Disrupted Murine Gut to Human Liver Signaling Alters Bile Acid Homeostasis in Humanized Mouse Liver Models

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Abbreviations:
α, β, and ωMCA, α-, β-, and ω-muricholic acids; Abca1/ABCA1, Abcg5/ABCG5, Abcg8/ABCG8, murine/human ATP-binding cassette efflux cholesterol transporter; ACN, acetonitrile; ALT, alanine aminotransferase; Asbt, apical sodium dependent bile acid transporter; Baat/BAAT, murine/human Bile acid CoA:amino acid N-acyltransferase; Bcrp/BCRP, murine/human breast cancer resistance protein; Bsep/BSEP, murine/human bile acid export pump; CA, cholic acid; Car/CAR, murine/human constitutive androstane receptor; CDCA, chenodeoxycholic acid; CV, coefficient of variation; Cxcl16, chemokine (C-X-C motif) ligand 16; Cyp/CYP, murine/human cytochrome P450 enzyme; DCA, deoxycholic acid; ddH2O, double distilled water; ELISA, enzyme-linked immunosorbent assay; ESI, electrospray; Fah, fumarylacetoacetate hydrolase; Fgf15/FGF19, rodent or human fibroblast growth factor 15/19; Fgfr4/FGFR4, murine/human fibroblast growth factor receptor 4; FRGN, triple knockout of Fah, Rag2, and Il2rg genes on the non-obese diabetic strain background; Fxr/FXR, murine/human farnesoid X receptor; gCA, glycocholic acid; gCDCA, glycochenodeoxycholic acid; gDCA, glycodeoxycholic acid; gLCA, glycolithocholic acid; gUDCA, glycoursodeoxycholic acid; GAPDH, human glyceraldehyde 3-phosphate dehydrogenase; Gsta/GSTA, murine/human glutathione S-transferase; HepG2, human hepatocellular carcinoma cell; hFRGN/h-chimeric, humanized liver mouse or FRGN mouse with human hepatocytes repopulated liver; Hmg CoA Reductase/HMG CoA Reductase, murine/human 3-hydroxy-3-methyl-glutaryl-CoA reductase; Hnf-1α/HNF-1α, murine/human hepatocyte nuclear factor 1 alpha; Hnf-4α/HNF-4α, murine/human hepatocyte nuclear factor 4 alpha; HPLC, high pressure lipid chromatography; Ibabp, mouse ileum bile acid binding protein; IL-2rg, Interleukin 2 Receptor Subunit Gamma; LC/MS/MS, liquid chromatography-tandem mass spectrometry; LCA, lithocholic acid; Lrh-
1/LRH-1, murine/human liver receptor homolog 1; Lxrα/LXR, murine/human liver X receptor alpha; MCA, muricholic acid; Mdr1a/MDR1, murine multidrug resistance protein 1a or human multidrug resistance protein 1; MeOH, methanol; mFRGN, FRGN mouse with foreign mouse hepatocytes [Rosa (129S7) strain] repopulated liver; Mrp/MDR, murine/human multidrug resistance associated protein; MRM, multiple reaction monitoring; NH4OH; ammonium hydroxide; NTBC, nitisinone or 2-(2-nitro-4-trifluoro-methylbenzoyl)-1,3-cyclohexanedione; Ntcp/NTCP, murine/human sodium-taurocholate cotransporting polypeptide; Oat, mouse organic anion transporter; Oatp/OATP, murine/human organic anion-transporting polypeptide; Ost-α/OST-α, murine/human organic solute transporter α; Ost-β/OST-β, murine/human organic solute transporter β; PBS, phosphate buffer saline; PE tubing, polyethylene tubing; PepT1, mouse oligopeptide transporter; PPARγ, Peroxisome proliferator-activated receptor gamma; Pxr/PXR/SXR, murine/human pregnane X receptor, also known as steroid X receptor; PVDF, polyvinylidene fluoride; qPCR, quantitative real-time PCR; Rag2, recombination activating gene 2; Rosa, mouse hepatocytes derived from 129S7 mouse strain; SMRM, scheduled multiple reactions monitoring; SPE, solid phase extraction; Shp/SHP, murine/human small heterodimer partner; tα, tβ, and tωMCA, tauro-α-, tauro-β- acid, tauro-ω-muricholic acid; tCA, taurocholic acid; tCDCA, taurochenodeoxycholic acid; tDCA, taurodeoxycholic acid; TGF-β1/TGF-β, transforming growth factor beta; TGFBR2, transforming growth factor beta receptor 2; Tgr5, plasma membrane-bound, G protein-coupled receptor for bile acids; tLCA, tauroliothocholic acid; tUDCA, taourursodeoxycholic acid; UDCA, ursodeoxycholic acid; Ugt/UGT, murine/human UDP-glucuronosyltransferases; Vegf-a and Vegf-c, vascular endothelial growth factor a and c
ABSTRACT

The humanized liver mouse model is increasingly being exploited for human drug metabolism studies. However, its model stability, inter-communication between human hepatocytes and mouse non-parenchymal cells in liver and murine intestine, and changes in extrahepatic transporter and enzyme expression has not been investigated. We examined these issues in FRGN [Fah(-/-), Rag2(-/-), and IL-2rg(-/-) on NOD background] and chimeric mice: mFRGN, and hFRGN (repopulated with mouse or human hepatocytes, respectively). hFRGN mice showed markedly higher levels of liver cholesterol, biliary bilirubin, and bile acids (liver, bile, and plasma; mainly human forms, but also murine bile acids), but lower TGFBR2 mRNA expression (10%) in human hepatocytes and other proliferative markers in mouse non-parenchymal cells (Tgf-β1) and cholangiocytes (Tgr5), suggestive of irregular regeneration processes in hFRGN livers. Changes in the murine intestine, kidney, and brain of hFRGN mice: in particular, induction of intestinal Fxr genes: Fgf15, Ibabp, Shp, and Ost-α, were observed. Proteomics revealed persistence of remnant murine proteins (Cyp7a1, other enzymes, and transporters) in hFRGN livers and likelihood of mouse activity. When compared to normal human liver tissues, hFRGN livers showed lower SHP mRNA and higher CYP7A1 (300%) protein expression, consequences of βMCA- and αMCA-mediated inhibition of the FXR-SHP cascade and miscommunication between intestinal Fgf15 and human liver FGFR4, as confirmed by the unchanged hepatic pERK/total ERK ratio. Dysregulation of hepatocyte proliferation and bile acid homeostasis in hFRGN livers led to hepatotoxicity, gallbladder distension, liver deformity, and other extrahepatic changes, questioning use of the preparation for drug metabolism studies.
INTRODUCTION

Humanized (h-chimeric) liver mouse models are preclinical tools that are used for the prediction of human drug metabolism (Sanoh and Ohta, 2014). Unlike the transgenic mouse models, h-chimeric liver mouse models that are repopulated with human hepatocytes consist of the full complementary array of human liver genes. The triple genetic knockout (Fah−/−, Rag2−/− and Il2rg−/− on non-obese diabetic strain background) FRGN mouse model is one of these models where the absence of Fah gene results in liver accumulation of the toxic metabolite, fumarylacetoacetate, that induces hepatic injury and allows for repopulation of foreign hepatocytes (Azuma et al., 2007). Toxicity is alleviated upon giving these mice daily supplements of 2-(2-nitro-4-trifluoro-methylbenzoyl)-1,3-cyclohexanedione (NTBC), which blocks fumarylacetoacetate production and allows for a controllable environment to achieve a series of hepatocyte implantation for successful engraftment.

The h-chimeric liver mouse models are increasingly being used for human drug metabolism studies (Katoh et al., 2004; Tateno et al., 2004; Katoh et al., 2005a; Katoh et al., 2005b; Katoh et al., 2007; Ohtsuki et al., 2014) and have provided preliminary insights on human metabolite formation in vivo (Liu et al., 2011; Sanoh et al., 2012; Bateman et al., 2014; Kitamura and Sugihara, 2014). These mice respond to selective induction of ligands for human specific nuclear receptors (Katoh et al., 2005a; Emoto et al., 2008; Sanoh and Ohta, 2014). In addition, h-chimeric liver mouse models have been used to study pre-clinical drug-interaction (Jaiswal et al., 2014) and drug-induced liver injury studies for troglitazone (Barnes et al., 2014; Samuelsson et al., 2014), bosentan (Xu et al., 2015), and fialuridine (Xu et al., 2014), and have provided an essential in vivo safety testing tool for potential human toxic metabolites (Strom et al., 2010; Cohen, 2014; Kitamura and Sugihara, 2014; Xu et al., 2015).
A sound comparison of the pharmacokinetic and metabolic data with \textit{h}-chimeric liver mouse models is dependent on the premise that the model is stable, with little or no change in transporters and enzymes in extrahepatic tissues that are also under regulation by nuclear receptors (Makishima, 2005). The farnesoid X receptor (FXR), liver X receptor (LXR), pregnane/steroid X receptor (PXR/SXR), and constitutive androstane receptor (CAR) are major nuclear receptors that are responsible for the regulation of transporters and enzymes in the body. Although these nuclear receptors in human and other species share common goals, species differences in nuclear receptor activation exist, and this will pose as a major concern in ligand-specific nuclear receptor activation (Chiang et al., 2001; Handschin and Meyer, 2003; Katoh et al., 2005a; Sayin et al., 2013).

