Bile salt biotransformations by human intestinal bacteria

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Abstract Secondary bile acids, produced solely by intestinal bacteria, can accumulate to high levels in the enterohepatic circulation of some individuals and may contribute to the pathogenesis of colon cancer, gallstones, and other gastrointestinal (GI) diseases. Bile salt hydrolysis and hydroxy group dehydrogenation reactions are carried out by a broad spectrum of intestinal anaerobic bacteria, whereas bile acid 7-dehydroxylation appears restricted to a limited number of intestinal anaerobes representing a small fraction of the total colonic flora. Microbial enzymes modifying bile salts differ between species with respect to pH optima, enzyme kinetics, substrate specificity, cellular location, and possibly physiological function. Crystallization, site-directed mutagenesis, and comparisons of protein secondary structure have provided insight into the mechanisms of several bile acid-biotransforming enzymatic reactions. Molecular cloning of genes encoding bile salt-modifying enzymes has facilitated the understanding of the genetic organization of these pathways and is a means of developing probes for the detection of bile salt-modifying bacteria. The potential exists for altering the bile acid pool by targeting key enzymes in the 7α/β-dehydroxylation pathway through the development of pharmaceuticals or sequestering bile salts also is associated with increased toxic and metabolic effects. High concentrations of secondary bile acids in feces, blood, and bile have been linked to the pathogenesis of cholesterol gallstone disease and colon cancer. We present here a current review of the microbiology of bile acid metabolism in the human GI tract, focusing on understanding the biochemical mechanisms and physiological consequences of such metabolism on both the bacterium and the human host.

SUPPLEMENTARY KEY WORDS bile acids • deoxycholic acid • 7α-dehydroxylation • gallstone disease • colon cancer • bile salt hydrolase • probiotics • hydroxysteroid dehydrogenase • hydrogen sulfide

The human large intestine harbors a complex microbial flora. Bacterial density in the human colon is among the highest found in nature, approaching $10^{12}$ bacteria/g wet weight of feces. In contrast, the host suppresses significant bacterial colonization of the small intestine by a variety of mechanisms, including rapid transit times, antimicrobial peptides, proteolytic enzymes, and bile. Failure of these mechanisms leads to bacterial overgrowth of the small intestine, resulting in malabsorption as bacteria compete with the host for nutrients. Under normal conditions, bacterial fermentation in the colon represents an important salvage mechanism. Complex carbohydrates, which are intrinsically indigestible or which escape digestion and absorption in the proximal gut, are fermented by colonic bacteria to yield short-chain fatty acids. It has been estimated that these short-chain fatty acids constitute 3–9% of our daily caloric intake. Colonic bacteria also contribute to the salvage of bile salts that escape active transport in the distal ileum. The major bile salt modifications in the large intestine include deconjugation, oxidation of hydroxy groups at C-3, C-7, and C-12, and 7α/β-dehydroxylation. Deconjugation and 7α/β-dehydroxylation of bile salts increases their hydrophobicity and their $P_K_a$, thereby permitting their recovery via passive absorption across the colonic epithelium. However, the increased hydrophobicity of the transformed bile salts also is associated with increased toxic and metabolic effects. High concentrations of secondary bile acids in feces, blood, and bile have been linked to the pathogenesis of cholesterol gallstone disease and colon cancer. We present here a current review of the microbiology of bile acid metabolism in the human GI tract, focusing on understanding the biochemical mechanisms and physiological consequences of such metabolism on both the bacterium and the human host.

THE ENTEROHEPATIC CIRCULATION OF BILE ACIDS

Bile acids are saturated, hydroxylated C-24 cyclopentanopenanthrene sterols synthesized from cholesterol in hepatocytes. The two primary bile acids synthesized in the

