Continuum of Host-Gut Microbial Co-metabolism: Host CYP3A4/3A7 are Responsible for Tertiary Oxidations of Deoxycholate Species

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Received November 26, 2018; accepted December 31, 2018

ABSTRACT

The gut microbiota modifies endogenous primary bile acids (BAs) to produce exogenous secondary BAs, which may be further metabolized by cytochrome P450 enzymes (P450s). Our primary aim was to examine how the gut adapts to the stress of microbe-derived secondary BAs by P450-mediated oxidative modifications on the steroid nucleus. Five unconjugated tri-hydroxy BAs that were structurally and/or biologically associated with deoxycholate (DCA) were determined in human biologic samples by liquid chromatography-tandem mass spectrometry in combination with enzyme-digestion techniques. They were identified as DCA-19-ol, DCA-6α-ol, DCA-5β-ol, DCA-6α-ol, DCA-1β-ol, and DCA-4β-ol based on matching in-laboratory synthesized standards. Metabolic inhibition assays in human liver microsomes and recombinant P450 assays revealed that CYP3A4 and CYP3A7 were responsible for the regioselective oxidations of both DCA and its conjugated forms, glycodeoxycholate (GDCa) and taurodeoxycholate (TDCA). The modification of secondary BAs to tertiary BAs defines a host liver (primary BAs)-gut microbiota (secondary BAs)-host liver (tertiary BAs) axis. The regioselective oxidations of DCA, GDCa, and TDCA by CYP3A4 and CYP3A7 may help eliminate host-toxic DCA species. The 19- and 4β-hydroxylation of DCA species demonstrated outstanding CYP3A7 selectivity and may be useful as indicators of CYP3A7 activity.

Introduction

The drug-metabolizing enzyme (DME) genes are a group of environmental susceptibility genes that have existed on this planet for billions of years (Nebert et al., 1988; Gonzalez and Nebert, 1990), long predating the use of drugs. The expression and activity of these DME genes are decisive in defining our susceptibility to toxic substances or diseases. Based on the roles of nuclear receptors, also known as xenobiotic sensors, that modulate DMEs in growth signal transduction pathways, Nebert (1990, 1991) proposed in 1990 that DMEs are responsible for controlling the steady-state levels of small organic, oxygenated molecules that act as signals for growth, differentiation, virulence, and even tumor promotion. Examples of DME functions beyond drug metabolism are manifested in the synthesis and disposition of steroid hormones by human cytochrome P450 enzymes (P450s). In this current era of metagenomics, it remains

AABBREVIATIONS: 3-dehydroCA, 7α,12α-dihydroxy-3-oxo-5β-cholan-24-oic acid; 3-dehydroCDCA, 7α-hydroxy-3-oxo-5β-cholan-24-oic acid; 3-dehydroDCA, 12α-hydroxy-3-oxo-5β-cholan-24-oic acid; 3-dehydroLCA, 3-oxo-5β-cholan-24-oic acid; 3-deoxyCA, 7α,12α-dihydroxy-5β-cholan-24-oic acid; 12-epiCA, 3α,7α,12β-trihydroxy-5β-cholan-24-oic acid; 12-epiDCA, 3α,12β-dihydroxy-5β-cholan-24-oic acid; AUC, area under the curve; BA, bile acids; bar genes, bile acid-inducible genes CA, cholate, 3α,7α,12α-trihydroxy-5β-cholan-24-oic acid; CDDCA, chenodeoxycholate; 3α,7α-dihydroxy-5β-cholan-24-oic acid; DCA, deoxycholate, 3α,12α-dihydroxy-5β-cholan-24-oic acid; AUC, area under the curve; BA, bile acids; bar genes, bile acid-inducible genes CA, cholate, 3α,7α,12α-trihydroxy-5β-cholan-24-oic acid; CDDCA, chenodeoxycholate, 3α,7α-dihydroxy-5β-cholan-24-oic acid; DCA, deoxycholate, 3α,12α-dihydroxy-5β-cholan-24-oic acid; DCA-19-ol, 3α,12α,19-trihydroxy-5β-cholan-24-oic acid; DCA-6α-ol, 3α,6α,12α-trihydroxy-5β-cholan-24-oic acid; DCA-1β-ol, 1β,3α,12α-trihydroxy-5β-cholan-24-oic acid; DCA-2β-ol, 2β,3β,12α-trihydroxy-5β-cholan-24-oic acid; DCA-4β-ol, 3α,4β,12α-trihydroxy-5β-cholan-24-oic acid; DCA-5β-ol, 3α,5β,12α-trihydroxy-5β-cholan-24-oic acid; DCA-6β-ol, 3α,6β,12α-trihydroxy-5β-cholan-24-oic acid; DCA-4β-ol, 3α,4β,12α-trihydroxy-5β-cholan-24-oic acid; DDA, 3,12α-dihydroxy-5β-cholan-24-oic acid; DCA, deoxycholate; 3α,12α-dihydroxy-5β-cholan-24-oic acid; DCA, deoxycholate; 3α,12α-dihydroxy-5β-cholan-24-oic acid; DCA-19-ol, 3α,12α,19-trihydroxy-5β-cholan-24-oic acid; DCA-1β-ol, 1β,3α,12α-trihydroxy-5β-cholan-24-oic acid; DCA-2β-ol, 2β,3β,12α-trihydroxy-5β-cholan-24-oic acid; DCA-4β-ol, 3α,4β,12α-trihydroxy-5β-cholan-24-oic acid; DCA-5β-ol, 3α,5β,12α-trihydroxy-5β-cholan-24-oic acid; DCA-6β-ol, 3α,6β,12α-trihydroxy-5β-cholan-24-oic acid; DCA, deoxycholate; 3α,12α-dihydroxy-5β-cholan-24-oic acid; DCA, deoxycholate; 3α,12α-dihydroxy-5β-cholan-24-oic acid; DDA, 3,12α-dihydroxy-5β-cholan-24-oic acid; isodCA, 3β,12α-dihydroxy-5β-cholan-24-oic acid; isoDCA, isoCDCA, isoDCA, 3β,7β-dihydroxy-5β-cholan-24-oic acid; LCA, lithocholate, 3α,12α-dihydroxy-5β-cholan-24-oic acid; LC-MS/MS, liquid chromatography-tandem mass spectrometry; MDCA, 3α,6β-dihydroxy-5β-cholan-24-oic acid; NMR, nuclear magnetic resonance; P450, cytochrome P450; p450, recombinant human cytochrome P450; TDCA, taurodeoxycholate; UCA, ursodeoxycholate, 3α,7β,12α-trihydroxy-5β-cholan-24-oic acid; UGT, UDP-glucuronosyltransferase.
poorly understood whether and how P450s participate in the host-gut microbial co-metabolism networks. Such an inquiry is intriguing because the microbiota-derived small molecules are “exogenous” to the host genome, may be candidates for metabolism by P450s, and might be mediators of important microbe-host metabolic interactions (Donia and Fischbach, 2015).