We recently observed remarkable physiological and metabolic zonation differences between hFRGN and FRGN mouse livers. Notably, hFRGN livers contain remnant mouse transporters and enzymes that are present at similar or even higher levels than those in FRGN livers (Chow et al., 2016). One possible explanation is that changes in intrinsic factors such as hormone and bile acid productions, as regulated by intestinal and hepatic nuclear receptors, will lead to changes in liver transporter and enzyme levels. At high bile acid concentrations, human cytochrome P450 \textit{\textit{7a}}-hydroxylase (CYP7A1), the rate-limiting enzyme for bile acid synthesis, is negatively regulated by the FXR-SHP cascade (Chiang, 2003; Chiang, 2009) and by the fibroblast growth factor 15/19 (Fgf15, human ortholog, FGF19), a hormone that is secreted by the ileum and binds to liver FGFR4 (receptor 4) to activate the \textit{c}-Fos pathway for repressing CYP7A1 in a negative feedback mechanism (Inagaki et al., 2005; Lin et al., 2007). In \textit{h}-chimeric livers, CYP7A1 and human bile acid levels were observed to be dramatically elevated, but the
condition was corrected upon administration of exogenous FGF19 or transfection of FGF19 gene in h-chimeric mice (Ellis et al., 2013; Naugler et al., 2015).

In order to fully understand bile acid dysregulation in these h-chimeric mice, we examined individual mouse and human bile acid species by liquid chromatography coupled with tandem mass spectrometry (LC-MS/MS) and employed proteomics to quantify changes in mouse and human CYP7A1 and other proteins. We included mFRGN mice (FRGN liver with foreign mouse hepatocytes repopulation) as another control to ensure that observations are independent of the diet/housing and surgical manipulation (Chow et al., 2016). Because the status of each organ/tissue could potentially impact the functionality of others (Naud et al., 2007; Naud et al., 2008), we appraised key biological, physiological parameters, and human and mouse gene expression in liver, including proliferative genes in non-parenchymal cells and in extrahepatic tissues to fully understand the inter-organ communication between human and mouse organs in hFRGN mice.
METHODS

Materials. Cholic acid (CA), chenodeoxycholic acid (CDCA), deoxycholic acid (DCA), lithocholic acid (LCA), ursodeoxycholic acid (UDCA), taurocholic acid (tCA) sodium salt hydrate, taurochenodeoxycholic acid (tCDCA) sodium salt, taurodeoxycholic acid (tDCA) sodium salt, and taurlithocholic acid (tLCA) sodium salt were obtained from Sigma Aldrich (Oakville, ON). Muricholic acid (MCA), α-, β-, and ω-muricholic acids (α, β, and ωMCA), tauro-α-, tauro-β- acid, tauro-ω-muricholic acid (tα-, tβ-, and tω-MCA) sodium salts, and taouroursodeoxycholic acid (tUDCA) sodium salt were obtained from Steraloids (Newport, RI). Glycocholic acid (gCA), glycochenodeoxycholic acid (gCDCA), glycodeoxycholic acid (gDCA), glycolithocholic acid (gLCA), and glycoursodeoxycholic acid (gUDCA) were kind gifts from Dr. Alan F. Hofmann (University of California, San Diego). Cholic-2,2,4,4-d₄ acid (CA-d₄), chenodeoxycholic-2,2,4,4-d₄ acid (CDCA-d₄), deoxycholic-2,2,4,4-d₄ acid (DCA-d₄), and lithocholic-2,2,4,4-d₄ acid (LCA-d₄) were purchased from CDN isotopes (Pointe-Claire, QC).

Mice and housing conditions: Male FRGN, mFRGN [FRGN livers repopulated with pooled mouse hepatocytes of the Rosa (129S7) mouse strain] and hFRGN [FRGN livers with hepatocytes repopulation] mice (4-6 months old from three different human donors of ages 5, 7 and 17, Table 1) were supplied by Yecuris Corporation (Tualatin, OR). All hFRGN livers were >80% human repopulated, as stated by Yecuris. The housing and diet conditions of FRGN, mFRGN, and hFRGN mice have been previously described in detail by Chow et al. (2016). All animal studies were performed in accordance to approved animal protocols at the University of Toronto.
**Biological Assays:** Plasma alanine transaminase (ALT) from fresh plasma samples were immediately quantified by ALT kit (Bioquant, Nashville, TN), and plasma and liver cholesterol and liver triglyceride levels were measured using total cholesterol (Wako Diagnostics, Richmond, VA and Thermo Scientific, Rockford, IL, respectively) and triglyceride (Thermo Scientific, Rockford, IL) kits as previously described by Chow et al. (2014). Total and conjugated bilirubin in bile were assayed by the bilirubin assay kit (Sigma Aldrich Canada) after a 15-fold dilution with saline.

**Oil Red O Staining:** After flushing of the mouse liver with ice-cold saline, livers were fixed with 10% formalin overnight for Haemotoxylin and Eosin staining, followed by Oil Red O staining, as performed by the Toronto Centre for Phenogenomics (Toronto, ON).

**Bile acid quantification from bile, plasma, liver and pool size using LC-MS/MS:** The bile ducts of FRGN, mFRGN, and hFRGN mice were cannulated after gallbladder ligation under anesthesia with PE10 tubing for continuous bile collection *in vivo* for 10-20 min. Blood was then collected by cardiac puncture with a heparinized needle, and the sample was centrifuged to obtain plasma. Ice-cold saline was pushed through the vena cava to flush blood out of tissues. Enterocytes from intestine [duodenal, proximal jejunal and ileal segments, with enterocytes removed by a scrapper as described (Chow et al., 2014)], kidney, liver, and brain were collected, cut into pieces, snapped frozen in liquid nitrogen, and stored at -80°C freezer for future analyses.

Detailed description of sample preparation for bile acids in plasma, bile, and liver tissue, and the bile acid pool size study was as follows. For bile acid sample preparation, the aliquot was first diluted 100-fold with ddH₂O, and added 20 μL MeOH:ddH₂O (50:50 v/v) containing a mixture of the internal standards [1 μg/mL of CDCA-d₄, CA-d₄, DCA-d₄, and LCA-d₄ (C/D/N Isotopes, Pointe-Claire, Canada)]. The resultant mixture was added to the pre-conditioned SPE
(Strata-X) column (2 mL MeOH, followed by 2 mL ddH2O). The column was then eluted with 2 mL ddH2O for clearing unwanted debris. Outflow collection was initiated upon elution with 4 mL MeOH. The eluent was dried under nitrogen gas, reconstituted with 100 µL MeOH:ddH2O mixture, vortexed for 1 min, and centrifuged at 12,000 g for 10 min at room temperature prior to injection. For plasma sample preparation, 100 µL of sample was added 20 µL of the internal standard mixture and precipitated with 1 mL ice-cold acetonitrile (ACN) containing 5% 1N NH4OH. The mixture was vortexed for 1 min, centrifuge at 12,000 g for 10 min at 4°C, and 1 mL supernatant was pipetted and dried under nitrogen gas. The residue was reconstituted with 100 µL MeOH:ddH2O mixture, vortexed for 1 min and centrifuged at 12,000 g for 10 min at room temperature prior to injection. For liver tissue preparation, the liver was homogenized with ddH2O (1:2, w:v) on ice. Then, 100 µL homogenate was added with 20 µL of the internal standard mixture and precipitated with 1 mL ice-cold ACN (in 5% 1N NH4OH). The mixture was vortexed for 1 min, centrifuged at 12,000 g for 10 min at 4°C, and 1 mL supernatant was pipetted and dried under nitrogen gas. The residual homogenate mixture was re-precipitated with another 1 mL of ice-cold ACN (in 5% 1N NH4OH), vortexed, spun, collected in the same collection tube, and dried under nitrogen gas. The residue was reconstituted with 100 µL MeOH:ddH2O mixture, vortexed for 1 min and centrifuged at 12,000 g for 10 min at room temperature prior to injection.

For preparation of the calibration curves for bile acids in each of the biological matrix, blank sample matrix was first prepared. Blank bile (diluted 100-fold with ddH2O), plasma, and liver homogenates (1:2, w:v with saline) were incubated with charcoal (100 mg/mL final concentration) and the sample was kept shaking overnight at 4°C. The resulting solution was spun at 9,000 g for 10 min, and the supernatant was filtered through a 0.45 µm pore size Supor
polyethersulfone membrane syringe filter (Pall Life Sciences). When these blank sample matrices were tested for the presence of bile acids with LC-MS/MS, none was found (below limit of quantification). Hence, for standard curves, bile, plasma, and liver homogenate tissue were first stripped of endogenous bile acids, then the appropriate standards were added (0.01-10 µg/mL final concentration for each bile acid species) and processed in the same manner as described above.

Bile acid extraction for bile acid pool size. The bile acid extraction procedure was similar to that previously described by Chow et al. (2014). On the last day of treatment with sterile water, mice were fasted for 4 h (from 9 a.m. to 1 p.m.). The intact gallbladder, liver, and intestine were removed together under anesthesia. Tissues were minced into pieces in a beaker containing 50 mL of anhydrous ethanol added with 50 µL of the internal standards [a mixture of 0.25 mg/mL of CDCA-d₄, CA-d₄, DCA-d₄, and LCA-d₄]. The content was boiled at 80°C (ethanol boiling point) for 1 h, and ethanol was added during the heating/incubation to replace evaporated liquid. After cooling, the extracts were filtered through Whatman filter paper and adjusted to 50 mL with ethanol in a volumetric flask. Some extracts were centrifuged at 10,000 g for 10 min and filtered through an Ultra-free-MC centrifugal filter device containing 0.22 µm PVDF membrane (Millipore, Billerica, MA) prior to analysis. Bile acid standards, prepared in different concentrations (0.1-50 µM), were extracted in a similar manner.

