Gut Microbiota Dysbiosis Is Associated with Altered Bile Acid Metabolism in Infantile Cholestasis

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ABSTRACT The co-occurrence of gut microbiota dysbiosis and bile acid (BA) metabolism alteration has been reported in several human liver diseases. However, the gut microbiota dysbiosis in infantile cholestatic jaundice (CJ) and the linkage between gut bacterial changes and alterations of BA metabolism have not been determined. To address this question, we performed 16S rRNA gene sequencing to determine the alterations in the gut microbiota of infants with CJ, and assessed their association with the fecal levels of primary and secondary BAs. Our data reveal that CJ infants show marked declines in the fecal levels of primary BAs and most secondary BAs. A decreased ratio of cholic acid (CA)/chenodeoxycholic acid (CDCA) in infants with CJ indicated a shift in BA synthesis from the primary pathway to the alternative BA synthesis pathway. The bacterial taxa enriched in infants with CJ corresponded to the genera Clostridium, Gemella, Streptococcus, and Veillonella and the family Enterobacteriaceae and were negatively correlated with the fecal BA level and the CDCA/CA ratio but positively correlated with the serological indexes of impaired liver function. An increased ratio of deoxycholic acid (DCA)/CA was observed in a proportion of infants with CJ. The bacteria depleted in infants with CJ, including Bifidobacterium and Faecalibacterium prausnitzii, were positively and negatively correlated with the fecal levels of BAs and the serological markers of impaired liver function, respectively. In conclusion, the reduced concentration of primary BAs in infants with CJ likely upregulates the conversion from primary to secondary BAs.

IMPORTANCE Liver health, fecal bile acid (BA) concentrations, and gut microbiota composition are closely connected. BAs and the microbiome influence each other in the gut, where bacteria modify the BA profile, while intestinal BAs regulate the growth of commensal bacteria, maintain the barrier integrity, and modulate the immune system. Previous studies have found that the co-occurrence of gut microbiota dysbiosis and BA metabolism alteration is present in many human liver diseases. Our study is the first to assess the gut microbiota composition in infantile cholestatic jaundice (CJ) and elucidate the linkage between gut bacterial changes and alterations of BA metabolism. We observed reduced levels of primary BAs and most secondary BAs in infants with CJ. The reduced concentration of fecal BAs in infants with CJ was associated with the overgrowth of gut bacteria with a pathogenic potential and the depletion of those with a potential benefit. The altered gut microbiota of infants with CJ likely upregulates the conversion from primary to secondary BAs. Our study provides a new perspective on potential targets for gut microbiota intervention directed at the management of infantile CJ.

KEYWORDS bile acids, cholestatic jaundice, dysbiosis, gut microbiota, infants

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Infantile cholestatic jaundice (CJ) is defined as jaundice caused by elevated conjugated bilirubin in infancy (1). The incidence of CJ is about 1 in 2,500 term infants (2). Infantile CJ is uncommon but potentially indicates severe liver disease (3). It was shown that several life-threatening disorders may have cholestasis as a major presenting sign of underlying neonatal liver disease (4). The early detection of CJ and accurate diagnosis are very important for optimal treatment and a favorable prognosis (3). The etiologies of CJ are complex, including biliary obstruction/structural abnormalities and genetic/metabolic disorders (3, 4). The major biliary obstruction/structural abnormality causes of CJ in infancy are biliary atresia, duct gallstones, and choledochal cyst (3, 4). Genetic/metabolic disorder causes of CJ include alpha-1 antitrypsin deficiency, galactosemia, hypopituitarism, tyrosinemia, progressive familial intrahepatic cholestasis, cystic fibrosis, and panhypopituitarism (3). Additional causes of CJ are neonatal hepatitis, viral infection, and parenteral nutrition-associated liver disease (3, 5).