Bile acids (BAs) are an important class of metabolites that undergo cometabolism by host and symbiotic gut microbiota. BA-mediated host-gut microbial crosstalk is important for maintaining a healthy gut microbiota (Theriot et al., 2014; Buffie et al., 2015), balanced lipid and carbohydrate metabolism (de Aguiar Vallim et al., 2013; Wahlström et al., 2016), insulin sensitivity (Maruyama et al., 2002; Kawamata et al., 2003), and innate immunity (Inagaki et al., 2006; Vavassori et al., 2009). Within the BA metabolic network, much attention has been given to defining the hepatic biosynthesis of the primary BAs CDCA and CA (Russell, 2003) and the 7-dehydroxylation of primary BAs by bacteria with bile acid–inducible (bai) genes as secondary BAs, lithocholate (LCA) and DCA (Ridlon et al., 2006; Dawson and Karpen, 2015). In contrast, tertiary BA metabolism, which may be termed the redox modification on the steroid skeleton of secondary BAs by the host, remains underinvestigated. The unclear pathways are fundamentally associated with how the host diagnostically adapts to the stress of the more host-toxic secondary BA stress, along with the colonization, development, homeostasis, and dysbiosis of gut microbiota.

In the last century, several unusual BAs with hydroxylations sites at C-1, -2, -4, -6, and -19 have been identified by gas chromatography–mass spectrometry in humans (Sjövall et al., 2010). Among them, two pathways, 6α-hydroxylation of LCA as hyodeoxycholate (HDCA) (Träulzsch et al., 1974; Araya and Wikvall, 1999; Deo and Bandiera, 2009) and 1β-hydroxylation of DCA as DCA-1β-ol (Gustafsson et al., 1985; Bodin et al., 2005; Hayes et al., 2016), were found to be catalyzed by CYP3A. Thus, HDCA and DCA-1β-ol may be considered tertiary BAs. The other unusual BA species are seldom detected and discussed in recent literature. We developed a liquid chromatography-tandem mass spectrometry (LC-MS/MS) method for the analysis of the human BA metabolome and have been able to detect several new unconjugated tri-hydroxy BAs in adult human serum and urine samples. In this work, we determined that these tertiary BA species are uniquely derived from the oxidation metabolism of DCA, glycodeoxycholate (GDCA), and taurodeoxycholate (TDCA). The P450 enzymes CYP3A4 and CYP3A7 were shown to be exclusively responsible, in a regioslective manner, for the relevant oxidation reactions. This finding has provided a new landscape for the functional biology of CYP3A, one of the most important DMEs, in the host-gut microbial cometabolism of BAs in human.

Materials and Methods

Materials and Reagents. The BA standards obtained from Steraoids (Newport, RI), TRC (Toronto, Canada), Santa Cruz (Dallas, TX), or Sigma-Aldrich (St. Louis, MO) are summarized in Supplemental Table S1. The P450 inhibitors were obtained from TCI (Nihonbashi-honcho, Chuo-ku, Japan), TRC, Santa Cruz, or Sigma-Aldrich. Sulfatase from Helix pomatia type H-1, β-glucuronidase from Helix pomatia type H-1, and cholyglycine hydrolase from clostridium perfringens were purchased from Sigma-Aldrich. Human liver microsomes from 150 mixed-gender pooled donors, human intestinal microsomes from seven mixed-gender pooled donors, NADPH-regenerating system solution A (NADPH-A, containing 26 mM NADP+, 66 mM glucose-6-phosphate and 66 mM MgCl₂ in water), NADPH-regenerating system solution B (NADPH-B, containing 40 U/ml glucose-6-phosphate dehydrogenase in 5 mM sodium citrate), and 0.5 M, pH 7.4, taurodeoxycholate (TDCA) were purchased from Corning (Tewksbury, MA). Human kidney microsomes from five mixed-gender pooled donors were obtained from BioIVT (West Sussex, UK). Recombinant human cytochrome P450 enzymes (p450s) prepared from baculovirus-transfected insect cells (Supersomes), including 1A2, 1B1, 2A6, 2B6, 2C8, 2C9*, 2C18, 2C19, 2D6*, 3A4, 3A5, 3A7, 2E1, 2J2, 4A11, 4F2, 4F3B, and 4F12, were purchased from Corning. Another set of p450s, prepared from plasmid-transfected Escherichia coli (Bactosomes), including 2B6, 2C8, 2C9, 2C19, 3D4, 3A5, and 3A7, were obtained from Cypex Ltd. (Dundee, Scotland). Sodium acetate, glacial acetic acid, and LC-MS grade methanol, acetonitrile, and formic acid were purchased from Sigma-Aldrich. Dimethysulfone (DMSO) was purchased from Thermo Fisher Scientific (Waltham, MA). Ultrapure water was obtained by using a Milli-Q system (Millipore, Bedford, UK).

Synthesis of DCA-19-ol. We synthesized 3α, 12α, 19-trihydroxy-5β-cholan-24-oic acid (DCA-19-ol, CAS 171524-54-2) from CA according to the same procedure described in reference (Kurosawa et al., 1995). The LC-MS/MS and nuclear magnetic resonance (NMR) data of DCA-19-ol are summarized in Supplemental Fig. S1. The following are the spectra data for the methyl 3α, 12α, 19-triacytoxy-5β-cholan-24-oic acid (DCA-19-ol) in deuterated chloroform (CDCl₃) δ 174.8, 72.9, 71.7, 68.0, 51.5, 48.3, 47.6, 47.1, 46.5, 35.5, 35.1, 34.9, 34.4, 32.8, 31.1, 30.8, 29.7, 28.7, 28.4, 27.4, 23.6, 23.1, 17.2, and 12.6.