LC-MS/MS for bile acid quantification. Extracted samples from bile, plasma, liver homogenate, and bile acid pool size were analyzed by LC-MS/MS using AB Sciex API 4000 Triple Quad LC/MS instrument (Applied Biosystems, ON) with ESI source in negative ion mode. Samples (10 µL) were injected and separated by a Kinetex 2.6µm C18 100A 100 x 4.6 mm column (Phenomenex Inc.), with a SecurityGuard pre-column (Phenomenex Inc.) at 600 µL/min flow
rate. The mobile phase consisted of pre-filtered 10 mM ammonium acetate (A) and HPLC grade ACN (B). A gradient was utilized over 20 min: 0-6 min, 35-35% solvent B; 6-14 min, 35-58% solvent B; 14-15 min, 58-95% solvent B; 15-16.5 min, 95-95% solvent B; 16.5-17 min, 95-35% solvent B; 17-20 min, 35-35% solvent B. MS parameters are listed in Table 2, and a gas temperature of 500°C, ion spray voltage of 3500V, and column ambient temperature were used. Selective ion monitoring was employed to detect the conjugated and unconjugated bile acids. Bile acids were quantified based on area of the peak of the standard, corrected by area of the appropriate deuterated internal standard, in calibration curves (Table 2).

**FGF19 in plasma, liver, bile and gallbladder.** Minced, frozen hFRGN liver tissues were homogenized in PBS (10 mM, pH 7.0; 1:1, w:v) over ice. The resultant homogenate was sonicated with a cell disrupter and then centrifuged at 5000 g for 5 min to provide a supernatant for later analysis. Plasma (diluted 2-fold), liver supernatant (diluted 5-fold), bile (diluted 40-fold) and gallbladder (diluted 5-fold) samples were diluted with PBS and assayed using a human FGF19 ELISA kit (R&D Systems Inc., Minneapolis, MN), following the manufacturer’s protocol. We also attempted at quantifying Fgf15 levels using ELISA. However, these levels were not reported since the values were considered unreliable (personal communication with Dr. S.A. Kliewer).

**Proteomics.** Protein expression (Karlgren et al., 2012) was determined by peptide-based LC-MS/MS measurements (Vildhede et al., 2014). Quantification of proteins in hFRGN (n=9), FRGN (n=8) and human liver (n=3) tissues (see Table 1 for donor information) was accomplished by mass spectrometry-based targeted proteomics using validated LC–MS/MS methods, as previously described (Groer et al., 2013).
Tissue was homogenized by a Cellcrusher tissue pulverizer (Schull, Ireland). The peptide sequence and multiple reaction monitoring (MRM) transition for each peptide and isotopically-labeled peptide as internal standard for quantification are listed in Table 3. An isotope-labeled peptide was used as the internal standard for each peptide (Table 3). Protein was determined by the protein bicinchoninic acid kit from Pierce Biotechnology (Rockford, IL). An aliquot of 200 μg membrane protein was digested with Mass Spec Grade Trypsin/Lys-C Mix (Promega, Madison, WI) (Qiu et al., 2013). Membrane fractions were extracted using ProteoExtract Native Membrane Extraction Kit (Calbiochem, San Diego, CA) according to the manufacturer’s protocol, and 20 μL of the reconstituted, digested sample was injected into Shimadzu LC system (LC-30A) coupled with ABSciex 6500 QTrap equipped with Turbo Spray ion source operating at positive-ion mode. All chromatographic separations were performed by gradient elution with a Waters Acquity UPLC Peptide BEH C18 130A 1.7 μm 2.1x150 mm column, maintained at 60°C at a flow rate of 300 μL/min. The gradient program started as 10% mobile phase A (0.1% formic acid in water), increased to 30% B (0.1% formic acid in ACN) over 25 min, followed by a sharp increase to 90% B in the next 0.5 min, then maintain at 90% B for 2 min. The gradient was then decreased to 10% B in 0.2 min, and held for 2 min. The mass spectrometer ion spray voltage was 5000V; temperature was 450°C; curtain gas was at 40 psi; and ion source gas 1 and ion source gas 2 were at 85 and 60 psi, respectively. Detection was accomplished by mass spectrometry-based targeted proteomics using validated LC−MS/MS methods, as previously described (Groer et al., 2013). Analyses were performed in scheduled multiple reactions monitoring (SMRM) mode. Peak integration and quantification were performed using the Analyst 1.6.2 software. The final protein expression (fmol per μg membrane protein) was calculated upon normalization of
the isolated membrane fraction as described by Li et al. (2008) and the accuracy (error) and precision (CV) were <20%. Each sample was analyzed in duplicate (technical repeats).

**Real-Time quantitative PCR (qPCR):** Total mRNA in tissue was isolated with the standard TRIzol extraction procedure and quantified as previously described (Chow et al., 2009; Chow et al., 2014). Specific mouse and human primers were designed in Primer-BLAST (http://www.ncbi.nlm.nih.gov/tools/primer-blast/) with exon-exon junction span and specificity check for appropriate species (Chow et al., 2016). Primers were checked against mouse liver and human hepatocellular carcinoma cell (HepG2) or human liver tissue mRNA and water for specificity, and the sequences are summarized in Supplementary Table 1. mRNA was synthesized into cDNA by a High Capacity cDNA synthesis kit; cDNA was quantified by the Applied Biosystems 7500 series system using SYBR Green or Taqman for detection. For human liver mRNA analysis, genes were normalized to human GAPDH (detects human). For mouse liver mRNA analysis, genes for mouse hepatocytes and non-parenchymal cells were normalized to mouse β-actin (detects mouse), and genes for cholangiocytes were normalized to mouse CK19 (detects mouse), a marker for cholangiocytes. Target gene data in mouse tissue was normalized to cyclophilin for brain, and kidney, and to villin for the small intestine.

**Western blotting:** Protein isolation was performed as previously described (Chow et al., 2009). Liver homogenate protein samples (25 μg) were loaded and separated by 12% SDS-polyacrylamide gels, transferred onto nitrocellulose membranes (Amersham Biosciences, Piscataway, NJ), then blocked with 5% (w/v) skim milk in Tris-buffered saline and 0.1% Tween 20 (TBS-T; Sigma Aldrich Canada) for 1 h at room temperature, followed by washes with 0.1% TBS-T. Primary antibody solutions of total ERK (1:5000 dilution) (Cell Signaling Technology, Danvers, MA; cat#9102) and pERK (1:1000 dilution) (Cell Signaling Technology, Danvers,
MA; cat#9101) or GAPDH/Gapdh (1:10000 dilution) (Abcam, Cambridge, MA; Cat#ab8245) were prepared in 5% skim milk in 0.1% TBS-T and were incubated overnight at 4°C. On the next day, the membrane was washed with 0.1% TBS-T and then incubated with secondary antibody (1:2000 anti-rabbit or 1:10000 anti-mouse) in 2% skim milk in 0.1% TBS-T for 2 h at room temperature, and then washed with 0.1% TBS-T. Bands were visualized with chemiluminescence reagents (Amersham Biosciences) and quantified by scanning densitometry. Two bands appeared for pERK and total ERK. The intensity for each of the 44 and 42 kDa band was normalized individually to GAPDH/Gapdh (37 kDa) and summed for pERK or total ERK. The ratio of pERK/total ERK was then calculated.

**Statistics:** Data were expressed as mean ± SEM for all data. For comparison between two groups, the 2-tailed paired Student’s *t*-test was used. The value of *P*<0.05 was set as the level of significance.
RESULTS

Characteristics of FRGN, mFRGN, and hFRGN mice. Liver weights of hFRGN mice were significantly greater than those of FRGN mice, although the liver and brain weights of mFRGN mice were smaller than those of FRGN mice (Table 4). The gross appearance of hFRGN livers appeared necrotic and consisted of multiple smaller lobes with distended gallbladders from that of FRGN livers (Figure 1A). mFRGN livers appeared abnormal and different, but less frequent. All three mouse species showed higher than baseline plasma ALT levels due to the withdrawal of NTBC (the supplement that rescues the liver from toxicity due to absence of Fah gene) 3 days prior to experimentation. For the FRGN mouse, NTBC withdrawal resulted in some degree of liver damage, shown by the slightly elevated ALT levels. For the mFRGN mice, the ALT level was close to baseline values (~20 IU/ml), suggesting that the foreign mouse hepatocytes were able to replace damaged naïve mouse hepatocytes and restore liver function and lessen toxicity. For the hFRGN mice, the ALT level was the highest, relative to those in FRGN and mFRGN mice. The reason may be the presence of excessive bile acids (discussed later) found in the hFRGN system. Moreover, hFRGN mice contained much higher levels of liver cholesterol and conjugated bilirubin than those for FRGN mice (Table 4). Hepatic triglyceride levels in mFRGN livers (7.36 ± 1.57 mg/g) were higher than those of FRGN and hFRGN livers, although levels in hFRGN livers (3.3±1.06 mg/g) were not significantly different from that in FRGN livers (1.81±0.62 mg/g; Table 4). These observations were consistent with histopathological images with lipid Oil Red O staining (Figure 1B).

Bile acid composition in bile, plasma, and liver and bile acid pool sizes in FRGN, mFRGN, and hFRGN mice. The composition of bile acids among FRGN, mFRGN, and hFRGN mice was compared (Figure 2A; Table 5). In FRGN and mFRGN mice, muricholic acids (αMCA,
βMCA, and ωMCA) and their taurine-conjugates, as well as taurocholic acid (tCA) were major murine bile acids present in all biological matrices (Figure 2A; Table 5). Overall, implantation of foreign mouse hepatocytes into FRGN recipients did not greatly alter the bile acid composition in all biological matrices (Figure 2A), although bile acid concentrations in mFRGN mice were about 1.5 to 3.3-fold higher than those in FRGN mice (Table 5). For hFRGN mice, dramatic changes in bile acid composition and dramatically higher (by several to a few thousand-fold) bile acid levels than those of FRGN mice were observed in bile, plasma, and liver (Figure 2A, Table 5). There were considerably greater glycine-conjugates, as expected of human bile acids, although taurine conjugates also observed. Muricholic acid and the variants of murine designation, persisted in hFRGN plasma, bile and liver. These differences in bile acid composition may be contributed by many external factors. For the comparison in humans, it must be mindful that the human hepatocytes repopulated in these hFRGN livers were from either male or female donors, and gender difference in bile acid metabolism can occur (Fisher and Yousef, 1973), but humanization of these livers originating from both male and female donors were indeed highly variable (Chow et al., 2016). In addition, gut microbiota are different between mice and humans, and this may contribute further to differences in secondary bile acid production (Wahlstrom et al., 2016).

