frequently observed in proximity to disrupted canals of Hering (Figure 7), we searched for communications between the bile duct system and bile infarcts as previously seen in CBDL mice.18 Fluorescent UDCA injected into the inferior vena cava was rapidly secreted into canaliculi and bile ducts in CBDL and LCA-fed mice (Figure 10, A and C). After retrograde injection into the common bile duct, India ink
filled bile ducts and bile canaliculi and accumulated in bile infarcts (Figure 10B). On the other hand, in LCA-fed mice, India ink filled bile ducts but accumulated in only few bile infarcts (Figure 10D). Taken together, these findings support our concept 1) that bile infarcts have connection to the bile duct system, 2) that bile infarcts contain high concentrations of bile acids, and 3) most importantly, that these experiments are consistent with partial biliary obstruction in LCA-fed mice.

**LCA Feeding Leads to Adaptive Hepatocellular Transporter and Metabolic Enzyme Expression in Mice**

To study the functional importance of the described partial biliary obstruction, we determined bile flow in LCA-fed mice on days 1 and 4 (Table 3). Bile flow tended to increase on day 1 and was significantly decreased on day 4. After an initial increase on day 1, biliary secretion of cholesterol, phospholipids, glutathione, and bile acids was significantly reduced in LCA-fed mice. The predominating bile acids in bile were LCA and chenodeoxycholic acid (Table 4).

Because previous studies in isolated-perfused rat liver have indicated significant alterations in expression and localization of ABC transporter proteins, which may, at least in part, explain cholestasis in LCA-treated rodents, we studied hepatobiliary transport proteins in our model. LCA feeding led to a significant reduction in the expression of the main basolateral bile acid uptake systems Ntcp and Oatp1 (Figures 11 and 12B). Canalicular

| Table 2. Hepatic mRNA Expression of Several Genes in 4-Day 1% LCA-Fed Mice |
|-----------------------------|-----------------------------|
| Control (n = 5) | 4-day 1% LCA (n = 5) |
| Collagen 1a1 | 1.00 ± 0.62 | 51.7 ± 43.6* |
| TIMP-1 | 1.00 ± 0.41 | 424 ± 233* |
| MMP-2 | 1.00 ± 0.23 | 9.5 ± 6.3* |
| Ntcp | 1.00 ± 0.28 | 0.26 ± 0.10* |
| Bsep | 1.00 ± 0.56 | 1.19 ± 0.47 |
| Mrp2 | 1.00 ± 0.61 | 0.83 ± 0.23 |
| Mrp3 | 1.00 ± 0.94 | 3.10 ± 1.03 |
| Cyp3a11 | 1.00 ± 0.89 | 3.33 ± 2.18 |
| Cyp7a1 | 1.00 ± 0.84 | 0.07 ± 0.04* |
| Cyp8b1 | 1.00 ± 0.33 | 0.03 ± 0.01* |
| Sult2a1 | 1.00 ± 0.96 | 7.11 ± 2.69* |

*P < 0.05 compared with standard diet-fed controls (Student’s t-test). Values are normalized to controls as 1.00 and 28S rRNA expression and are expressed as mean ± SD.

Bsep protein levels and localization were not altered in LCA-fed mice compared with controls (Figures 11 and 13B), and there was no evidence of canalicular mistargeting or retrieval of Bsep (Figure 13, E and F). Total Mrp2 protein levels were also unchanged in LCA-fed mice (Figure 11), but the expression of Mrp2 protein was limited to acinar zone 3 (Figure 13, H and I), where it remained localized to the canalicular membrane (Figure

![Figure 9](image1.png) & ![Figure 10](image2.png)

Figure 9. Proliferation of periductal myofibroblasts and bile duct epithelial cells in LCA-fed mouse liver. Immunohistochemistry for (A and C) CK19-positive cholangiocytes and proliferation marker Ki-67 (B and D). A: In the control liver, only some positive nuclei are seen in the portal field. B: In contrast, there are numerous Ki-67-positive (CK19-negative) periductal myofibroblasts surrounding the bile duct and also few proliferating Ki-67-positive BECs in LCA-fed mouse liver. bd, bile duct. Original magnification, ×40.

Figure 10. Bile infarcts in LCA-fed mouse liver fill-up with fluorescent UDCA injected into the inferior vena cava, but partial obstruction of the biliary tree prevents retrograde filling of bile infarcts with India ink. Fluorescent UDCA injected into the inferior vena cava (A and C) and India ink injected into the common bile duct (B and D) in livers from CBDL mice (A and B) and 4-day LCA-fed mice (C and D). A and C: Bile infarcts in CBDL and LCA-fed mouse livers are impregnated by fluorescent UDCA, suggesting that fluorescent UDCA is secreted into bile canaliculi and regurgitates into bile infarcts (surrounded by the white circles) due to outflow obstruction in both models. B: India ink injected into the common bile duct after CBDL fills most bile infarcts (white asterisks). D: India ink injected into the common bile duct enters interlobular bile ducts (arrows) in LCA-fed mouse. India ink stains only the minority of bile infarcts (white asterisk), whereas most remain negative for India ink (black asterisks), suggesting that partially biliary obstruction prevents complete retrograde filling of bile infarcts with India ink in LCA-fed mice. Original magnification, ×10.
Biliary Bile Acid Composition in Control and LCA-Fed Mice  