Synthesis of DCA-6α-ol and DCA-6β-ol. We synthesized 3α, 6α, 12α-trihydroxy-5β-cholan-24-oic acid (DCA-6α-ol, CAS 21066-18-2), and 3β, 6β, 12α-trihydroxy-5β-cholan-24-oic acid (DCA-6β-ol, CAS 63266-90-0) from CA according to the same procedure described in reference (Iida et al., 1991b). The LC-MS/MS and NMR data of DCA-6α-ol and DCA-6β-ol are summarized in Supplemental Fig. S2 and S3, respectively. The following are the spectra data for the methyl 3α, 6α, 12α-trihydroxy-5β-cholan-24-oic acid (DCA-19-ol) in deuterated chloroform (CDCl₃) δ 174.8, 72.9, 71.7, 68.0, 51.5, 48.3, 47.6, 47.1, 46.5, 35.5, 35.1, 34.9, 34.4, 32.8, 31.1, 30.8, 29.7, 28.7, 28.4, 27.4, 23.6, 23.1, 17.2, and 12.6.

Synthesis of DCA-5β-ol. We synthesized 3α, 5β, 12α-trihydroxy-cholan-24-oic acid (DCA-5β-ol, CAS 58678-36-7) were synthesized from DCA according to the same procedure described in reference (Iida et al., 2001). The LC-MS/MS and NMR data of DCA-5β-ol were summarized in Supplemental Fig. S4. The following are the spectra data for the methyl 3α, 12α-dehydroxy-5β-hydroxycholan-24-oic acid (DCA-5β-ol) in deuterated chloroform (CDCl₃) δ 174.8, 72.9, 71.7, 68.0, 51.5, 48.3, 47.6, 47.1, 46.5, 35.5, 35.1, 34.9, 34.4, 32.8, 31.1, 30.8, 29.7, 28.7, 28.4, 27.4, 23.6, 23.1, 17.2, and 12.7.

Synthesis of DCA-1β-ol. We synthesized 1β, 3α, 12α-trihydroxy-5β-cholan-24-oic acid (DCA-1β-ol, CAS 80434-32-8) from DCA according to the same procedure described in Hayes et al. (2017). The LC-MS/MS and NMR data of DCA-1β-ol were summarized in Supplemental Fig. S5. The following are the spectra data for the methyl 1β, 3α-dihydroxy-12α-acetoxy-5β-cholan-24-oic acid (DCA-1β-ol) in deuterated chloroform (CDCl₃) δ 174.8, 72.9, 71.7, 68.0, 51.5, 48.3, 47.6, 47.1, 46.5, 35.5, 35.1, 34.9, 34.4, 32.8, 31.1, 30.8, 29.7, 28.7, 28.4, 27.4, 23.6, 23.1, 17.2, and 12.7.

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Human Serum and Urine. Postprandial human serum and urine were collected from 13 healthy adult volunteers (Ferslew et al., 2015). After ingestion of the standardized high-fat breakfast, urine samples were collected and pooled over the 2-hour period; blood samples were collected in untreated glass tubes at 0.0, 0.5, 1.0, 1.5, and 2.0 hours and allowed to clot for 30–60 minutes to separate the serum. This study was approved by the University of North Carolina at Chapel Hill (UNC-CH) Biomedical Institutional Review Board and published in ClinicalTrials.gov (NCT01766096). Overnight fasting spot urine samples were collected at West China Hospital of Sichuan University from 45 healthy volunteers including 30 men and 15 women (18–40 years old, body mass index 19–26). Briefly, the inclusion criteria for healthy subjects were normal blood, liver and kidney functions; negative test results for the biomarker of infectious diseases including hepatitis B, hepatitis C, HIV and Treponema pallidum; no abnormalities in electrocardiogram, abdominal ultrasonography and chest radiography; no history of gastrointestinal surgery except for appendectomy; and no ingestion of any medications or dietary supplements 2 weeks before urine collections. The studies were approved by the Institutional Review Board of West China Hospital of Sichuan University. All serum and urine samples were stored at −80°C until analysis.

Sample Preparation for BAs Analysis. Analysis of BAs metabolome were performed using the enzyme digestion techniques published in our recent work (Zhu et al., 2018). For the postprandial human serum and urine samples from 13 healthy adults, aliquot (50 μl) was transferred to 700-μl round-well 96-well plate in quadruplicate. The first aliquot was prepared with treatment-1 (T1) in which 150 μl of sodium acetate buffer (pH 5.0) was added. T1 treatment without the use of enzymes was designed to determine the free unconjugated form. The second aliquot was prepared with treatment 2 (T2), in which 150 μl of buffer containing 100 U of cholyglycine hydrolase was added to deconjugate the N-acetylated forms. The third aliquot was prepared with treatment 3 (T3), in which 150 μl of buffer containing 50 U of sulfatase and 500 U of β-glucuronidase was added to hydrolyze sulfates and glucuronides. The final aliquot was prepared with treatment 4 (T4), in which 150 μl of buffer containing all three enzymes was added. In brief, the free unconjugated form (Free) was always detected in T1–T4; glycine/taurine-conjugates were combinedly observed in T2; sulfates/glucuronides were combinedly detected in T3; glycine/taurine conjugates, sulfates/glucuronides, and “double-conjugates” linked with both glycine/taurine and sulfate/glucuronide were collectively detected in T4 (Total). The fasting spot-urine samples were prepared and determined only with T4 treatment.