**Bile.** In hFRGN bile, CA, CDCA, and DCA and their taurine- and glycine-conjugates, major forms found in humans, were observed to be significantly higher when compared to those in FRGN bile (Table 5); tUDCA, gUDCA, tLCA, gLCA, αMCA and tαMCA were also considerably higher in hFRGN bile. A detailed comparison of the bile contents for hFRGN (including FRGN and mFRGN bile) and human bile was further conducted (Table 6). Even though levels of the glycine-conjugates of CA, DCA, CDCA, and UDCA were present in
hFRGN bile, suggesting human metabolic activities, these concentrations were lower relative to those in human bile (Table 6). Moreover, higher proportions of taurine-conjugates as well as muricholic acids (10% total), rodent specific bile acids formed from Cyp2C70 (Takahashi et al., 2016), were observed in hFRGN bile, but were virtually absent in human bile. Species differences in the bile acid conjugation pathways between human and rodents raised serious questions about the contribution of murine hepatic activities in these highly humanized livers, inasmuch as bile acids are mostly glycinated in human livers but are taurine-conjugated in rodent livers (Sayin et al., 2013). The higher taurine to glycine conjugates in hFRGN mice suggests the presence of remnant mouse hepatocyte activities. In addition, βMCA and CA, which are normally conjugated with taurine in rodents (Sayin et al., 2013), were present abundantly as 58% of biliary bile acids in hFRGN mice, contrasting the 6.5% CA and 0% MCA observed for human bile (Table 6). These facts again suggest the persistence of murine hepatic activities.

**Plasma.** In plasma, higher levels of human bile acids were observed in hFRGN mice than those in FRGN mice, though those for MCA and tMCAs (murine bile acids) in hFRGN plasma were also higher (Table 5).

**Liver.** In liver, primary and secondary bile acids, such as MCAs and CDCA were significantly lower in hFRGN mice when compared to those in FRGN mice, but DCA and taurine- and glycine-conjugated bile acids were higher (Table 5). Rodent bile acids were higher in hFRGN livers: tαMCA, tβMCA and tωMCA concentrations in hFRGN livers were 29, 42 and 1.9-fold those in FRGN liver, respectively (Table 5).

**Bile acid pool sizes.** The extremely high levels of bile acids in bile, plasma and liver observed in hFRGN mice prompted us to compare the bile acid pool sizes between hFRGN and FRGN mice. The bile acid pool size for hFRGN mice (n=4) was found to be 4.2-fold that of FRGN mice.
(n=4) (Figure 2B). Human conjugated bile acids such as tCDCA, tDCA, tUDCA, tLCA, gCA, gCDCA, gDCA, gDCA were significantly higher in hFRGN mice, and the MCA content in hFRGN mice was proportionately lower, although not absent. Glyco-conjugates, tauro-conjugates, and DCA were almost 1000-fold (gCDCA, 2220-fold; gCA, 943-fold; gDCA, 1140-fold), 4.2-fold (tDCA, 26-fold; tCA, 6.3-fold), and 8.7-fold higher in hFRGN mice, respectively (Figure 2B).

Comparison of basal human mRNA expression between hFRGN livers and normal human liver tissues. hFRGN livers and human liver tissues were examined to compare basal human mRNA and other underlying changes. For examination of whether any of the bile acid synthetic pathways was altered, mouse and human enzymes in hFRGN livers and human liver tissues were measured (Figure 3). CYP7A1, or CYP 7α-hydroxylase, the rate limiting enzyme for the classical (major) pathway in bile acid synthesis, oxysterol 7α-hydroxylase or CYP7B1, the rate limiting enzyme in the alternative (acidic) pathway for CDCA production, and CYP27A1, the alternative enzyme for CDCA production in both the classical and alternative pathways, and BAAT (conjugative enzyme) (Chiang, 2009), were evaluated. As shown in Figure 3A, there was no difference in human hepatic CYP27A1, CYP7B1, and BAAT mRNA expression between hFRGN livers and human livers, except for human CYP7A1 mRNA expression, which was 120-fold higher in hFRGN over human liver tissue (Figure 3A). By contrast, mouse liver Cyp7b1, Cyp27a1, and Baat mRNA expression (Figure 3B) in hFRGN livers were much lower than those for FRGN and mFRGN mice.

The mRNA expression of human CAR, HNF-4α, PPARγ, albumin, β-Klotho, CYP2E1, GSTA4-4, UGT1A1, OATP1B1, OATP1B3, BSEP, ABCG5, ABCG8, OST-α–OST-β, MDR1, and MRP2 were also higher (1.5-2.5-fold) in hFRGN livers (Figure 3A), whereas those for FXR
and SHP, were lower in hFRGN livers than those of human liver tissues. One plausible reason for the lower FXR target gene expression is the presence of extremely high levels of taurine-conjugated MCAs (Table 5), FXR antagonists (Sayin et al., 2013); inhibition of FXR would lead to the reduction in SHP expression (Figure 3A). Those for CYP1A1, CYP2C19, BCRP, OST-α and OST-β mRNA levels in hFRGN livers were 2.5-fold higher, with a notable 100-fold higher CYP7A1 mRNA level in hFRGN livers when compared to human liver tissue (Figure 3A). Despite the high bile acid contents in hFRGN livers (Table 5), the mRNA expression of MRP3, bile acid-associated FXR target, was unexpectedly lower (Figure 3A).

**Miscommunication among human hepatocytes and murine nonparenchymal cells (mouse Kupffer cells, stellate cells, and cholangiocytes) in hFRGN vs. FRGN and mFRGN livers**

The communication between human hepatocytes and mouse non-parenchymal cells in hFRGN livers was evaluated to assess the proliferation status. Since the receptor II of TGF-β1, TGFBR2, in human hepatocytes and TGF-β1 in mouse that is produced by a multitude of non-parenchymal liver cells, including Kupffer cells, sinusoidal endothelial cells, dendritic cells and stellate cells (De Bleser et al., 1997; Schon and Weiskirchen, 2014; Weiskirchen and Tacke, 2014) are both antimitogenic, the termination signal was evaluated. TGFBR2 mRNA expression in hFRGN livers was significantly lower (only 10%) than that in normal human liver tissues (Figure 3A), suggesting continuous hepatocyte proliferation in hFRGN livers. In addition, mouse TGF-β1 mRNA level in hFRGN was not elevated compared to FRGN livers (Figure 3B), suggesting that mouse stellate/Kupffer cells failed to respond to the high number of human hepatocytes present in hFRGN livers (Yoshizato et al., 2012). These results suggest a lack of communication between mouse stellate cells and human hepatocytes.
To examine further whether proliferation of hepatocytes had influenced cholangiocyte growth and intrahepatic bile acid mass, Cxcl16 (Omenetti et al., 2009) and other vascular endothelial growth factors (Vegf-a and Vegf-c) (Glaser et al., 2014; Meng et al., 2014), key cell growth proliferators in cholangiocytes, were measured (Figure 3B). The results showed that these markers were significantly decreased, suggesting that murine cholangiocyte proliferation was inhibited. In addition, expression of a transmembrane G-protein coupled receptor (Tgr5) that is activated by bile acids (especially tLCA and tDCA) (Keitel and Haussinger, 2013; Duboc et al., 2014; Reich et al., 2016) and is localized in the cholangiocytes, modulating cholangiocyte proliferation and bile flow, was lower (Figure 3B). Asbt, regulated by cAMP and Tgr5 (Xia et al., 2006; Keitel and Haussinger, 2013) in cholangiocytes, was lower in hFRGN livers (Figure 3B). The results suggest a decrease in cholangiocyte proliferation and bile acid reuptake (Lazaridis et al., 1997). The expected consequence of downregulated Tgr5 in hFRGN livers is a reduction of the cholehepatic shunt, leading to higher liver and plasma bile acid concentration and the possibility of generating cystic fibrosis or primary sclerosing cholangitis characteristics (Glaser and Alpini, 2009; Maroni et al., 2015).

**Proteomics of human/murine liver protein abundances in hFRGN livers.** To investigate the potential role of bile acid dysregulation in the hFRGN liver, we performed proteomic analysis, the more quantitative tool to differentiate between human and mouse proteins in hFRGN livers, to evaluate the absolute transporter and enzyme changes (Figure 4). Similar to the mRNA data (Figure 3A), the protein expression of human CYP7A1, determined by LC-MS/MS, was higher (>3-fold) in hFRGN livers than that in human liver tissue (Figure 4A). The protein expression of mouse Cyp7a1, the rate-limiting enzyme in cholesterol metabolism, in hFRGN livers was only slightly lower when compared to that in FRGN livers, suggesting that the contribution of murine
enzyme for bile acid synthesis in hFRGN liver would not be negligible despite that Cyp27a1 and Cyp7b1 mRNA expression levels were much lower (Figure 4B). Remnant protein expression of murine Oatp1b2, the ortholog of human OATP1B1 and OATP1B3 (Evers and Chu, 2008), was also observed (Figure 4B). Due to a lack of species specificity in protein sequence (Ntcp/NTCP, Bcrp/BCRP, Bsep/BSEP, Mrp2/MRP2, Mrp3/MRP3, Mrp4/MRP4, and P-gp) of the peptides, a comparison between the amounts of human and mouse specific protein in hFRGN and FRGN livers could not be made in the proteomic analyses. Rather, the comparison was conducted using either mouse-specific or cross-reactive peptide for the target proteins in FRGN and hFRGN livers. In Figure 4C, protein contents in Ntcp/NCTP, Bcrp/BCRP were found to be lower for hFRGN livers than those for FRGN livers whereas levels for the ATP transporters (Mrp2/MRP2, Mrp3/MRP3 and Mrp4/MRP4 and P-gp) were all higher. Protein differences of Ntcp/NTCP, Mrp4/MRP4, and murine/human P-gp in FRGN and hFRGN livers were consistent to those protein levels previously measured with immunoblotting (Chow et al., 2016), which may also be reflective of cross-reactive peptides. In addition, these protein data were consistent with those for mRNA expression that were published (Chow et al., 2016).