The common clinical feature of CJ is cholestasis, which is defined as reduced bile formation or flow, resulting in the retention of substances normally excreted into bile (1). Bile is produced by the liver and stored and concentrated in the gallbladder. The main components of bile are water, electrolytes, and organic molecules, including bile acids (BAs), cholesterol, phospholipids, and bilirubin (6). The BAs in bile are derived from the catabolism of highly insoluble cholesterol, which facilitates the digestion and absorption of dietary lipids and fat-soluble vitamins in the small intestine (7). The primary BAs, cholic acid (CA) and chenodeoxycholic acid (CDCA), are synthesized in the liver, stored in the gallbladder, and discharged into the small intestine after eating and play important physiological functions in food digestion and waste product elimination (6). The majority (about 95%) of primary BAs are efficiently reabsorbed from the terminal ileum (8). The remainder (about 5%) reach the colon, where they are deconjugated, dehydrogenated, and dehydroxylated by the intestinal bacteria to form secondary BAs and passively absorbed into the portal circulation (9). Once they return to the liver, the BAs are reconjugated and then resecreted together with newly synthesized bile salts. This overall process constitutes one cycle of the enterohepatic circulation. BAs have multiple endocrine functions as regulators of hepatic glucose and lipid metabolism, liver regeneration, inflammation, and the gut microbiota (7, 10). Cholestasis results in the abnormal accumulation of bile salts, bilirubin, and lipids in liver and the blood, impairing bile-mediated physiological functions (5).

The gut microbiota is a large and diverse community of microorganisms that contribute to human health and disease (11). The human gut microbiota facilitates harvesting of nutrients and energy from the ingested food and produces numerous metabolites to regulate host metabolism and immune functions (11). The gut microbiota plays a central role in the metabolism of BAs by regulating deconjugation, dehydroxylation, dehydrogenation, and epimerization to convert primary BAs into unconjugated secondary and free BAs (12). It has been shown that the gut microbiota modulates BA synthesis by regulating the expression of enzymes for BA formation (13). BA deconjugation is catalyzed by bacteria with bile salt hydroxylase activity (14). Deconjugated primary BAs that escape reuptake in the small intestine are then deconjugated and dehydroxylated into secondary BAs by the 7α-dehydroxylating microbiome (15). Conversely, BAs also act as inhibitors of the gut microbiome via antimicrobial effects on gut microbes by damaging bacterial membranes and altering the intracellular macromolecular structure (14) and via indirect effects through the activation of innate immune genes in the small intestine, such as nuclear farnesoid X receptor (FXR)-induced antimicrobial peptides (16).

The co-occurrence of gut microbiota dysbiosis and BA metabolism alteration has been shown in many human liver diseases, including nonalcoholic fatty liver diseases (17, 18), primary biliary cholangitis (19), primary sclerosing cholangitis (20), and liver cholestasis (21–25), but data on pediatric populations are lacking. In the current study, we aimed to assess the mutual influences between the altered gut microbial community and BA metabolism in infantile CJ.
RESULTS

Clinical characteristics of infants with CJ. As shown in Table 1, of those 56 infants with CJ enrolled in the study, 34 (34/56, 60.7%) were boys. The median age at diagnosis was 67 days (interquartile range [IQR], 55.0, 93.5 days). Fourteen infants (14/56, 25.0%) were diagnosed with biliary atresia, and eight infants (8/56, 14.3%) were confirmed to carry a gene mutation reported to cause CJ (3). There was no statistically significant difference in gender, age, and feeding type among infants with CJ and impaired hepatic function (IHF) and healthy control (HC) infants. Blood tests showed elevated levels of BA (158.5 μmol/liter; IQR, 115.3, 213.5 μmol/liter), direct bilirubin (DB; 93.0 μmol/liter; IQR, 70.5, 123.7 μmol/liter), total bilirubin (TB; 163.1 μmol/liter; IQR, 120.2, 214.2 μmol/liter), and gamma-glutamyltransferase (GGT; 149.0 U/liter; IQR, 99.5, 316.8 U/liter) in infants with CJ, and these were significantly higher than those in infants with IHF (Table 1). High levels of alanine aminotransferase (ALT; 214.0 U/liter; IQR, 127.3, 349.0 U/liter) and aspartate aminotransferase (AST; 121.0 U/liter; IQR, 76.5, 233.8 U/liter) were also observed in infants with CJ, and the ALT level was significantly higher in infants with CJ than in subjects with IHF. Additionally, slightly lower levels of albumin (ALB) and hemoglobin (HB) were detected in infants with CJ than in subjects with IHF.