The calibration and quality control samples were prepared with T1 treatment. The plate was incubated at 37°C for 6 hours and subsequently lyophilized. In each well, 200 μl of acetone or containing 1% formic acid and 100 nM internal standards (LCA-2,2,4,4-D4, DCA-2,2,4,4-D4, UDCA-2,2,4,4-D4, and CA-2,2,4,4-D4) was added in each well. The plate was vortex-mixed at 1500 rpm at 10°C for 30 minutes and centrifuged at 4°C at 3000g for 20 minutes. Two hundred microliters of supernatant was vacuum-evaporated at 30°C. The residue was reconstituted with 50 μl of acetone and 50 μl water by vortex-mixing at 10°C at 900 rpm for 20 minutes. After centrifugation, the plate was placed into autosampler for subsequent analysis.

Quantitative LC-MS/MS Analysis. Quantitative analysis of unconjugated BAs was performed using the method published in our recent work on ACQUITY ultraperformance LC coupled to a Xevo TQ-S MS (Waters, Milford, MA) (Yin et al., 2017; Zhu et al., 2018). The mobile phases consisted of 0.01% formic acid and 100 mM, acetonitrile (V4.1; Waters) and UNIFI (V1.8; Waters). DCA-19-ol, DCA-6-a-ol, a-glucuronidase was added to hydrolyze sulfates and glucuronides. The final aliquot was prepared with treatment 4 (T4), in which 150 μl of buffer containing all three enzymes was added. In brief, the free unconjugated form (Free) was always detected in T1–T4; glycine/taurine-conjugates were combinedly observed in T2; sulfates/glucuronides were combinedly detected in T3; glycine/taurine conjugates, sulfates/glucuronides, and “double-conjugates” linked with both glycine/taurine and sulfate/glucuronide were collectively detected in T4 (Total). The fasting spot-urine samples were prepared and determined only with T4 treatment. The calibration and quality control samples were prepared with T1 treatment. The plate was incubated at 37°C for 6 hours and subsequently lyophilized. In each well, 200 μl of acetone or containing 1% formic acid and 100 nM internal standards (LCA-2,2,4,4-D4, DCA-2,2,4,4-D4, UDCA-2,2,4,4-D4, and CA-2,2,4,4-D4) was added in each well. The plate was vortex-mixed at 1500 rpm at 10°C for 30 minutes and centrifuged at 4°C at 3000g for 20 minutes. Two hundred microliters of supernatant was vacuum-evaporated at 30°C. The residue was reconstituted with 50 μl of acetone and 50 μl water by vortex-mixing at 10°C at 900 rpm for 20 minutes. After centrifugation, the plate was placed into autosampler for subsequent analysis.

In Vitro Metabolism Studies of BAs. In vitro metabolisms of BAs were performed according to the guidelines released by Corning. In brief, the working solutions of selective P450 inhibitors were prepared in DMSO at a concentration of 10.0 mM for all BA substrates except for LCA (4.0 mM). The working solutions of selective P450 inhibitors were prepared in DMSO at the preset concentrations, including ketoconazole (100 μM), fluconazole (2.0 mM), diethylidithiocarbamate (DEDPC, 4.0 mM), quinidine (200 μM), sulfaphenazole (200 μM), quercetin (200 μM), montelukast (100 μM), sertraline (2.0 mM), trans-2-phenylcyclopropylamine (2-PCPA, 40 μM), and α-naphthoflavone (α-NF, 20 μM). The working solutions of time-dependent inhibitors were also prepared in DMSO, including verapamil (5.0 mM), paroxetine (2.0 mM), ticlopidine (200 μM), tienilic acid (2.0 mM), phenelzine (200 μM), thio–TIOPA (1.0 mM), and furafylline (200 μM). Incubation with an initial substrate concentration of 50 μM (20 μM for LCA) was performed on 96-well plates in a shaking incubator at 37°C. The total incubation volume was 100 μl containing 5.0 μl of NADPH-A, 1.0 μl of NADPH-B, 0.5 μl of working solution of BA substrates, 0.5 μl of working solution of inhibitors or blank solvent, and 2.5 μl of microsomes (protein concentration of 20 mg/ml) or rP450s (1.0 nmol of protein/ml) in 0.1 M, pH 7.4, phosphate-buffered saline. The protein concentration in the incubation media was 0.5 mg/ml for microsomes or 50 pmol/ml for rP450s. All incubations were performed in triplicate. For the inhibition assay using selective P450 inhibitors, the reactions were initiated by adding enzymes (microsomes or rP450s) after a preincubation at 37°C for 5 minutes. For the inhibition assay using time-dependent P450 inhibitors, the reactions were initiated by adding working solution of BA substrates after a preincubation at 37°C for 30 minutes. The reactions were stopped at a preset time point by adding 300 μl of ice-cold acetone containing 0.1% formic acid and 50 μM CA-2,2,4,4-D4 as an internal standard. The reaction mixture was then centrifuged at 4°C at 4000g for 20 minutes. For incubations using unconjugated BAs as substrates, the supernatant (50 μl) was diluted with an equal volume of water and subjected to LC-MS/MS analysis. For incubations using GDCA or TDCA as substrates, the supernatant was vacuum-evaporated and prepared according to the enzyme digestion technique of T2 to hydrolyze and analyze the oxidized metabolites in their unconjugated forms.

Data Processing. The LC-MS/MS raw data were processed by MassLynx (V4.1; Waters) and UNIFI (V1.8; Waters). DCA-19-ol, DCA-6-a-ol, DCA-6-β-ol, DCA-6-oxo, DCA-1-β-ol, and DCA-β-ol were semiquantified using the calibration curve for CA because they produced equivalent responses at the same level at selective ion recording of 407 > 407. The high-definition MS/MS raw data were processed by Xcalibur (V4.2; Thermo Fisher Scientific, San Jose, CA). The MS/MS spectrum of each BA was obtained by combining the scans within the peak width. Multivariate analysis of the total urinary BA profiles was carried out using SIMCA P+ (v13; Umetrics, Umeå, Sweden) and R software (version 2.15.0, www.r-project.org). Analysis of in vitro metabolism data was performed using GraphPad prism (version 7.0; GraphPad software, La Jolla, CA). The apparent renal clearance was calculated with the 0–2 hours total urinary excretion amount (nanomole)/the 0–2 hours total serum area under curve (AUC) (nanomole*minutes per milliliter).