Miscommunication between human hepatocytes and murine intestine

The potential Fgf15 and FGF19 signaling pathway on hepatic human CYP7A1 regulation in hFRGN mice was evaluated to assess the communication between murine intestine and human hepatocytes. Levels of FGF19 in liver, bile, gallbladder, and plasma, when assayed with ELISA, were found absent (data not shown). Levels of Fgf15 in plasma, though measured with ELISA, remained uncertain since the measured values are unreliable (personal communication with Dr. Steven A. Kliewer). Intestinal murine Fgf15 mRNA expression was therefore served as a biomarker of what the circulating Fgf15 levels would have been. In hFRGN intestine, Fgf15
expression was significantly induced (500-fold), likely by intestinal Fxr activation (Figure 5A), a line of reasoning that is supported by induction of all other intestinal Fxr targets such as Ost-α, Ibabp, and Shp, since high levels of bile acid in hFRGN mice were observed (Figure 2B). In addition, ileal Asbt expression (Figure 5A) was lower in hFRGN intestine, likely due to inhibition by the Fxr-Shp-Lrh-1 cascade (Chen et al., 2003) or inhibition by Fgf15 (Sinha et al., 2008). Moreover, analyses of phosphorylated ERK (pERK) and total ERK protein (Figure 5B), an indicator of FGFR4 activation in human hepatocytes, showed that both pERK and the ratio of pERK/total ERK were similar in FRGN and hFRGN livers, suggesting that the increase in Fgf15 did not result in higher activation of FGFR4/β-Klotho signaling pathway in human hepatocytes. As a result, CYP7A1 protein expression in hFRGN remained elevated compared to that for human liver tissue (Figure 4A), drastically increasing the bile acid pool size in hFRGN mice (Figure 2B).

**Changes in basal mRNA expression in other mouse tissue.** Changes in nuclear receptor, transporter, and enzyme expression in the mouse intestine, kidney, and brain were also evaluated among FRGN, mFRGN and hFRGN mice to assess the potential impact of bile acid dysregulation on inter-organ communication.

**Intestinal genes:** The basal mRNA levels of different intestinal genes were generally similar between FRGN and mFRGN intestine (Figure 6). However, levels were much different between FRGN and hFRGN intestine. For important xenobiotic intestinal genes, higher Car and Pxr levels as well as Mdr1a, Mrp4, Gsta4-4, Sult1a1, and Ugt1a1 were observed in hFRGN intestine (Figure 6). Fluctuations in intestinal Oatp2b1 and Gsta3-3 were found in mFRGN and hFRGN intestine, and slightly lower intestinal Mrp3 (all three segments) and Bcrp (ileal segment) were observed for the hFRGN intestine. The basis for these differences is currently unknown since
both higher LCA (a Pxr agonist) and human bile acids (strong Fxr agonist) co-existed in the intestine.

**Renal genes.** Only minor differences in renal genes between mFRGN and FRGN mice were observed (Supplementary Figure 1), and variations in Lxra, Pxr, Oapt1a1, Oatp1a4, Ost-β, Oat1, Oat3 and Sult1a1 expression were small. However, when the basal expression of FRGN and hFRGN kidneys was compared, greater changes were observed. Renal Pxr and Car levels were to be increased, and Lxra level decreased in hFRGN mice. In addition, renal mRNA expression of Oatp1a4, Ost-α, Ost-β, Mrp2, Mrp3, Mrp4, and Gsta4-4 were all elevated in hFRGN mice, though Vdr, Oapt1a1, Oat1, PepT2, and Cyp2e1 levels in hFRGN mice were lower as compared to those of FRGN mice. The data suggest that many of the observed changes could have been the results of Fxr- and Pxr-mediated regulation due to the high levels of bile acids.

**Brain genes.** Only minor changes in brain nuclear receptors and transporters were observed between mFRGN and FRGN mice (Supplementary Figure 2). Changes in transporter were small, although expression of Oapt1a4, Oatp2b1, and Mdr1a levels were increased. Levels of Cyp1a2, Cyp2e1, Cyp3a11, Gsta3-3, and Ugt1a1 were significantly lower in hFRGN brain when compared to those of FRGN brain.
DISCUSSION

Development of the h-chimeric mouse liver model has been hailed as a useful in vivo tool for the study of human liver metabolism. Many studies supported the presence of human enzyme and transporter expression in humanized mouse livers (Katoh et al., 2005a; Katoh et al., 2005c; Nishimura et al., 2005) while pharmacokinetic studies in humanized mice showed the presence of in vivo human metabolites (Okumura et al., 2007; Grompe and Strom, 2013; Bateman et al., 2014). However, other studies using chimeric models showed that the levels of in vivo human metabolites were unpredictable and different from that in humans (De Serres et al., 2011; Liu et al., 2011; Sanoh et al., 2012).

In our previous study, we have identified two key issues on discordance of the humanized model. We unveiled the presence of remnant native mouse hepatocytes and loss of metabolic zonation in hFRGN livers (Chow et al., 2016). In our present report, histopathologic, biological, mRNA, and proteomics (using LC-MS/MS) data provided us additional evidence to suggest instability of the model, including uncontrollable human hepatocyte proliferation and liver deformity (Figures 1 and 3), high bile acid dysregulation (Table 5; Figure 2) and toxicity (Table 4), enlarged gallbladder filling (Figure 1A), and significant changes in extrahepatic transporters and enzymes (Figures 5 and 6, Supplementary Figures 1 and 2). Although the model is increasingly being used as a tool for drug discovery and development to relate to human drug metabolism and toxicity, it is recognized that many of these deficiencies have been previously overlooked and not properly addressed.

Dysregulated liver regeneration and deformity in hFRGN livers was evidenced by the miscommunication between human hepatocytes and murine stellate cells, leading to substantial hepatocyte proliferation, reduced intracellular spaces (Chow et al., 2016) and inhibited
cholangiocyte growth. The signaling in hepatocytes and stellate cells between TRGBR/TGF-β plays a critical role in liver regeneration (Yoshizato et al., 2012). Studies with cultured hepatocytes have shown that TGF-β1 and TGFBR2 are antimitogenic for hepatocytes. Under normal resting (physiological) condition, hepatocytes express TRGFBR2, whereas nonparenchymal cells express low levels of Tgf-β1. When the liver is injured, TRGBR2 expression in hepatocytes is reduced to initiate proliferation. Once an adequate number of hepatocytes is produced, the stellate cells surround the hepatocytes and produce TGF-β, which signals to hepatocytes to terminate proliferation and negatively inhibit TGF-β expression (Yoshizato et al., 2012). Our results showed that hFRGN livers consisted of remarkably lower TGFBR2 expression, but an unchanged TGF-β1 expression vs. that for FRGN (Figure 3A) livers, even though the livers were highly humanized (Table 1). Histopathologic images from current (Figure 1A) and previous studies (Chow et al., 2016) support the evidence of unchecked hepatocyte proliferation and densely populated liver (Figure 1A). Invasive hepatocyte proliferation and inhibited cholangiocyte growth (Figure 3B) resulted in reduced bile flow (Chow et al., 2016) and cholehepatic shunting and increased in hepatic bile acid accumulation and toxicity (Tables 4 and 5). The densely populated hFRGN livers may further lead to ischemia within the acinus, a condition that leads to inhibition of FXR (Cheng et al., 2013).

The inability of humanized mouse livers to regulate its CYP7A1 expression led to high bile acid levels and toxicity. Normally, Cyp7a1/CYP7A1 in liver is under Fxr/FXR and Fgf15/FGF19 control (Figure 7A). At high bile acid levels, Fxr/FXR in the intestine activates intestinal Fgf15/FGF19, which forges an interaction with Fgfr4/FGFR4 to repress Cyp7a1/CYP7A1 (Inagaki et al., 2005). In liver, the FXR-SHP-LRH-1 or FXR-SHP-HNF-4α cascade negatively controls CYP7A1 (Goodwin and Kliewer, 2002). hFRGN mice displayed
higher levels of CDCA, DCA, and CA and their conjugates (Figure 3B), which are strong FXR/Fxr ligands (Makishima et al., 2002), leading to intestinal Fxr activation and intestinal Fgf15 induction (Figure 5A). However, hFRGN livers exhibited high CYP7A1 expression (Figure 4A), suggesting that the signaling pathway associated with Fgf15 appeared to be non-functional, as shown by the unchanged ratio of pERK/total ERK (Figure 5B), suggestive of human hepatocytes and murine intestine miscommunication. Additionally, SHP levels in hFRGN livers (Figure 3A) were lower, suggesting that high bile acid concentration (Table 5) failed to activate hepatic human FXR to suppress CYP7A1 in hFRGN livers. A plausible explanation is due to the bile acid composition changes in hFRGN livers: absence of CDCA and high tβ-MCA (29-folds) and tα-MCA (42-folds) levels (Table 5), which are FXR antagonists (Sayin et al., 2013). These differential abundances in tMCA and CDCA appear to be the driving forces for decreased hepatic FXR activation in hFRGN livers. High bile acid levels could also induce toxicity by activating nuclear factor-kappaB (NF-κB), which is associated with higher interleukin-6 and COX-2 expression towards enhanced growth and apoptosis resistance in cholangiocarcinoma cells; the release of cytokines can also cause biliary damage (Liu et al., 2014).