Fecal BA profiles are significantly altered in infants with CJ. We used a targeted metabolomics approach to evaluate the fecal BA profiles in the 95 individuals enrolled in this study, including 54 infants with CJ, 25 infants with IHF, and 16 HC infants. The levels of all primary BAs were decreased in infants with CJ, and these decreases were significant for allocholic acid (ACA), CA, CDCA, and α- and β-muricholic acid (α- and

### TABLE 1 Characteristics of infants with CJ, IHF, and HC

| Characteristic | Value for the following group: | HC (n = 45) | IHF (n = 25) | CJ (n = 56) | $X^2/ Z^a$ | P  |
|---------------|---------------------------------|------------|-------------|-------------|------------|----|
| No. (%) of infants by gender | | | | | 2.000 | 0.955 |
| Boys | 26 (57.8) | 15 (60.0) | 34 (60.7) | | | |
| Girls | 19 (42.2) | 10 (40.0) | 22 (39.3) | | | |
| Median (IQR) age (days) | 90.0 (60.0, 140.0) | 79.0 (61.0, 145.0) | 67.0 (55.0, 93.5) | | 2.650 | 0.080 |
| No. (%) of infants with the following feeding type: | | | | | 4.000 | 0.215 |
| Breast-feeding | 19 (42.2) | 11 (44.0) | 12 (21.4) | | | |
| Formula | 7 (15.6) | 5 (20.0) | 13 (23.2) | | | |
| Combination | 19 (42.2) | 9 (36.0) | 31 (55.4) | | | |
| No. (%) of infants with the following diagnosis: | | | | | 3.000 | 0.001 |
| CMV infection | | 6 (24.0) | 3 (5.4) | | | |
| Genetic disorders | | 0 (0) | 8a (14.3) | | | |
| Biliary atresia | | 0 (0) | 14 (25.0) | | | |
| Unknown | 19 (76.0) | | 31 (55.3) | | | |
| Median (IQR) concn | | | | | | |
| ALB (g/liter) | 41.0 (39.0, 43.0) | 39.0 (36.0, 41.3) | | | −2.573 | 0.010 |
| ALT (U/liter) | 138.0 (62.0, 201.0) | 214.0 (127.3, 349.0) | | | −2.689 | 0.007 |
| AST (U/liter) | 118.0 (56.0, 188.0) | 121.0 (76.5, 233.8) | | | −0.854 | 0.393 |
| BA (μmol/liter) | 33.0 (15.0, 72.8) | 158.5 (115.3, 213.5) | | | −4.944 | 0.000 |
| DB (μmol/liter) | 4.0 (2.1, 7.0) | 93.0 (70.5, 123.7) | | | −7.157 | 0.000 |
| HB (g/liter) | 112 (99.0, 119.0) | 101.5 (93.8, 111.0) | | | −1.663 | 0.096 |
| GGT (U/liter) | 78.0 (49.0, 167.0) | 149.0 (99.5, 316.8) | | | −2.986 | 0.003 |
| TB (μmol/liter) | 12.4 (7.4, 18.3) | 163.1 (120.2, 214.2) | | | −6.748 | 0.000 |

aALB, albumin; ALT, alanine aminotransferase; AST, aspartate aminotransferase; BA, bile acid; DB, direct bilirubin; HB, hemoglobin; GGT, gamma-glutamyltransferase; TB, total bilirubin; IQR, interquartile range; CMV, cytomegalovirus; HC, health control; IHF, impaired hepatic function; CJ, cholestatic jaundice.

bTwo infants had Alagille syndrome with a Jagged 1 (JAG1) mutation, four had citrin deficiency with a solute carrier family 25 member 13 (SLC25A13) mutation, and two had BA synthesis defects with aldo-keto reductase family 1 member D1 (AKR1D1) mutation.

cThe data were compared by the nonparametric Mann-Whitney test (two groups) or Kruskal-Wallis H test (multiple groups). The distribution of dichotomous variables was compared by the chi-squared test.

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Reduced levels of most of the secondary BAs were found in infants with CJ, with the exceptions being \( \beta \)-hyodeoxycholic acid (\( \beta \)-HDCA) (Fig. 1). The patterns of decrease were also observed for the primary and secondary conjugated BAs in infants with CJ, although statistical significance was not reached. In order to
evaluate the enzymatic processes in BA metabolism that may underlie the differences noted in infants with CJ, we investigated the two ratios (Table 2) reflective of enzymatic activities in the liver and the gut microbiome proposed previously (26). The CA/CDCA ratio was measured and was found to be significantly lower in infants with CJ, reflecting a shift in BA synthesis from the primary to the alternative BA pathway that occurs in the liver. The deoxycholic acid (DCA)/CA ratio was increased in infants with CJ, although the difference was not significant, indicating that the enhanced production of secondary BAs occurred in a proportion of infants with CJ.