Table 4. Biliary Bile Acid Composition in Control and LCA-Fed Mice

<table>
<thead>
<tr>
<th>Bile acid</th>
<th>Cholesterol (mmol g⁻¹ LW min⁻¹)</th>
<th>Phospholipids (mmol g⁻¹ LW min⁻¹)</th>
<th>Glutathione (nmol g⁻¹ LW min⁻¹)</th>
<th>Bile acids (mmol g⁻¹ LW min⁻¹)</th>
</tr>
</thead>
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<tr>
<td>Control (n = 6)</td>
<td>1.6 ± 0.4</td>
<td>0.4 ± 0.1</td>
<td>4.1 ± 0.9</td>
<td>6.0 ± 2.0</td>
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<tr>
<td>1-day LCA (n = 6)</td>
<td>1.7 ± 0.5</td>
<td>0.9 ± 0.3</td>
<td>7.7 ± 1.5*</td>
<td>7.2 ± 1.8</td>
</tr>
<tr>
<td>4-day LCA (n = 6)</td>
<td>0.6 ± 0.4*</td>
<td>0.3 ± 0.2*</td>
<td>2.4 ± 1.9*</td>
<td>2.7 ± 2.2*</td>
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*P < 0.05 compared with standard diet-fed controls; †P < 0.05 1- versus 4-day LCA (ANOVA with Bonferroni post test). Values are expressed as mean ± SD. LW, liver weight.

13. K and L. These data suggest that the observed reduction in biliary glutathione secretion in LCA-fed mice (Table 3) was related to impaired Mrp2 expression in acinar zone 1 and 2. Basolateral Mrp3 expression was significantly induced by LCA feeding (Figures 11 and 12D). Interestingly, reduced Ntcp staining was reciprocally accompanied by enhanced Mrp3 staining in LCA-fed mouse liver (Figure 12). The changes in transporter protein levels were accompanied by respective changes in mRNA expression (Table 3), indicating that these changes may be mediated largely at a transcriptional level. However, these findings do not exclude additional posttranscriptional effects of LCA. Expression of Sult2a1 was also induced in LCA-fed mice (Table 2), indicating that LCA-fed mice may, at least in part, metabolize LCA by sulfation.

Discussion

The emerging interest in the molecular mechanisms of bile acid transport and detoxification has resulted in several studies on LCA-induced liver injury in knockout mice for nuclear receptors and their respective target genes.10-15 These studies have mainly focused on specific molecular aspects of LCA detoxification, whereas the cholestatic phenotype induced by LCA remains functionally and morphologically poorly understood.29,30 Moreover, the critical question arises of whether feeding
Figure 13. Bsep expression is preserved throughout the liver lobule, whereas Mrp2 expression is restricted to acinar zone 3 in LCA-fed mouse liver. Double-immunofluorescence labeling of CKs (green; A, C, D, F, G, I, and L) and Bsep (B, C, E, and F) or Mrp2 (H, I, K, and L) (red) in 1% (w/w) LCA-fed (for 4 days) mouse liver. B and C: Bsep expression is preserved throughout the liver lobule in LCA-fed mouse liver. H and I: In contrast, Mrp2 expression is reduced to hepatic acinar zone 3. A and G: Note also the increased density of the CK-IF network in acinar zone 1 (indicated by the arrowheads). E, F, K, and L: Note that there is no evidence for canicular retrieval or targeting defect for Bsep and Mrp2 in LCA-fed mouse liver. bd, bile duct. Bar for A–C and G–I = 100 μm. Bar for D–F and J–L = 5 μm.
of a toxic bile acid is capable of inducing bile duct injury as the primary mechanism of hepatotoxicity. We herein show that feeding hydrophobic LCA to mice induces partial bile duct obstruction, bile infarcts, destructive cholangitis, and periductal fibrosis within days (Figure 14).

What Is the Pathogenesis of Cholestasis in LCA-Fed Mice?

In the past, many pathogenetic mechanisms of cholestasis in LCA-treated rodents (ie, mainly rats) have been proposed,3,5-9 and most of these studies have concluded that LCA-induced cholestasis is canalicular in origin. We herein demonstrate that cholestasis in LCA-fed mouse liver results from partial obstruction of bile ducts by crystals (presumably LCA precipitates). This novel conclusion is also supported by the described differential distribution pattern of biliary-secreted fluorescent UDCA and India ink, injected into the common bile duct in LCA-fed and CBDL mice. In addition, we found that Bsep expression was preserved and there was no evidence for canalicular retrieval of this transporter. Mrp2 tissue protein levels were unchanged, but its localization was limited to acinar zone 3, which could explain reduced biliary glutathione excretion in our model. Moreover, expression of Ntcp and Oatp1 was significantly reduced whereas Mrp3 (accompanied by Sult2a1 overexpression) was induced in response to LCA feeding, suggesting that these findings reflect an adaptive response protecting hepatocytes from ongoing hepatocellular accumulation of bile acids as previously observed in CBDL mice.22 Although it cannot be entirely excluded that these transporter alterations may be directly caused by LCA, the findings of the current study are consistent with the alterations of hepatic transport systems observed previously in mice and rats with obstructive cholestasis due to common bile duct ligation.22,31-33 Thus, it is unlikely that the transporter alterations observed in the current study are the primary cause for cholestasis in LCA-fed mice but rather may represent a secondary adaptive response to the cholestatic phenotype.