High bile acid concentrations in hFRGN bile (Table 5) may lead to gallbladder filling in hFRGN mice (Figure 1A), in events that are related to bile acid-mediated activation of Tgr5, the transmembrane G-protein-coupled receptor expressed in cholangiocytes and gallbladder epithelial cells (Li et al., 2011; Keitel and Haussinger, 2013; Duboc et al., 2014). Activation of the Tgr5 in gallbladder smooth muscle cells by bile acids, especially lithocholic and deoxycholic acids and their tauro-conjugates, would result in smooth muscle cell relaxation, gallbladder filling and gallbladder stasis (Lavoie et al., 2010; Jones et al., 2015).
Even though high bile acid levels in hFRGN mice were observed, plasma cholesterol level, which is maintained by the low-density lipoproteins (LDL) and high-density lipoproteins (HDL), remained unchanged, whereas liver cholesterol was increased (Table 4). The exact mechanism for these changes is currently unknown. Ellis et al. (2013) reported that LDL, VLDL, and HDL fractions were shifted in chimeric mice, suggesting possible differences of lipid formation and uptake mechanism. Although HMG CoA reductase (that synthesizes cholesterol) levels (Figure 3A) in hFRGN livers were similar to those in human livers, cholesterol synthesis rates between human and mouse may be different. Certainly, studies are needed to elucidate the upstream pathways of cholesterol synthesis in chimeric livers.

The recent TK-NOG chimeric model may be a better option than hFRGN mice for control of the level of humanization in the mouse liver (Kim et al., 2014). However, the same disrupted signaling between the murine intestine and human hepatocytes and mouse non-parenchymal cells in TK-NOG mouse would also exist. To alleviate elevated bile acids levels, Naugler et al. (2015) suggest that human bile acid production can be controlled by administration of exogenous FGF19 in chimeric mice. However, the dose and dosing regimen of FGF19 administration and the associated effects on drug metabolism and transport are unknown. In addition, the disrupted signaling of human hepatocyte proliferation in the mouse liver requires the introduction of human Kupffer cells implantation in mouse liver (Wilson et al., 2014) or exogenous administration of human TGF-β. Certainly, TGF-β and FGF19 are some of the important signaling mechanisms worth investigation for the optimization of humanized mouse liver models.

The distension of the gallbladder, the unchecked hepatocyte proliferation and liver deformity, and miscommunication between liver cell types in the humanized liver and between
the mouse intestine and humanized liver, as well as the lack of negative feedback control of the FXR-SHP cascade on bile acid homeostasis (Figure 7B) are likely to bring about liver toxicity and model instability. These problems will exist not only in the hFRGN mice, but also for other chimeric (humanized) mice such as the PXB and TK-NOG chimeric mice. The dysregulation of bile acid production in h-chimeric mice and physiological changes that accompany the accumulation of highly toxic bile acids will further contribute to changes in expression of transporter and enzyme in extrahepatic tissues (Figure 6 and Supplementary Figures 1 and 2), which would constitute another important consideration in pharmacokinetics and drug disposition studies. Activation of other nuclear receptors, Pxr and Car, due to elevated bile acids, LCA and the metabolite, 3-keto-5β-cholanic acid (Goodwin and Kliewer, 2002) may result in changes in transporters and enzymes. Bile acids, especially tMCAs and elevated bilirubin levels in hFRGN livers will also contribute to varying extents of Fxr and Car induction in extrahepatic tissues (Huang et al., 2003; Chen et al., 2011). These translate to induction of targeted transporters and enzymes such as Mrp2, Mrp3, Mrp4 and Mdr1a, Gst4-4, Sult1a1, and Ugt1a1 (Huang et al., 2003; Zollner et al., 2006; Zollner and Trauner, 2009; Wagner et al., 2011) in hFRGN intestine, kidney or brain (Figures 6 and Supplemental Figures 1 and 2) and drug dispositional changes. To conclude, instability of the hFRGN liver and extrahepatic tissue with respect to nuclear receptor activation by the dramatic production of differential human (CDCA and CA) and murine (tMCAs) bile acids, both FXR agonists and antagonists, followed by stimulation of the murine intestine and miscommunication among liver cell types and between intestine and humanized liver will induce toxicity and instability issues in this preparation for human drug metabolism studies.
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Author contributions

Participated in research design: Chow, Evans, Silva, Pang
Conducted experiments: Chow, Quach, Wang, Pang
Contributed new reagents or analytic tools: Chow, Quach, Wang, Zhang, Li, Lai, Pang
Performed data analysis: Chow, Quach, Zhang, Wang, Lai, Pang
Wrote or contributed to the writing of the manuscript: Chow, Quach, Zhang, Wang, Li, Evans, Silva, Lai, Tirona, Pang
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Footnotes

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Figure Legends

Figure 1. Appearances of the FRGN, mFRGN and hFRGN livers. (A) Gross appearance: Note the abnormal gross appearance the distended gall bladder, small lobes, and the necrotic tissue in hFRGN livers. (B) Oil Red O staining at 100X (scaling from 0-250 µm with interval of 50 µm) and 400X magnifications (scaling from 0-50 µm with interval of 10 µm) of FRGN, mFRGN, and hFRGN livers (one representative preparation each). Normal morphology was observed in FRGN livers, whereas mFRGN livers showed higher lipid deposits than FRGN livers while lipid deposits in hFRGN livers were only slightly higher. Hepatocytes appeared tightly packed in hFRGN livers.

Figure 2. Bile acid composition in (A) bile, plasma, and liver and (B) bile acid pool size (n=4-8). Note that for hFRGN mice, there were much higher proportions of CDCA, CA, and DCA and their taurine-conjugates (>85% composition), which are strong FXR ligands, whereas for FRGN and mFRGN mice, these were 17-44%. By contrast, murine bile acids: muricholic acid (MCA), αMCA, βMCA and ωMCA and their taurine-conjugates, were 53-81% in FRGN and mFRGN mice, and were 13-15% for hFRGN mice. The bile acid pool size was 4.2-fold higher for hFRGN than that of FRGN mice. Data was mean±SEM; #, P < .05, between FRGN and hFRGN mice, using a 2-tailed Student’s t-test.

Figure 3. Hepatic mRNA expression of (A) human genes in hFRGN livers (n=11) and those in human liver tissues (n=6) and (B) in murine genes present in FRGN, mFRGN, and hFRGN livers (n=4-8). Hepatic human mRNA expression in hFRGN livers were generally higher than those in human liver tissues, whereas FXR and SHP mRNA expression were significantly lower (A). Mouse Tgf-β1, the anti-proliferative gene in non-parenchymal cells, was unchanged while mRNA expression of other murine bile acid metabolic enzymes in mouse hepatocytes and murine proliferative markers and Asbt in cholangiocytes were lower in hFRGN livers (B). Data was mean±SEM; †, P < .05, between human livers and hFRGN livers; *, P < .05, between FRGN and mFRGN livers; #, P < .05, between FRGN and hFRGN livers, using a 2-tailed Student’s t-test.

Figure 4. Proteomics of liver proteins. (A) for human-specific peptide, liver CYP7A1 protein level was > 3-fold higher in hFRGN livers (n=9) than in human liver tissues (n=3); (B) for mouse-specific peptides, murine Cyp7a1 level was comparable to that of FRGN livers, while murine Oatp1b2 protein persisted in hFRGN liver; (C) Cross-reactive (mouse+human) peptides were also used to detect liver proteins in hFRGN and FRGN (n=6-8) livers. Data was mean±SEM; ND denotes not detected/below detection limit; †, P<0.05, between hFRGN and human liver; †, P<.05, between FRGN and mFRGN livers; #, P<.05, between FRGN and hFRGN livers, using a 2-tailed Student’s t-test.

Figure 5. (A) mRNA expression of murine intestinal nuclear receptors, enzymes, and transporters related to bile acid transport/metabolism and (B) the ratio of pERK / total ERK protein levels (in liver) as an activation indicator of the Fgf15-FGFR4 signaling pathway in FRGN, mFRGN, and hFRGN mice (n=4-8). *, P < .05, between FRGN and
mFRGN intestine of same segment or liver; #, \( P < .05 \), between FRGN and hFRGN intestine of same segment or liver; \(^a\) \( P < 0.05 \), between FRGN duodenum and FRGN jejunum or ileum using 2-tailed Student’s \( t \)-test.

**Figure 6.** In intestine, mRNA expression of mouse nuclear receptors, transporters, and enzymes in FRGN, mFRGN, and hFRGN mice (n=4-8). Higher levels of Car and Pxr, in hFRGN intestine, were observed. Higher levels of Mdr1a, Mrp4, Gsta4-4, Sult1a1, and Ugt1a1 in were present in most intestinal segments of hFRGN intestine, although higher Gsta4-4 and Sult1a1, though lower Mrp3 and ileal Bcrp expression were observed in mFRGN intestine (mechanism unknown); other changes between FRGN and mFRGN were minor. *, \( P < 0.05 \) denotes comparison between FRGN and mFRGN intestine of same segment using 2-tailed Student’s \( t \)-test. #, \( P < 0.05 \) denotes comparison between FRGN and hFRGN intestine of same segment using 2-tailed Student’s \( t \)-test. \(^a\) \( P < 0.05 \), between FRGN duodenum and FRGN jejunum or ileum using 2-tailed Student’s \( t \)-test.