**CJ is associated with gut microbiota dysbiosis.** To investigate the changes in the gut microbiome, we performed 16S rRNA gene sequencing on 126 fecal samples from infants with CJ (n = 56), infants with IHF (n = 25), and HC infants (n = 45). Alpha diversity, measured by the Shannon and Chao1 indexes, was significantly lower in infants with CJ than those with IHF and HC (Fig. 2A and B). Principal-coordinate analysis (PCoA) of a Bray-Curtis distance matrix generated from genus-level taxa showed a separation between the microbiomes of infants with CJ and HC infants (Fig. 2C), whereas the samples from infants with IHF were not able to be distinguished from those of the infants in the other two groups.

We obtained 17 operational taxonomic units (OTUs) with a significantly different abundance among the three groups (P < 0.05 by one-way analysis of variance [ANOVA]; these OTUs consisted of taxa that had an abundance of greater than 0.1% across all samples and that occurred in at least 20% of the samples) (Fig. 3; see Table S1 in the supplemental material). CJ patients showed increased abundances of taxa assigned to the genera *Clostridium sensu stricto* (OTU2 and OTU46), *Streptococcus* (OTU13), and *Veillonella* (OTU40) and the family Enterobacteriaceae (OTU24) compared with the HC. Some taxa belonging to the genus *Enterococcus* (OTU68), *Gemella* (OTU45), and *Streptococcus* (OTU91) were more abundant in infants with IHF than in HC. The levels of potentially beneficial bacteria, such as *Bifidobacterium* (OTU96) and *Faecalibacterium prausnitzii* (OTU151), were significantly decreased in infants with CJ. The levels of the taxon *Faecalibacterium prausnitzii* (OTU151) were also significantly reduced in children with IHF. In addition, the relative abundances of some taxa belonging to the genera *Bacteroides*, *Blautia*, *Coprococcus*, *Eggerthella*, and *Flavonifractor* and the Lachnospiraceae were also significantly reduced in infants with CJ compared with the HC.

**CJ-related gut microbiome functional alterations.** BugBase software was applied to analyze the community-wide phenotypes of the stool microbiome. We observed that the proportion of aerobic bacteria was increased modestly in infants with IHF (Fig. 4A). The proportions of anaerobic bacteria (Fig. 4B) were significantly reduced and those of facultatively anaerobic bacteria (Fig. 4C) were enriched in infants with CJ. Bacterial function-associated oxidative stress tolerance (Fig. 4D) was found to be enriched in the CJ subjects. Gram-negative bacteria (Fig. 4E) were decreased and Gram-positive bacteria (Fig. 4F) were greatly increased in infants with IHF in comparison with their levels in HC and CJ subjects. In addition, the bacterial function-associated mobile element content (Fig. 4G), biofilm formation (Fig. 4H), and pathogenesis (Fig. 4I) were found to be enriched in the CJ subjects. These findings were mainly attributed to the proteobacteria (most likely the *Enterobacteriaceae* family).

### TABLE 2 Ratios of BAs reflective of liver and gut microbiome and enzymatic activities<sup>a</sup>

| Ratio informative about metabolic processes | Ratio calculated | Mean (95% CI) | P (Tukey’s test) |
|--------------------------------------------|------------------|--------------|-----------------|
| **BA synthesis: primary vs alternative pathway** | CA/CDCA | 6.745 (1.905) | 6.683 (1.27) | 1.818 (0.408) | 1 | 4.2E−09 |
| **Conversion from primary to secondary BA by gut bacteria** | DCA/CA | 0.207 (0.159) | 0.29 (0.22) | 1.179 (0.636) | 0.9 | 0.16 |

<sup>a</sup>BA, bile acid; CA, cholic acid; CDCA, chenodeoxycholic acid; DCA, deoxycholic acid; HC, health control; IHF, impaired hepatic function; CJ, cholestatic jaundice; CI, confidence interval.