Results

Tri-Hydroxyl BAs Exclusively Derived from DCA Oxidations. As shown in Fig. 1, A–C, we found five unknown tri-hydroxyl BA species (TBA05, TBA09, TBA10, TBA13 and TBA18) in the
post-prandial urine and serum of 13 healthy volunteers collected at UNC-CH Hospitals (Zhu et al., 2018). Enzyme digestion techniques demonstrated that they were extensively conjugated, thus, explaining their "invisibility" to those direct analysis techniques without consideration of BA conjugation. The Quad time-of flight MS/MS spectra captured in the digested urine samples showed that they had fragment ions possibly associated with 12-hydroxylation on the skeleton (Lan et al., 2016; Zhu et al., 2018). They were therefore supposed to be structurally associated with DCA and CA, the two major BAs with 12-hydroxylation. Additional analysis of the total BA profiles in spot urine samples from 45 healthy Chinese adults confirmed that they are common BA species in human. Metabolomic analysis of the total urinary BA profiles from 58 healthy adults was employed to examine the biologic associations of the unidentified tri-hydroxyl BAs with known BA species. Despite vast individual differences, the potential biologic relationships were observed using hierarchical cluster analysis of Pearson correlation coefficients between the urinary total unconjugated BAs profile in the test population of 58 healthy volunteers (36 men and 22 women).

Characterization of the Stereoselective Oxidation of DCA. Besides the 3α- and 12α-hydroxyl groups, there is only one unknown oxidation site on the steroid skeleton of TBA04, TBA05, TBA09, TBA10, TBA13 and TBA18. The unusual hydroxylation sites at C-1, C-2, C-4, C-6 and C-19 have been reported in earlier gas chromatography-mass spectrometry-based studies of human BAs and were particularly discussed as being present in early human life (Sjövall et al., 2010). We gratefully acknowledge the contribution from Takashi Iida who shared his negative electrospray ionization/time-of-flight MS data of some early synthesized compounds in the MASSBANK database (Horai et al., 2010), which enabled us to have a relatively reasonable guide for our own synthetic efforts. By matching the high-definition MS/MS spectra (Fig. 2) and retention data (Fig. 1 and 3) of the synthesized standards to those of metabolites captured in vivo or in vitro, TBA04, TBA05, TBA09, TBA10, TBA13 and TBA18 were eventually identified as DCA-19-ol, DCA-6β-ol, DCA-5β-ol, DCA-6α-ol, DCA-1β-ol and DCA-4β-ol, respectively. As far as we know, this is the first report confirming the occurrence of DCA-5β-ol and DCA-19-ol in humans.

Preliminary fragmentation analysis identified common mass losses corresponding to [M-H$_2$O-CO]$^-$ (m/z 345.3) and/or [M-H$_2$O-CO]$^-$ (m/z 343.3) in the spectra of DCA-6β-ol, DCA-6α-ol, DCA-1β-ol and DCA-4β-ol. A diagnostic fragment ion, [C$_{19}$H$_{27}$O]$^-$ (m/z 271.2), was found in the spectra of DCA-5β-ol, which is similar as the ring cleavage fragment of CA, [C$_{19}$H$_{29}$O]$^-$ (m/z 289.2). Diagnostic fragment ions associated with other ring-cleavage mechanisms,
were detected in the spectra of DCA-4\textsubscript{b}-ol and DCA-19-ol, respectively. These observations were to some extent consistent with the previously identified key role of 12-hydroxylation in the negative fragmentation of unconjugated C24 BAs (Lan et al., 2016). The detailed analysis is to be summarized in our later analytical report.

We have detected both in vivo and in vitro (Figs. 1C and 3A; Supplemental Fig. S19), another major metabolite of DCA, 3-dehydroDCA (Zhu et al., 2018). Consistent with previous reports (Amuro et al., 1986; Deo and Bandiera, 2008, 2009), the 3-dehydrogenation reaction was also observed for LCA, CDCA, UDCA and CA (Supplemental Fig. S20–S23). The 3-dehydrogenation mechanism was proposed to be involved with a 3\textsubscript{a},3\textsubscript{b}-diol intermediate formed by 3\textsubscript{b}-hydroxylation, based on the evidence that the incubation of isoDCA in human liver microsomes did not produce 3-dehydroDCA (Bodin et al., 2005). Our work has provided additional evidence for the 3\textsubscript{b}-oxidation mechanism because C3 was the major oxidation site of 3-deoxyCA (Supplemental Fig. S18). To this end, we have

![Fig. 2. High-definition MS/MS spectra of the oxidized DCA metabolites captured from the synthesized standards (a) and the metabolites detected in vivo or in vitro (b). All standards were synthesized in the laboratory except 3-dehydDCA was obtained from TRC (cat. no. O856870). The data of endogenous DCA-6\textsubscript{b}-ol, DCA-5\textsubscript{b}-ol, DCA-6\textsubscript{a}-ol, DCA-1\textsubscript{b}-ol, and DCA-4\textsubscript{b}-ol were captured from a representative digested urine sample. The in vitro data of DCA-19-ol and 3-dehydroDCA were captured from the DCA incubation samples in human liver microsomes.](https://dmd.aspetjournals.org/content/287/1/2286.full.html)
provided strong evidence that DCA is regioselectively hydroxylated at C-1β, -3β, -4β, -5β, -6α, -6β, and -19. As shown in Fig. 3C, the identified oxidation sites cluster on the same plane around the 5β-hydrogen. We have compared the stereoselective oxidation rates of DCA in the pooled microsomes prepared from human liver, intestine and kidney. The human kidney microsomes showed no reactivity to DCA oxidations at any sites (data not shown). Our data shows that the DCA oxidations happen primarily in human liver microsomes and secondarily in human intestinal microsomes (Fig. 3D). No formation of DCA-19-ol and DCA-4β-ol was observed in human intestinal microsomes.