**Figure 7.** Schematics depicting interaction between (A) the murine intestine and murine liver and bile acids (denoted as BA), and (B) the murine intestine and humanized liver. (A) Normally, there are two major pathways for Cyp7a1 regulation: (i) in liver, by the Fxr-Shp-Lrh-1 or Hnf-4\( \alpha \) cascade and (ii) in murine intestine by Fgf15, secreted by the ileum and under Fxr stimulation, to react with Fgfr4 in liver to repress Cyp7a1. (B) Due to the production of human bile acids by hFRGN livers, strong FXR antagonists, the tMCAs inhibit liver FXR, whereas strong FXR ligands stimulate intestinal Fxr targets: Shp, Fgf15, Ibabp, and Ost-\( \alpha \). However, miscommunication between the murine intestinal Fgf15 and human FGFR4 and inhibition of liver FXR activated mediated cascades failed to repress CYP7A1 expression. Consequently, CYP7A1 remained unrepressed and high BAs prevailed in this vicious circle. Note that blue lines represent stimulation whereas red line represents inhibition. As well, solid line denotes the activation/increase pathway whereas dot lines denote the absence/decrease of the pathway.
Table 1: **hFRGN human donor and human liver tissue information.** All hFRGN livers were >80% human repopulated, as indicated by Yecuris.

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<td>Liver mRNA &amp; protein</td>
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<tr>
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<td>hFRGN13</td>
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<td>Liver mRNA</td>
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<td>Internal Standard (IS)</td>
<td>Declustering Potential (V)</td>
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<td>CA-d₄</td>
<td>-125</td>
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<td>βMCA</td>
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<td>CA-d₄</td>
<td>-125</td>
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<td>CA-d₄</td>
<td>-125</td>
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<td>CDCA</td>
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<td>CDCA-d₄</td>
<td>-130</td>
</tr>
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<td>DCA-d₄</td>
<td>-130</td>
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<td>gUDCA</td>
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<td>CDCA-d₄</td>
<td>-80</td>
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<tr>
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<td>gDCA</td>
<td>448.3→73.9</td>
<td>CDCA-d₄</td>
<td>-80</td>
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<td>CDCA-d₄</td>
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<td>tCA</td>
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<td>CA-d₄</td>
<td>-120</td>
</tr>
<tr>
<td>Tauro-α-muricholic acid</td>
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<td>CA-d₄</td>
<td>-120</td>
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<tr>
<td>Tauro-β-muricholic acid</td>
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<td>CA-d₄</td>
<td>-120</td>
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<td>Tauro-ω-muricholic acid</td>
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<td>CA-d₄</td>
<td>-120</td>
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<td>CDCA-d₄</td>
<td>-110</td>
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<tr>
<td>Taurochenodeoxycholic acid</td>
<td>tCDCA</td>
<td>498.3→79.9</td>
<td>CDCA-d₄</td>
<td>-110</td>
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<td>DCA-d₄</td>
<td>-110</td>
</tr>
<tr>
<td>Taurolithocholic acid</td>
<td>tLCA</td>
<td>482.3→79.9</td>
<td>LCA-d₄</td>
<td>-130</td>
</tr>
<tr>
<td>Cholic-2,2,4-4-d₄ Acid</td>
<td>CA-d₄</td>
<td>411.3→411.3</td>
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<td>-25</td>
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<tr>
<td>Deoxycholic-2,2,4-4-d₄ Acid</td>
<td>DCA-d₄</td>
<td>395.3→395.3</td>
<td>-130</td>
<td>-25</td>
</tr>
<tr>
<td>Chenodeoxycholic-2,2,4-4-d₄</td>
<td>CDCA-d₄</td>
<td>395.3→395.3</td>
<td>-130</td>
<td>-25</td>
</tr>
<tr>
<td>Lithocholic-2,2,4-4-d₄ Acid</td>
<td>LCA-d₄</td>
<td>379.3→379.3</td>
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<td>-23</td>
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Table 3. Specific peptide sequence for detection of surrogate proteins and MRM conditions

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<tr>
<th>Protein</th>
<th>Species Cross-Reactivity</th>
<th>Peptide Sequence</th>
<th>Q1 (Da)</th>
<th>Q3 (Da)</th>
<th>Declustering Potential (V)</th>
<th>Collision Energy (eV)</th>
<th>Cell Exist Potential (V)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CYP7A1/Cyp7a1</td>
<td>Human/Mouse/Rat</td>
<td>LSSASLNIR</td>
<td>480.8</td>
<td>760.4</td>
<td>66</td>
<td>23</td>
<td>32</td>
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<tr>
<td>CYP7A1</td>
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<td>AHILNNLDNFK</td>
<td>433.4</td>
<td>523.1</td>
<td>44</td>
<td>15</td>
<td>20</td>
</tr>
<tr>
<td>Cyp7a1</td>
<td>Mouse/Rat</td>
<td>LFAVQEIK</td>
<td>474.2</td>
<td>616.2</td>
<td>66</td>
<td>23</td>
<td>26</td>
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<td>Oatp1b2</td>
<td>Mouse/Rat</td>
<td>SVQPELK</td>
<td>400.7</td>
<td>614.3</td>
<td>80</td>
<td>17</td>
<td>26</td>
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<tr>
<td>NTCP/Ntcp</td>
<td>Human/Mouse/Rat</td>
<td>GIYDGDLK</td>
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<td>710.2</td>
<td>80</td>
<td>17</td>
<td>26</td>
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<tr>
<td>BSEP/Bsep</td>
<td>Human/Mouse/Rat</td>
<td>STALQILIQR</td>
<td>515.4</td>
<td>657.5</td>
<td>90</td>
<td>25</td>
<td>26</td>
</tr>
<tr>
<td>BCRP/Bcrp</td>
<td>Human/Mouse/Rat</td>
<td>SSLDVLAAR</td>
<td>522.9</td>
<td>644.3</td>
<td>100</td>
<td>25</td>
<td>26</td>
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<tr>
<td>P-gp</td>
<td>Human/Mouse/Rat</td>
<td>AGAVAEVLAAR</td>
<td>635.3</td>
<td>971.4</td>
<td>95</td>
<td>31</td>
<td>25</td>
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<tr>
<td>MRP2/Mrp2</td>
<td>Human/Rat</td>
<td>LTIIQPDPILFGSGLR</td>
<td>885.8</td>
<td>665.5</td>
<td>90</td>
<td>37</td>
<td>25</td>
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<tr>
<td>MRP3/Mrp3</td>
<td>Human</td>
<td>IDGLNVADIGLHDLR</td>
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<td>697.2</td>
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<td>20</td>
<td>18</td>
</tr>
<tr>
<td>MRP4/Mrp4</td>
<td>Human/Mouse/Rat</td>
<td>APVLFFDR</td>
<td>482.8</td>
<td>697.4</td>
<td>66</td>
<td>28</td>
<td>21</td>
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</table>
Table 4: Physiological parameters of FRGN, mFRGN and hFRGN mice (n=4 to 9)

<table>
<thead>
<tr>
<th>Parameter</th>
<th>FRGN</th>
<th>mFRGN</th>
<th>hFRGN</th>
</tr>
</thead>
<tbody>
<tr>
<td>Body Weight (g)</td>
<td>32.1 ± 3.6</td>
<td>33.0 ± 7.6</td>
<td>30.6 ± 4.0</td>
</tr>
<tr>
<td>Liver (% Body Weight)</td>
<td>7.61 ± 0.27</td>
<td>4.95 ± 0.14†</td>
<td>10.6 ± 0.31*</td>
</tr>
<tr>
<td>Brain (% Body Weight)</td>
<td>1.55 ± 0.07</td>
<td>1.27 ± 0.06†</td>
<td>1.56 ± 0.11</td>
</tr>
<tr>
<td>Kidney (% Body Weight)</td>
<td>2.08 ± 0.18</td>
<td>1.82 ± 0.14</td>
<td>2.24 ± 0.16</td>
</tr>
<tr>
<td>Plasma ALT (IU/ml)</td>
<td>113 ± 15.5</td>
<td>52.2 ± 17.0†</td>
<td>215 ± 30.1*</td>
</tr>
<tr>
<td>Plasma Cholesterol (mg/dl)</td>
<td>151 ± 11.8</td>
<td>161.1 ± 11.0</td>
<td>152 ± 15.6</td>
</tr>
<tr>
<td>Liver Cholesterol (mg/g)</td>
<td>2.17 ± 0.22</td>
<td>2.63 ± 0.20</td>
<td>3.14 ± 21*</td>
</tr>
<tr>
<td>Liver Triglyceride (mg/g)</td>
<td>1.81 ± 0.62</td>
<td>7.36 ± 1.57†</td>
<td>3.31 ± 1.05</td>
</tr>
<tr>
<td>Total Biliary Bilirubin in Bile (mg/dl)</td>
<td>5.0 ± 0.8</td>
<td>NM</td>
<td>11.6 ± 4.7</td>
</tr>
<tr>
<td>Conjugated Bilirubin in Bile (mg/dl)</td>
<td>1.0 ± 0.2</td>
<td>NM</td>
<td>4.9 ± 1.8*</td>
</tr>
</tbody>
</table>