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PICRUSt (Phylogenetic Investigation of Communities by Reconstruction of Unobserved States) software was used to predict the relative abundance of KEGG functions, which were compared between children with CJ and IHF and the HC. We found that bacterial transporters were increased in children with altered liver function (both infants with CJ and infants with IHF) compared to the HC subjects (Fig. 5). In addition, ATP-binding cassette (ABC) transporters were particularly more abundant in CJ subjects than in the HC.

**Covariance between serum BAs, fecal BAs, and the gut microbiome.** To examine the relationship between members of the gut microbiota and BAs, we calculated Spearman’s rank correlation coefficient for the 17 OTUs (those that were significantly changed in infants with CJ and/or IHF), serological indexes, and fecal BAs. We performed unsupervised clustering of OTUs, serological parameters, and fecal BAs, which revealed four distinct OTU clusters (clusters O1 to O4; Fig. 6A) and a clear separation of fecal BAs and serological indexes. OTUs in the first cluster, cluster O1 (which included taxa assigned to the genera *Clostridium sensu stricto*, *Veillonella*, and *Streptococcus* and the family *Enterobacteriaceae*), were positively and negatively correlated with the levels of serum BAs (and most of the serological indexes) and fecal BAs, respectively. In contrast, the OTUs in cluster O4 (which included the taxa *Blautia*, *Eggerthella*, *Faecalis*,...
FIG 3  Significant differences between the gut bacterial taxa of the cholestatic jaundice (CJ), impaired hepatic function (IHF), and healthy control (HC) groups. Comparisons among the groups were performed using one-way analysis of variance (ANOVA; \( P < 0.05 \)). The Wilcoxon test was performed for pairwise comparisons, *, \( P < 0.05 \); **, \( P < 0.01 \); ***, \( P < 0.001 \). g, genus; s, species; f, family.
bacterium, Flavonifractor, Lachnospiraceae incertae sedis, and Ruminococcus) were negatively and positively correlated with the levels of serum BAs and fecal BAs, respectively. Notably, clusters O1 and O4 were positively and negatively correlated with the ratio of secondary/primary BAs (DCA/CA), respectively.

The associations between the functional profiles of the gut microbiota and hepatic function and fecal BAs were also investigated. Two distinct functional clusters (clusters F1 and F2) resulted from applying Spearman correlations between the PICRUSt-predicted KEGG pathways representative of each group (linear discriminant analysis [LDA] score > 3, \( P < 0.05 \)), serological indexes, and fecal BAs (Fig. 6B). The KEGG pathways in cluster F1, which included components related to other ion-coupled transporters, the phosphotransferase system, the secretion system, transcription factors, transporters, ABC transporters, and two-component systems, were found to be positively associated with most serological indexes and inversely associated with the fecal levels of BAs, which were stronger for primary and secondary BAs than for their conjugated components. The KEGG pathways in cluster F2, which included alanine aspartate and glutamate metabolism, DNA replication proteins, DNA repair and recombination proteins, other glycan degradation, purine metabolism, pyrimidine metabolism, ribosome, and transcription machinery, were negatively and positively correlated with serum BAs (as well as other serological indexes of liver functions) and fecal BAs, respectively. The levels of KEGG pathways in cluster F1 were found to be positively and negatively correlated with DCA/CA and CDCA/CA, respectively, whereas the functions of cluster F2 showed the opposite trend of correlations.
DISCUSSION

The primary BAs are synthesized in the liver through two pathways, namely, the classical (neutral) and the alternative (acidic) pathways. The classical BA synthetic pathway produces about 90% of total BAs, composed of CA and CDCA in approximately equal amounts. In contrast to the classical pathway, the alternative pathway produces only CDCA (6). Alterations in BA synthesis and composition are important factors that modulate hepatotoxicity in chronic liver diseases. Liver health, fecal BA concentrations, and the gut microbiota composition are closely connected (22). BAs and the microbiome influence each other in the gut, where bacteria modify the BA profile, while intestinal BAs regulate the growth of commensal bacteria, maintain barrier integrity, and modulate the immune system (22). An inborn error of BA synthesis can produce abnormal BA metabolites, which can lead to cholestatic liver disease in infancy and progressive neurological disease in childhood and into adulthood (27, 28). In this study, we demonstrated the association between the changes of the gut microbiome and BA metabolism and estimated the potential impact on host physiology in infant CJ patients. We observed reduced levels of the primary BAs and most of the secondary BAs in stool samples from infants with CJ, a finding which is consistent with the typical symptoms of cholestasis. Infants with CJ showed a decreased fecal CA/CDCA ratio, indicating that the classic pathway was impaired, while the alternative one was preserved. In addition, an increase in the ratio of secondary to primary BAs (DCA/CA) was observed in infants with CJ, which may suggest altered activity of bacterial 7α-dehydroxylases, leading to an excess production of secondary BAs, many of which are cytotoxic to the liver. Actually, the concentrations of two secondary BAs, α- and β-HDCA, were found to be increased in infants with CJ compared with the HC and infants with IHF.