**CYP3A Inhibitors Selectively Inhibit DCA Oxidations in Human Liver Microsomes.** A series of in vitro inhibition assays were performed to begin the characterization of the P450 isoenzymes responsible for DCA oxidations. Figure 3, E and F, shows the inhibitory effects of various P450 inhibitors, either time-dependent or not, on the regioselective oxidations of DCA. No inhibition effects were observed for the selective inhibitors for CYP1A2 (furafylline and a-naphthoflavone), CYP2C8 (phenelzine, montelukast, and quercetin), CYP2C9 (sulfaphenazole), and CYP2D6 (quinidine), and the nonselective inhibitors for CYP2A6 (2-PCPA), CYP2C19 (nootkatone), and CYP2E1 (DEDC). Tiencilic acid, a selective inhibitor for CYP2C9, showed a mild inhibition to the oxidations at C19 (14%) and C4-β (13%). Thio-TEPA, a nonselective inhibitor for CYP2B6, showed a mild inhibition to the oxidation at C5-β (12%). Ticlopidine, a nonselective inhibitor for CYP2C19, showed mild inhibition to the oxidation at C5-β (7%).

Paroxetine (10 μM), a selective time-dependent CYP2D6 inhibitor, showed mild inhibition to the oxidations at C3-β (17%), C1-β (22%), C5-β (30%), C6-α (7%), and C6-β (26%). Sertraline (10 μM), a nonselective CYP2B6 inhibitor with time-dependent inhibitory effects on CYP3A4 (Masubuchi and Kawaguchi, 2013), showed mild inhibition to the oxidations at C3-β (31%) and C1-β (28%). Fluconazole (10 μM), a strong inhibitor of CYP2C19 and moderate inhibitor of CYP2C9 and CYP3A, showed mild to moderate inhibition of oxidation at C3-β (33%) and C6-β (23%). Verapamil (25 μM), a selective time-dependent CYP3A inhibitor, showed strong inhibition of oxidation at C3-β (60%), C1-β (79%), C5-β (85%), C6-α (65%), and C6-β (100%) and moderately at C19 (27%) and C4-β (37%). Ketoconazole (0.5 μM), an index CYP3A inhibitor, showed strong inhibition of oxidation at C3-β (71%), C1-β (76%), C5-β (80%), C6-α (68%), and C6-β (76%) and, to a lesser extent, at C19 (49%) and C4-β (51%). These results clearly demonstrated that CYP3A is the major isoenzymes responsible for the tertiary stereoselective oxidation of DCA.

**CYP3A4 and CYP3A7 Are Responsible for DCA Oxidations Regardless of N-Acylamidation.** We investigated DCA oxidation in a
panel of rP450s (0.5 nmol protein/ml) compared with that in human liver microsomes (0.5 mg protein/ml). As shown in Fig. 4A, CYP3A4 and CYP3A7 showed the highest catalytic activity at all oxidation sites, whereas CYP3A5 showed a much lower activity level of 3-dehydrogenation and trace or no oxidation at the other sites. For the other P450s, except for CYP3A, only trace amounts of 3-dehydroDCA and DCA-6β-ol were detected in incubations of CYP2C8. As shown in Supplemental Fig. S24, the specific oxidations of DCA through CYP3A4 and CYP3A7 were consistently observed in two rP450 systems, Supersomes and Bactosomes. The metabolic enzyme phenotyping results were consistent with those of the human liver microsome inhibition assay. We thus concluded that CYP3A4 and CYP3A7 were dominantly responsible for the tertiary oxidation of DCA.

The circulated DCA is conjugated in the liver through N-acyl linkage with either glycine as GDCA or taurine as TDCA. We compared the in vitro metabolite formation of DCA, GDCA, and TDCA in rP450s and human liver microsomes, in which the oxidized metabolites of GDCA and TDCA were detected after being digested by chocholyglycine hydrolase (Zhu et al., 2018). Typical ion chromatograms of DCA, GDCA, or TDCA incubated in human liver microsomes, rCYP3A4, rCYP3A5, and rCYP3A7 were shown in Supplemental Fig. S25–S28. Almost the same enzyme specificity with unique contributions of CYP3A4 and CYP3A7 was observed for the oxidation of GDCA and TDCA (Fig. 4). An exception was that the enzyme specificity was lost for 3β-oxidation of GDCA and TDCA, to whom CYP2E1, 2J2, 4A11, 4F2, 4F3B, and 4F12 demonstrated higher.
reactivity than the other tested P450s. The regioselective activities of CYP3A4, CYP3A5, and CYP3A7 to oxidize DCA, GDCA, and TDCA are comparatively highlighted in Fig. 5. CYP3A4 demonstrated a reverse activity toward 1β- and 6α-oxidation of DCA with and without the N-acylamidation. Except for the oxidation at C-3β, CYP3A4 oxidized DCA predominantly at C-1β, whereas GDCA and TDCA were oxidized by CYP3A4 most frequently at C-6α. Unlike CYP3A4, CYP3A7 exhibited a nearly consistent regioselectivity for DCA, GDCA, and TDCA. N-acylamidation of DCA significantly increased the CYP3A7-mediated 1β, 5β, 6α, 19, and 4β-hydroxylation, with the highest activity observed for TDCA. The activities of CYP3A7 for some reactions were significantly greater than those of CYP3A4. The metabolite formation ratios of CYP3A7/CYP3A4 decreased in the following order: TDCA 19-hydroxylation (81.4), GDCA 19-hydroxylation (70.9), DCA 19-hydroxylation (43.8), GDCA 1β-hydroxylation (11.8), TDCA 1β-hydroxylation (9.1), and DCA 4β-hydroxylation (8.7). Based on enzyme specificity data, it was believed that these reactions are promising as possible novel in vitro probe reactions for CYP3A7 activity. In summary, it was clearly demonstrated that CYP3A4 and CYP3A7 were responsible for the regioselective oxidation of DCA, regardless of its N-acylamidation status.

Conjugation Pattern of Tertiary BAs. Based on the characterized tertiary BA metabolism pathways, we showed the host-gut microbial cometabolism for the downstream metabolism of CA (Fig. 6A) and CDCA (Fig. 6B) in 13 healthy adults during the 2 hour-postprandial period after a high-fat diet. Enzyme digestion techniques demonstrated that all the tertiary BAs were extensively conjugated, resulting in an easily neglected level for their free unconjugated forms, particularly in serum samples. Despite the tertiary BAs having a low serum level, DCA-6α-ol, DCA-1β-ol, DCA-5β-ol, and HDCA were determined as the major tertiary species presenting with total urinary levels comparable with primary and secondary BAs in all 13 subjects. By using the enzyme-digestion technique, the total level of a BA may be categorized in four fractions (Fig. 6C): the free unconjugated form, the amidated form, the glucuronidated/sulfated form, and the “double-conjugates” linked with both glycine/taurine and glucuronide/sulfate. The latter two fractions are both glucuronidated/sulfated species. The average percentage conjugation patterns of serum AUC and urine level for each BA were shown in Fig. 6, A and B. Some data were not available because detections were with a trace level (<10 nM) in some samples. For example, DCA-19-ol was not found in any serum samples and was detected in only 2 of 13 urine samples.