Why Do LCA-Fed Mice Develop Bile Infarcts?

Bile infarcts defined as confluent perportal liver cell necrosis with retained pigment material are a characteristic type of liver cell damage in response to obstructive cholestasis in humans and related animal models (eg, CBDL in rodents).26 We have previously shown that maintained bile secretion and increased biliary pressure are critical determinants for the development of bile infarcts in CBDL mice and that these lesions result from rupture of the canals of Hering,18,22 with exposure of hepatocytes to bile acid concentrations in the millimolar range derived from bile leaking into the liver parenchyma.34 The development of bile infarcts in LCA-fed mice (recently also described in intraperitoneally treated mice10,11) resulting from leaky, disrupted cholangioles (canals of Hering) may have previously been misinterpreted as “hemorrhagic cysts resembling peliosis hepatis” by Bagheri et al23 in LCA-treated rats. Our concept is confirmed by the finding of LCA crystals within bile infarcts. This hypothesis is further supported by the fact that most bile infarcts in LCA-treated mice remained unstained by India ink injected retrogradely into the common bile duct (which contrasts the findings in CBDL mice). On the other hand, biliary (orthogradely) excreted fluorescent UDCA accumulated in bile infarcts as observed in LCA-fed animals. Both observations may primarily be related to proximal bile duct obstruction in LCA-fed mice. In addition, previous studies by Benedetti et al35 also suggested an obstructive component of LCA-induced cholestasis, at least in the isolated perfused rat liver.

Why Do LCA-Fed Mice Develop Destructive Cholangitis?

Because bile infarcts and destructive cholangitis are the morphological hallmarks of the cholestatic phenotype in LCA-fed mouse liver, it is most likely that the bile ducts represent an important Achilles’ heel in regard to the hepatotoxic potential of bile acids in mice analogous to our previous findings in Mdr2<sup>−/−</sup> mice.18,19 LCA appears to provoke cholangitis through its physicochemical properties, ie, high hydrophobicity and lithogenicity. The perception that bile duct obstruction in LCA-fed mice may be caused by LCA precipitates is also supported by the fact that bile duct stones previously observed in chronically LCA-fed rats primarily consist of LCA.2 Because we observed intraductal LCA crystals damaging BECs and
finally resulting in ulcerations, it is reasonable to assume that cholangitis is directly related to the local toxic effects of LCA in mouse liver.\textsuperscript{3,6,26} However, it cannot be excluded from the presented data that an LCA-induced inflammatory response of BECs could additionally perpetuate cholangitis and proliferation of periductal myofibroblasts resulting in periductal fibrosis (eg, via inducing cytokine/chemokine production in BECs).\textsuperscript{16} In addition, based on the recent observation of toxic bile acid-induced hepatic stellate cell proliferation by Svegliati-Baroni et al.\textsuperscript{37} it is attractive to speculate that the observed proliferation of periductal myofibroblasts in LCA-fed mice could also be directly related to bile acids. The demonstrated induction of collagen 1a1, together with the greater induction of TIMP-1 in relation to MMP-2, suggests that liver fibrosis in LCA-fed mice could result, at least in part, from increased matrix production in the presence of decreased matrix degradation. Because the prolonged feeding of 1\% LCA-supplemented diet resulted in a detrimental outcome, future studies will have to determine whether alternative feeding protocols (eg, alternating periods of LCA and standard-diet feeding) are also able of producing parenchymal liver fibrosis and ultimately cirrhosis.

\section*{Do These Experimental Findings Have Potential Implications for Human Liver Disease?}

The novel finding that LCA feeding in mice induces destructive cholangiitis supports the concept that “toxic” bile is able to induce cholangiitis and consequently periductal fibrosis.\textsuperscript{19} Despite the fact that there is no proof for a causative role of LCA in human cholestatic liver diseases, LCA has been used since the early 1960s in experimental models to study cholestasis. However, because we observed crystals in cholangioles, bile ducts, and bile infarcts detected by electron microscopy, such crystals may have been missed in previous studies in fixed liver samples (ie, solubilization of LCA crystals during formalin fixation). Therefore, it may also be worthwhile to perform ultrastructural studies of bile ducts and their contents in human liver samples (eg, liver explants).

In summary, we provide evidence that the cholestatic phenotype in LCA-fed mouse liver is the result of LCA-induced obstruction of bile ducts and suppurrative cholangitis leading to periductal fibrosis (Figure 14). This is accompanied by an adaptive transporter and enzyme response. This information should be valuable for understanding LCA-treated mice as a model system for cholestatic liver diseases.

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\section*{References}