We demonstrated that the CJ-associated gut microbiota dysbiosis was characterized by a decreased biodiversity, a decrease in the microbiota with a beneficial potential (the genera Bifidobacterium and Faecalibacterium), and an overgrowth of potentially pathogenic bacteria (the Enterobacteriaceae family). Species of the genus Bifidobacterium are believed to confer beneficial effects upon their host, and their high abundance in the infant gut indicates saccharolytic activity toward glycans. Bifidobacterium appears to influence short-chain fatty acid (SCFA) production, either directly, by modulating the synthesis of SCFAs, such as acetate and formate (29), or indirectly, by altering the gut microbiota composition and/or microbiome-microbiome interactions (30). In addition,
species of the genus *Bifidobacterium* also contribute to maintaining the normal function of the gut mucosa and protect the mucosa from injurious factors, such as allergens, pathogens, and toxins (31). The genus *Faecalibacterium*, which comprises the sole known species *Faecalibacterium prausnitzii*, consists of well-established protective bacteria with the capacity to reinforce the barrier integrity of epidermal cells and repress inflammation through the production of SCFAs (32, 33). Thus, declines in the abundances of *Bifidobacterium* and *Faecalibacterium prausnitzii* may evoke and enhance systemic inflammation in the host. A member of the family *Enterobacteriaceae* was significantly enriched in the gut microbiota of CJ infants, resulting in an increased level of bacterial gene contents with pathogenic potential, and its abundance was positively correlated with serum indicators of liver damage. Additionally, lower abundances of *Lachnospiraceae* and *Blautia* were detected in infants with CJ, which is similar to the findings for patients with cirrhosis (21, 22).

FIG 6 Correlation matrices constructed from gut bacterial taxa, fecal BAs, and serum indicators of hepatic function. Spearman correlation analysis between fecal BA serum indicators of hepatic function and the top 17 OTUs with significantly different abundances among the groups (A) and the gut microbiota functional categories (B) was employed. Four distinct OTU clusters (clusters O1 to O4) and two distinct functional clusters (clusters F1 and F2) were observed. Red and blue represent the positive and negative correlations, respectively. GHDC, glycohyodeoxycholic acid; TCA, taurocholic acid; GCA, glycocholic acid; LCA, lithocholic acid; GLCA, glycolithocholic acid; THDCA, taurohyodeoxycholic acid; TLCA-3S, tauroliithocholic acid-3-sulfate acid; GCDCA, glycochenodeoxycholic acid; GHCA, glycohyocholic acid; TUDCA, tauroursodeoxycholic acid; GDCA, glycodeoxycholic acid; THCA, taurohyocholic acid; TDCA, taurodeoxycholic acid; GUDCA, glycursohydroxycholic acid; ALB, albumin; ALT, alanine aminotransferase; AST, aspartate aminotransferase; BA, bile acid; DB, direct bilirubin; HB, hemoglobin; GGT, gamma-glutamyltransferase; TB, total bilirubin.
The known gut bacteria capable of processing \(7\alpha\)-dehydroxylases are the species of *Clostridium* (such as *Clostridium hiranonis*, *C. hylemonae*, *C. scindens*, and *C. sordelli*) (15). Due to a lack of redundancy in this system, any perturbations to the \(7\alpha\)-dehydroxylating bacteria are likely to induce an imbalance in the ratio between secondary and primary BAs and impact hepatic function. Although two taxa of *Clostridium* were found to be enriched in the infants with CJ, their identification to the species level could not be accurately determined based on the sequences of the 16S rRNA V3-V4 regions. Thus, further investigations of the microbiome profile with shotgun metagenomic sequencing are needed to identify the exact species that shift \(7\alpha\)-dehydroxylation during cholestasis.