Fig. 4. CYP3A4 and CYP3A7 are responsible for the oxidation of DCA, GDCA, and TDCA. The metabolite formation after incubations of 50 μM DCA (A), GDCA (B), and TDCA (C) for 60 minutes in a panel of 18 rCYPs (CYP1A2, 1B1, 2A6, 2B6, 2C8, 2C9*1, 2C18, 2C19, 2D6*1, 3A4, 3A5, 3A7, 2E1, 2J2, 4A11, 4F2, 4F3B, and 4F12; 50 pmol protein/ml) were compared with the data of human liver microsomes (0.5 mg protein/ml) acquired in parallel. The oxidized metabolites of GDCA and TDCA were detected after being digested by cholesterylhydrolase. Data were shown as mean ± S.D. (n = 3).
DCA-6α-ol and HDCA, the 6α-hydroxylated metabolites of DCA and LCA, were the most heavily glucuronidated/sulfated tertiary species. The glucuronidated/sulfated fractions of DCA-6α-ol and HDCA were significantly more elevated than DCA and LCA, respectively. HDCA circulated in serum and excreted in urine mainly in glucuronidated/sulfated forms and DCA-6α-ol existed mainly in “double-conjugated” forms. A similar conjugation pattern was observed for DCA-4β-ol and DBA03, an unidentified LCA hydroxylated metabolite that was detected in 6 of 13 subjects. In comparison, DCA-1β-ol and DCA-5β-ol, which did not undergo glucuronidation/sulfation as significantly as DCA-6α-ol did, were excreted in urine, mainly in amided forms. The glucuronidated/sulfated fractions of DCA-1β-ol and DCA-5β-ol in urine were both significantly lower than that of DCA. DCA-6β-ol and MDCA, the 6β-hydroxylated metabolites of DCA and LCA, were also excreted in urine mainly in amided forms. Therefore, we deduced that tertiary 6α-oxidation facilitates glucuronidation/sulfation metabolism, whereas 1β- and 5β-oxidation of DCA species do not. Further studies are required to investigate the glucuronidation and sulfation mechanisms of tertiary BA species under the different conjugation patterns of them.

**Tertiary Oxidation Facilitates the Renal Excretion of DCA Species.** The free and total 0- to 2-hour urinary levels and time-dependent serum concentrations (mean ± S.E.M.) of each BA detected by T1 and T4 are comparatively showed in Fig. 6 and Supplemental Table S2. The primary BAs, CDCA and CA, are synthesized from cholesterol, N-acylamidated, secreted into the bile, and recycled in the enterohepatic system. The free and total levels of CDCA and CA were significantly higher in serum than urine, manifesting a conservative renal disposition for the primary BAs to facilitate their enrichment in the enterohepatic system. After postprandial release into the intestine, most CDCA and CA (95%) are recovered in the bile by the ileal apical sodium-dependent bile acid transporter, the sinusoidal Na+-taurocholate cotransporting polypeptide, and the canicular bile salt export pump, and the remaining 5% enters the lower gut, where they are further metabolized by gut bacteria into secondary BAs. The major metabolic pathway for secondary BAs is deconjugation of the amidated forms by bacteria with bile salt hydrolase activity and the subsequent 7-dehydroxylation of unconjugated forms by bacteria that carries the bai gene. The resulting LCA and DCA are lipophilic and exogenous to the host. Most of the LCA is excreted in feces because it is much less soluble than DCA, which is consistent with the relative total AUC of DCA/CA (3.68 ± 1.08, mean ± S.E.M., n = 13) being about 40 times higher than that of LCA/CDCA (0.09 ± 0.01); however, the free and total levels of DCA were also significantly higher in serum than urine, indicating that DCA is also inclined to be recycled in the enterohepatic system, as seen for CA and CDCA. Therefore, there might be another mechanism to regulate BA homeostasis by disposing the “toxic” secondary BAs, particularly, DCA which is less hydrophobic and has a much greater exposure than LCA.

In contrast to CA and DCA, the total levels of major tertiary metabolites of DCA (DCA-6α-ol, DCA-1β-ol, DCA-5β-ol, and DCA-4β-ol) were significantly higher in serum than urine, indicating that they are preferentially excreted in urine compared with the primary and secondary BAs. We calculated the apparent renal clearance data based on the total urinary excretion data and the total serum AUC data during the 2 hour-postprandial period. The effects of conjugation metabolism (glucuronidation, sulfation, and N-acylamidation) on renal clearance were included in the apparent clearance data. As shown in Fig. 6D, the tertiary BAs demonstrated significantly higher apparent renal clearances than did the primary and secondary BAs, particularly for the downstream metabolites of DCA. In this regard, we believed that tertiary BA metabolism functions, at least in part, in regulating the BA pool size by facilitating the renal excretion of secondary BAs. Accordingly, the urinary total BA profile is an ideal footprint to understand the functional status of the host-gut microbial cometabolism of BAs in biomedical and transitional researches.