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241

review
human liver are cholic acid (CA; 3α,7α,12α-trihydroxy-5β-cholan-24-oic acid) and chenodeoxycholic acid (CDCA; 3α,7α-dihydroxy-5β-cholan-24-oic acid). Bile acids are further metabolized by the liver via conjugation (Nacyl amidation) to glycine or taurine, a modification that decreases the Pka to ~5. Thus, at physiological pH, conjugated bile acids are almost fully ionized and may be termed bile salts (6). Bile salts are secreted actively across the canalicular membrane and are carried in bile to the gallbladder, where they are concentrated during the interdigestive period. After a meal, release of cholecystokinin from the duodenum stimulates the gallbladder to contract, causing bile to flow into the duodenum (7). Bile salts are highly effective detergents that promote solubilization, digestion, and absorption of dietary lipids and lipid-soluble vitamins throughout the small intestine. High concentrations of bile salts are maintained in the duodenum, jejunum, and proximal ileum, where fat digestion and absorption take place. Bile salts are then absorbed through high-affinity active transport in the distal ileum (6). Upon entering the bloodstream, bile salts are complexed to plasma proteins and returned to the liver. Upon reaching the liver, they are cleared efficiently from the circulation by active transporters on the sinusoidal membrane of hepatocytes and rapidly secreted into bile. This process is known as the enterohepatic circulation. Figure 2 depicts the enterohepatic circulation in the context of the gastrointestinal anatomy and also indicates the relative numbers and genera of the predominant bacteria inhabiting each section of the GI tract.

During the enterohepatic circulation, bile salts encounter populations of facultative and anaerobic bacteria of relatively low numbers and diversity in the small bowel. Bile salt metabolism by small bowel microbes consists mainly of deconjugation and hydroxy group oxidation. Ileal bile salt transport is highly efficient (~95%), but approximately 400–800 mg of bile salts escapes the enterohepatic circulation daily and becomes substrate for...
significant microbial biotransforming reactions in the large bowel (6). Comparison of bile acid composition in the gallbladder and feces illustrates the extent of microbial bile acid metabolism in the large intestine (Fig. 3). The secondary bile acids deoxycholic acid (DCA; \(3\alpha,12\alpha\)-dihydroxy-5\(\beta\)-cholan-24-oic acid) and lithocholic acid (LCA; \(3\alpha\)-hydroxy-5\(\beta\)-cholan-24-oic acid) are produced solely by microbial biotransforming reactions in the human large intestine. DCA accumulates in the bile acid pool (LCA to a much lesser extent) as a result of passive absorption through the colonic mucosa and the inability of the human liver to \(7\alpha\)-hydroxylate DCA and LCA to their respective primary bile acids. LCA is sulfated in the human liver at the 3-hydroxy position, conjugated at C-24, and excreted back into bile (6). The resultant bile acid sulfate is poorly reabsorbed from the gut. Even though 3-sulfo-LCA/LCA is lost in feces and does not normally accumulate in the enterohepatic circulation (8).

DECONJUGATION OF BILE SALTS

Characteristics of bile salt hydrolase(s)

Deconjugation refers to the enzymatic hydrolysis of the C-24 N-acyl amide bond linking bile acids to their amino acid conjugates. This reaction is substrate-limiting and goes to completion in the large bowel. Bile salt hydrolases (BSHS) are in the cholyglycine hydrolase family (EC 3.5.1.24) and have been isolated and/or characterized from several species of intestinal bacteria (Table I). The importance of the position, charge, shape, and chirality of various analogs of taurine/glycine conjugates on the rate of hydrolysis by BSHs has also been investigated (9). BSHs differ in subunit size and composition, pH optimum,
of a lack of conservation observed in residues making up the substrate binding pocket of the conjugated bile acid hydrolase gene product of *C. perfringens* (CBAH-1) and the corresponding residues predicted in amino acid multiple sequence alignment with other BSHs (Fig. 4). The sterol moiety is bound primarily through hydrophobic interactions in the CBAH-1 (residues highlighted in gray in Fig. 4) as well as hydrogen bonds to the carboxylate group. Although the crystal structure of CBAH-1 did not reveal specific recognition of the taurine/glycine moiety, kinetic data from several BSHs suggest that the conjugates are important in substrate specificity (Table 1). Therefore, additional crystallization and site-directed mutagenesis (preferably with mutagenesis of Cys2) of BSHs from different species will be helpful in explaining the kinetic observations of substrate specificity.