† denotes P< 0.05 compared between mFRGN and FRGN mice using 2-tailed Student’s t-test
* denotes P< 0.05 compared between hFRGN and FRGN mice using 2-tailed Student’s t-test;
* not measured
### Table 5: Bile acid concentrations in bile, plasma and liver (data of Figure 2A) of FRGN, mFRGN and hFRGN livers (n=4 to 8, mean±SEM)

|                  | FRGN (μM) | mFRGN (μM) | hFRGN (μM) | FRGN hFRGN Ratio | mFRGN hFRGN Ratio | hFRGN hFRGN Ratio | FRGN (nmol/g) | mFRGN (nmol/g) | hFRGN (nmol/g) | FRGN hFRGN Ratio |
|------------------|-----------|------------|------------|------------------|------------------|------------------|--------------|--------------|--------------|----------------|-----------------|
| **Bile (μM)**    |           |            |            |                  |                  |                  |              |              |              |                  |
| Total BA        | 23,137 ± 4,290 | 77,369 ± 31,272 | 241,840 ± 47,453* | 10.5             | 6.6 ± 3.1        | 13.3 ± 4.7       | 321 ± 40.1* | 49           | 76.4 ± 24.2   | 121 ± 38.0      | 429 ± 42.2* |
| Primary+secondary BA | 252 ± 98   | 288 ± 88   | 2752 ± 698* | 10.9             | 2.4 ± 1.5        | 4.4 ± 2.5        | 394 ± 14.8* | 17           | 44.8 ± 15.2   | 75.9 ± 25.6     | 8.1 ± 4.0* |
| Tauro-conjugated BA | 22,857 ± 4,262 | 77,027 ± 31,201 | 183,520 ± 32,047* | 8               | 4.2 ± 1.6        | 8.9 ± 3.1        | 153 ± 29.1* | 37           | 31.3 ± 10.2   | 45.5 ± 12.6     | 382 ± 39.8* |
| Glyco-conjugated BA | 28 ± 10    | 52 ± 33    | 55,567 ± 17,757* | 2,003            | <0.01            | <0.01            | 129 ± 36.6* | ∞            | 0.16 ± 0.09   | <0.01           | 39.3 ± 9.7* |
| αMCA            | 9.3 ± 4.8  | 24.9 ± 11.3 | 88.0 ± 17.3* | 9.5              | 0.09 ± 0.05      | 0.56 ± 0.34      | 1.81 ± 0.97  | 20.1         | 1.08 ± 0.32   | 4.61 ± 0.97      | 0.35 ± 0.12* |
| MCA             | 522 ± 150  | 5,580 ± 1,904 | 9,536 ± 3,077* | 18.3             | 0.13 ± 0.04      | 0.55 ± 0.21      | 7.13 ± 2.93* | 55           | 0.32 ± 0.18   | 0.91 ± 0.30      | 9.17 ± 2.12* |
| βMCA            | 63.3 ± 22.9 | 143 ± 41.8  | 956 ± 192   | 1.5              | 1.22 ± 0.91      | 1.83 ± 1.21      | 3.11 ± 0.60  | 2.5          | 31.7 ± 11.83  | 54.6 ± 21.86     | 1.23 ± 0.24* |
| tβMCA           | 9,507 ± 2,720 | 35,606 ± 11,721 | 16,354 ± 3,530* | 1.7             | 1.93 ± 0.89      | 5.26 ± 1.98      | 11.8 ± 2.67* | 6.1          | 0.75 ± 0.38   | 8.14 ± 4.54      | 31.2 ± 3.52* |
| αMCA            | 61.6 ± 27.6 | 68.7 ± 21.9 | 24.5 ± 5.3  | 0.4              | 0.64 ± 0.39      | 0.98 ± 0.75      | 2.43 ± 0.65  | 3.8          | 5.84 ± 2.3    | 7.51 ± 2.88      | 3.05 ± 0.08* |
| tαMCA           | 4,863 ± 1,265 | 11,766 ± 5,849 | 1,021 ± 308* | 0.21             | 0.58 ± 0.22      | 1.31 ± 0.47      | 2.61 ± 0.48* | 4.5          | 1.71 ± 0.99   | 4.03 ± 1.67      | 3.23 ± 0.92 |
| MCA             | 0.1 ± 0.15 | <0.01       | 1.4 ± 0.68  | 14               | 0.01 ± 0         | 0.01 ± 0         | 0.10 ± 0.02* | 10          | 0.49 ± 0.17   | 0.63 ± 0.13      | 0.03 ± 0.01* |
| tMCA            | 117 ± 47   | 51 ± 16     | 2502 ± 680* | 21.3             | 0.2 ± 0.1        | 0.7 ± 0.6        | 16.2 ± 12.3* | 76          | 2.89 ± 1.05   | 3.22 ± 0.68      | 4.58 ± 3.36 |
| tMCA            | 7,318 ± 1,484 | 22,085 ± 11,681 | 109,010 ± 24,806* | 14.9             | 1.33 ± 0.49      | 1.47 ± 0.50      | 61.4 ± 16.9* | 46          | 25.1 ± 8.4    | 28.1 ± 7.5       | 165 ± 36.4* |
| gMCA            | 28 ± 10    | 52 ± 32     | 50,400 ± 15,742* | 1,816            | <0.01            | <0.01            | 24.4 ± 10.9* | ∞           | 0.08 ± 0.05   | <0.01           | 17.1 ± 7.0* |
| tMCA            | 160 ± 33   | 763 ± 12.3* | 8,503 ± 3,440* | 53              | <0.01            | <0.01            | <0.01        | ∞           | 0.01 ± 0.01   | 0.01 ± 0.01      | 0.01 ± 0.01 |
| gMCA            | 671 ± 287* | <0.01       | <0.01       | <0.01            | <0.01            | <0.01            | <0.01        | ∞           | <0.01 ± 0.01  | <0.01 ± 0.01      | <0.01 ± 0.01 |
| tMCA            | <0.01      | <0.01       | 38.5 ± 12.2* | <0.01            | 0.07 ± 0.01      | 0.12 ± 0.03      | 1.48 ± 2.41* | 221         | 0.16 ± 0.07   | 0.14 ± 0.02      | 1.54 ± 0.67 |
| tMCA            | 367 ± 262  | 581 ± 151   | 36,171 ± 12,732* | 99              | 0.09 ± 0.03      | 0.06 ± 0.02      | 66.8 ± 14.7* | 725         | 2.23 ± 1.16   | 1.46 ± 0.21      | 161 ± 18.8* |
| gMCA            | <0.01      | <0.01       | <0.01       | <0.01            | <0.01            | <0.01            | <0.01        | ∞           | <0.01 ± 0.01  | <0.01 ± 0.01      | <0.01 ± 0.01 |
| tMCA            | 2.7 ± 1.2  | 14.4 ± 3.7* | 269.2 ± 131* | 100              | <0.01            | <0.01            | 0.14 ± 0.07  | <0.01        | 0.08 ± 0.01   | 0.18 ± 0.06      | 1.74 ± 0.43* |
| tMCA            | <0.01      | <0.01       | 17.9 ± 8.1*  | <0.01            | <0.01            | <0.01            | 0.31 ± 0.11* | ∞           | 0.02 ± 0.02   | 0.07 ± 0.02      | 3.5 |

† and * denotes P < 0.05 compared between mFRGN and FRGN mice and hFRGN and FRGN mice, respectively, using 2-tailed Student’s t-test or Mann-Whitney test; a denotes ratio of hFRGN/FRGN.
Table 6: Comparison of biliary bile acid composition to that in healthy adults according to Rossi et al. (1987) in human bile (t and g denote taurine- and glycine-conjugated bile acids, respectively)

<table>
<thead>
<tr>
<th>Bile Species</th>
<th>FRGN</th>
<th>mFRGN</th>
<th>hFRGN</th>
<th>Humans</th>
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<tr>
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<td>0.09</td>
<td>1.41</td>
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<td>tCA</td>
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<td>6.5</td>
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<tr>
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<td>0.05</td>
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<td>34.1</td>
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</tr>
<tr>
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<td>0.98</td>
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Figure 2

(B)  

100x  400x

FRGN Liver

mFRGN Liver

hFRGN Liver
Figure 3

(A) Human Nuclear Receptors

- CAR
- FXR
- HNF-1α
- HNF-4α
- Lrhr1
- LXRα
- Pparγ
- PXR
- SHP

(B) Human Other Genes

- Albumin
- P4AT10
- FGF19
- FGF24
- TGFBR2

(C) Human Phase I Enzymes

- CYP1A1
- CYP1B1
- CYP2B6
- CYP2C19
- CYP2D6
- CYP2E1
- CYP3A4
- CYP7A1
- CYP8B1
- CYP9B1

(D) Human Phase II Enzymes

- BAAT
- GSTA4
- SULT1A1
- SULT1A2
- SULT1E1
- SULT2A1
- UGT1A1

(E) Human Influx Transporters

- OATP1B1
- OATP1B3
- OATP2B1
- NTCP

(F) Human Efflux Transporters

- ABCA1
- ABCG1
- ABCG2
- BCRP
- BSEP
- MRPP1
- MRPP2
- MRPP4
- OSTα
- OSTβ
Figure 3
Figure 4

(A) Human specific peptides

Protein Content (fmol/µg)

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(B) Mouse specific peptides

Protein Content (fmol/µg)

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Cross-reactive mouse and human peptides

Influx Transporter

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Efflux Transporters

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# indicates significant difference.
Figure 5

(A) Relative mRNA Expression in Mouse Intestine

- **Fxr**
  - FRGN
  - mFRGN
  - hFRGN

- **Shp**
  - Duodenum
  - Jejunum
  - Ileum

- **Lrh-1**
  - Duodenum
  - Jejunum
  - Ileum

- **Ost-α**
  - Duodenum
  - Jejunum
  - Ileum

- **Ost-β**
  - Duodenum
  - Jejunum
  - Ileum

- **Asbt**
  - Duodenum
  - Jejunum
  - Ileum

- **Ibabp**
  - Duodenum
  - Jejunum
  - Ileum

- **Fgf15**
  - Duodenum
  - Jejunum
  - Ileum

Legend:
- a
- #
Figure 6

Relative mRNA Expression in Mouse Intestine

Car
Pxr
Oatp2b1
PepT1
Bcrp
Mdr1a
Mrp2
Mrp3
Mrp4
Cyp3a11
Gsta3-3
Gsta4-4
Sult1a1
Ugt1a1

Duodenum
Jejunum
Ileum

FRGN
mFRGN
hFRGN