**Discussion**

This work has characterized the CYP3A-catalyzed metabolic pathway of tertiary BAs in the host-gut microbial cometabolism network of BAs in human adults (Fig. 6). Our work confirmed that the previously presumed “unusual” oxidized metabolites of DCA are ubiquitously occurring in the adult population but are often undetected in serum owing to their extensive conjugation metabolism, which allows them to undergo renal excretion. It was also determined in this study that CYP3A4, the most abundant P450 isoenzyme in adult liver, was responsible for the tertiary oxidations of DCA, GDCA, and TDCA. The fact that fetal-specific CYP3A7 plays an important role in the tertiary oxidation of DCA, GDCA, and TDCA may also explain why those BAs with “unusual” oxidation sites were preferably found in neonates, infants, and pregnant women (Sjøvall et al., 2010). Because of challenges in differentiating metabolic contributions from host or gut microbiomes in human subjects, more studies in germ-free animals are required to confirm whether gut bacteria participate the oxidation metabolism of DCA species. Based on the analysis of the postprandial BA profiles in serum and urine, it is believed that the tertiary BA metabolism functions, at least in part, to facilitate the renal excretion of secondary BAs and accordingly regulate the BA pool size and composition. As suggested by the in vitro inhibition data (Fig. 3, E and F), the inhibition of tertiary BA metabolism might...
be pathologically associated with cholestasis-associated liver injury induced by strong CYP3A inhibitors, such as ketoconazole (Stricker et al., 1986).

Enzyme digestion techniques were used in this work to investigate the conjugation metabolism of BAs (Zhu et al., 2018). The indirect method may not provide information of conjugation sites and pathways but clearly present different conjugation patterns of the investigated BA species. Based on composite analysis of the renal clearance data and conjugation pattern data in Fig. 6, we propose that the tertiary oxidation of DCA species contributes to host disposition of DCA species in two major ways. On one hand, the CYP3A4/7 mediated 6α-oxidation of DCA species facilitates the glucuronidation/sulfation metabolism and the subsequent renal excretion. Similar phenomena have been reported for the UDP-glucuronosyltransferase (UGT) 2B4- catalyzed glucuronidation and the subsequent urinary excretion of HDCA, the 6α-hydroxylated metabolite of LCA (Sacquet et al., 1983; Parquet et al., 1985; Radomi-Knska-Pyrek et al., 1987; Pillot et al., 1993). On the other hand, the CYP3A4/7 catalyzed 1β- and 5β-oxidation of DCA species does not help glucuronidation/sulfation but may directly facilitate the renal secretion. UGT1A3, UGT2B4, UGT2B7, and sulfotransferase 2A1 have been reported to be involved in the conjugation metabolism of BAs (Trottier et al., 2006). More efforts are required to investigate glucuronidation, sulfation and tubular secretion mechanisms of the hydroxylated metabolites of DCA species.

CYP3A4 is considered of great importance in general pharmacology, and CYP3A7 has become increasingly important in pediatric pharmacology (Stevens, 2006). Despite the high degree of sequence identity between the genes encoding human CYP3A4 and CYP3A7, it is still incompletely understood how they adapt to and regioselectively oxidize a wide range of compounds (Sevrioukova and Poulos, 2017). A lack of probe reactions is acknowledged as one of the key problems. Beyond the recently proposed DCA 1β-hydroxylation (Hayes et al., 2016), this work has provided the regioselective oxidations of DCA, GDCA, and TDCA as novel in vitro probe reactions for CYP3A4 and CYP3A7 activities. As shown in Fig. 3C, CYP3A4 and CYP3A7 both show C-H activation capacities around the 5β-hydrogen on the steroid skeleton of DCA, GDCA, and TDCA. Relative to CYP3A4, CYP3A7 favored the oxidations at C-19, C-4β, and C-1β. The 19-hydroxylation of DCA, GDCA, and TDCA demonstrated an overwhelming CYP3A7 selectivity than the currently used CYP3A7 probe reactions, 16α-hydroxylation of dehydroepiandrosterone and the 2α-hydroxylation of testosterone (Leeder et al., 2005). Therefore, future studies in combination with protein crystallography and simulation of molecular docking will be useful in promoting our understanding of the structural basis for the regioselective oxidation mechanisms of CYP3A4 and CYP3A7.

The fetal-specific CYP3A7 is also significantly expressed in a subset of adult livers partially associated with the CYP3A7*1C allele (Sim et al., 2005). In comparison with the reported data of human fetal livers
58 healthy volunteers demonstrated a high 4-categorized as dietary supplements (Zanger and Schwab, 2013), all of which may be (Goodwin et al., 1999), and constitutive androstane receptor (Goodwin 2004), vitamin D receptor (Drocourt et al., 2002), pregnane X receptor (19-hydroxylation of DCA species. are required to decipher the in vitro-in vivo inconsistency of the metabolism of 19-hydroxylated metabolites in vivo. More studies are regulated by both endogenous and exogenous factors. In fact, the time course of postnatal regulation of CYP3A7 toward CYP3A4 is intriguingly consistent with the colonization of infant gut bacteria responsible for secondary BA production, such as Bacteroides, Clostridium, Lactobacillus, Bifidobacterium, and Listeria spp., are responsible for BA deconjugation and Clostridium and Eubacterium spp. responsible for 7a-dehydroxylation (Gérard, 2013; Ridlon et al., 2016). We therefore hypothesize that the ontogeny of CYP3A4 and CYP3A7 responsible for tertiary BAs metabolism might be associated with a postnatal protective mechanism in response to the stress of secondary BAs.

In conclusion, this work disclosed the continuum mechanism in the host-gut microbial cometabolism of BAs in human adults. The primary BAs are synthesized from cholesterol in host liver, the secondary BAs are modified from primary BAs by gut microbiota, and the recovered secondary BAs are remodeled as tertiary BAs in host liver. CYP3A4 and CYP3A7 were shown to be predominantly responsible for the tertiary regioselective oxidation of DCA, GDCA, and TDCA at C-1β, -3β, -4β, -5β, -6α, -6β, and -19 on the steroid skeleton. The disclosed tertiary BA metabolic pathways have extended the biologic function of CYP3A4 and CYP3A7 from being an important DME that acts to detoxify pharmacologic agents to that of having an inherent role in the host response to the stress of secondary BAs. In summary, we propose that discovering the inherent roles of DMEs in host-gut microbial cometabolism, along with the colonization, development, homeostasis, and dysbiosis of the symbiotic microbiota will become an important future scientific endeavor for the prevention of drug and environment-induced toxicity.

Acknowledgments

We thank Dr. Takashi Iida (Nihon University) for the gift of isoUCA and contributions for the MS data in the public online resources MASSBANK.

Authorship Contributions

Participated in research design: Liu, Jia, Lan.
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