Functional analysis of the bacterial communities in the infants with CJ revealed the enhanced expression of ABC transporters. In the host system, ABC transporters serve as an important cytoprotective mechanism by eliminating toxins and drugs out of cells, thereby protecting the hepatocyte from toxicity due to BA overaccumulation. Bacterial ABC transporters are also able to protect the microorganisms from xenobiotic pressure (34). The apparent upregulation of ABC transporters and other ion-coupled transporters might implicate an enhanced antimicrobial pressure in the gut environment.

In summary, infantile CJ is associated with a significant dysbiosis of the gut microbiota, which is featured by a general decline in bacteria with beneficial potential as well as those capable of processing the necessary enzymes to convert primary to secondary BAs and an enrichment of potentially pathogenic taxa. These changes in the gut microbiota might modulate the composition of BAs, disrupt the gut barrier integrity, and provoke and enhance inflammation.

### MATERIALS AND METHODS

**Study cohort.** A total of 126 infants from Shanghai Children’s Hospital, Shanghai, China, were recruited to the study cohort between September 2016 and March 2019 (Table 1). The diagnosis of CJ in the infants was based on the guidelines described elsewhere (1, 3). An abnormal direct bilirubin (DB) level was defined as a value greater than 17 \(\mu\)mol/liter if the total bilirubin (TB) level was less than 85 \(\mu\)mol/liter or a value of DB that represented more than 20% of that of TB if the TB level was greater than 85 \(\mu\)mol/liter (1). Serological tests for liver function were performed. Levels of BA, DB, TB, gamma-glutamyltransferase (GGT), aspartate aminotransferase (AST), alanine aminotransferase (ALT), and albumin (ALB) were measured by using an AU5800 clinical chemistry analyzer (Beckman Coulter, Brea, CA, USA). Hemoglobin (HB) was measured by use of a sodium lauryl sulfate (SLS) hemoglobin test (Sysmex XN-1000 automatic hematology analyzer; Sysmex Corporation). Twenty-five infants with normal levels of DB and TB but with elevated AST and ALT levels were enrolled in the study as controls and were termed the impaired hepatic function (IHF) group. Forty-five healthy infants without any history of chronic diseases were enrolled as healthy controls (HC). None of the healthy infants had clinically relevant CJ or any symptoms at the time of fecal collection, and subjects who took antibiotics within 2 months before fecal collection or who had any inflammatory conditions were excluded from the study. Written informed consent was obtained from the parents or legal guardians of the infants eligible for study enrollment. This study was approved by the Regional Ethical Review Boards of Shanghai Children’s Hospital and carried out in accordance with the principles of the Declaration of Helsinki 1966 and later versions.

**Quantification of fecal BAs.** Targeted metabolomics profiling was performed to measure the concentrations of 43 BAs in fecal samples according to previously reported methods (35). In brief, each accurately weighed lyophilized fecal sample (~10 mg) was homogenized with 50 \(\mu\)l of water using a Bullet Blender tissue homogenizer (Next Advance Inc., Averill Park, NY). An aliquot of 150 \(\mu\)l of acetonitrile containing 9 internal standards was added, and the extraction was performed using the homogenizer. After centrifugation, 50 \(\mu\)l of each supernatant was transferred to a 96-well plate and diluted with 150 \(\mu\)l of a mobile phase mixture (mobile phase B-mobile phase A [50:50, vol/vol]). The injection volume was 5 \(\mu\)l. After centrifugation, 5 \(\mu\)l supernatant was used for measurement by liquid chromatography-tandem mass spectrometry analysis (LC-TQMS). An Acquity ultrasphere liquid chromatography (UPLC) system (Waters Corp., Milford, MA, USA) coupled with a Xevo TQ-S mass spectrometer (Waters Corp., Milford, MA, USA) was used to quantitate the BAs. MassLynx software (version 4.1; Waters Corp., Milford, MA, USA) was used for instrument control and data processing. Chromatographic separation was achieved with a Waters BEH C\(_{18}\) column (particle size, 1.7 \(\mu\)m; 2.1 mm by 100 mm [internal dimensions]). The UPLC-mass spectrometry (MS) raw data were acquired in negative mode and were processed using the TargetLynx application manager (Waters Corp., Milford, MA, USA) to obtain calibration equations and the measured concentration of each bile acid in the samples.

**Fecal microbiome analysis.** Stool samples were collected with sterile swabs by nursing staff at Shanghai Children’s Hospital and stored at ~80°C. Genomic DNA extraction, PCR amplification, library preparation, and Illumina sequencing were conducted according to a protocol described previously (36). In brief, total microbial DNA was extracted using a QIAamp DNA stool minikit (Qiagen, Germany). The extracted genomic DNA was PCR amplified with barcoded primers (forward primer, 5’-CCT ACG GGA GGC
AGC AG-3’; reverse primer, 5’-GGA CTA CHV GGG TWT CTA AT-3’) targeting the 16S rRNA V3-V4 region. Water samples that had undergone the same procedures of DNA extraction and PCR amplification were used as a control. An equal amount of DNA from each sample was pooled and verified using an Agilent 2100 bioanalyzer (Agilent, USA). Sequencing was performed using an Illumina MiSeq platform at HRK Bio-Tech Co., Ltd. (Shenzhen, China).

A total of 8,265,997 raw sequence reads were produced from 126 stool samples. The paired reads were merged. The average number of reads per sample was 65,603 ± 23,222. The average length after joining of the reads was 462 ± 8 bp per sample. Data processing was performed by using the USEARCH (version 10.0.240) program (37) with an open-source bioinformatics pipeline described at http://www.drive5.com/usearch. The depth of the sequencing data used for analysis was 10,000 reads per sample, based on the findings of the rarefaction analysis (see Fig. S1 in the supplemental material). We applied the Unoise error correction (denoising) algorithm to reconstruct a set of correct biological sequences in the reads and generate zero-radius operational taxonomic units (OTUs) (38). Taxonomic assignment was performed by using the Ribosomal Database Project (RDP) classifier. Statistical analysis was performed with the Calypsog web server (39). The alpha diversity of the fecal microbiome was measured by use of the Shannon and Chao1 indexes. The overall differences in microbiome structure were evaluated through principal-coordinate analysis (PCoA) of a Bray-Curtis distance. BugBase was applied to predict the organism-level microbiome phenotypes (40). The KEGG ortholog functional profiles of the microbial communities were predicted by PICRUSt (Phylogenetic Investigation of Communities by Reconstruction of Unobserved States) software (41). The linear discriminant analysis (LDA) effect size method (LEfSe) was applied to determine the PICRUSt-predicted functions that were enriched in the different groups. The differences in the relative abundances of the taxa and the concentrations of BAs among the groups were assessed for statistical significance by one-way ANOVA followed by Wilcoxon pairwise comparisons. The Spearman correlation was applied to investigate the associations between serological indexes, fecal BAs, and the composition and functions of the gut microbiome.

Statistical analysis. Statistical analysis was performed with SPSS (version 20.0) software. The data are presented as medians and interquartile ranges (IQR; 25th to 75th percentiles) and compared by use of the nonparametric Mann-Whitney test (for two groups) or the Kruskal-Wallis H test (for multiple groups). The distribution of dichotomous variables was compared by the chi-squared test. A P value of <0.05 was considered statistically significant.

Data availability. Raw metabonomic data for the bile acids are available in the Metabolights metabolomics repository under study accession number MTBLS1295. Raw sequencing data are available in the European Nucleotide Archive server under study accession number PRJEB33641.

SUPPLEMENTAL MATERIAL
Supplemental material for this article may be found at https://doi.org/10.1128/mSystems.00463-19.

FIG S1, TIF file, 1 MB.
TABLE S1, DOCX file, 0.02 MB.

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We declare no conflicts interest.
Y.W. and T.Z. designed the study. X.Z., Y.X., X.L., H.H., and T.G. analyzed and interpreted the clinical data. Y.W., X.G., J.H., and D.Y. performed the bioinformatic and statistical analyses. D.L. contributed to fecal sample collection. Y.W. and X.G. created the figures and wrote the manuscript, and T.Z. revised the manuscript. All authors read and approved the final manuscript.

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