**Distribution, genetic organization, and regulation of BSH**

Genes encoding BSHs have been cloned from *C. perfringens* (15), *Lactobacillus plantarum* (16), *La. johnsonii* (12, 17), *Bi. longum* (10), *Bi. bifidum* (18), *Bi. adolescentis* (19), and *Listeria monocytogenes* (20, 21). Homologs and putative *bsh* genes have also been identified recently through microbial genome analysis. The organization and regulation of genes encoding BSH differ between species and genera. Monocistronic BSH genes have been reported in *La. plantarum* (16), *La. johnsonii* (12), *Li. monocytogenes* (21), and *Bi. bifidum* (18). A gene encoding BSH (CBAH-1) cloned from *C. perfringens* (15) differed significantly in size and amino acid sequence from a BSH purified from a different strain of *C. perfringens* (13). The inactivation of the gene encoding CBAH-1 resulted in only partial reduction in BSH activity (BSH activity was 86% of that in the wild type), suggesting multiple BSH genes in *C. perfringens*. Furthermore, the crystal structure showed that the enzyme encoded by the CBAH-1 gene forms an active homotetramer (11). These observations, coupled with the detection of both intracellular and extracellular BSHs, provide further evidence for multiple isoforms, although the organization and regulation of the *bsh* gene(s) from *C. perfringens* are not known at present (22). Polycistronic operons encoding three genes involved in bile salt deconjugation (*cbsT1*, *cbsT2*, and *cbsH*) have been characterized in *La. johnsonii* and *La. acidophilus* (12). Genes *cbsT1* and *cbsT2* appear to be gene duplications that encode taurocholate/CA antiport proteins of the major facilitator superfamily, whereas *cbsH* encodes the BSH β-isofrom (23). In addition, an uncharacterized extracellular factor has been detected in *La. johnsonii* 100-100, which stimulates BSH activity and uptake of conjugated bile salts during the stationary growth phase (12, 24). BSH expression is also growth phase-dependent. Stationary phase expression has been reported in *Bacteroides fragilis* (25), and exponential phase expression was reported for *Bi. longum* (10).

**Benefits of BSHs to the bacterium**

BSHs appear to enhance the bacterial colonization of the lower gastrointestinal tract of higher mammals. The
Bile salt biotransformations by human intestinal bacteria

Physiological advantages of BSHs are not fully understood and may vary between bacterial species and genera. It has been hypothesized that deconjugation may be a mechanism of the detoxification of bile salts. De Smet et al. (26) observed significantly higher rates of deconjugation of glycodeoxycholate (GDCA) over TDCA in La. plantarum. Mutants lacking functional BSH (bsh-) exhibited pH- and concentration-dependent toxicity of GDCA compared with wild-type cells; this effect was not demonstrated with TDCA. De Smet et al. (26) hypothesized that the difference in dissociation constants between GDCA and TDCA resulted in the collapse of cellular proton motive force by intracellular protonation with the glycine conjugate. The presence of a functional BSH results in the intracellular accumulation of free bile acids, which become protonated in a stoichiometric manner, decreasing energy-lar accumulation of free bile acids, which become pro-

Deconjugation may provide a means of obtaining cellular carbon, nitrogen, and sulfur for some bacterial species. This has been demonstrated in bacteroides (34) and is suggested in Bi. longum (10). In fact, the bsh gene from Bi. longum is cotranscribed with the gene encoding glutamine synthetase adenyltransferase (glbE), a component of the nitrogen regulation cascade (10). In this regard, hydrolysis of the conjugated bile acid may provide amino nitrogen, providing a possible explanation for the coordinated regulation of these seemingly physiologically unrelated genes (10). Taurine utilization is also widespread and can serve as an energy source under both aerobic and anaerobic conditions (35). Glycine can be used as an energy source by certain clostridia by the Stickland reaction (36). The Stickland reaction is a form of amino acid fermentation in which one amino acid donates electrons that are accepted by another amino acid distinct from the electron donor. Another hypothesis suggests that BSHs are detergent shock proteins enabling survival during stress (37). De Smet et al. (26) found no evidence for this in lactobacilli after growth with various detergents.

The widespread distribution of BSHs across Gram-negative and Gram-positive intestinal bacteria coupled
Fig. 4. Multiple sequence alignment of cholangic hydrolases. Protein sequences were obtained from the National Center for Biotechnology Information (http://www.ncbi.nlm.nih.gov/). Alignments were made with the ClustalW program (http://www.ebi.ac.uk/clustalw/) using the GONNET 250 matrix. Residues highlighted in yellow are predicted active site amino acids based on the crystal structure of the BSH from *C. perfringens* (11) as well as on site-directed mutagenesis and biochemical data (10, 12–14). Residues highlighted in gray correspond to residues involved in substrate binding in the BSH from *C. perfringens* (11). The secondary structural elements, which are based on the conjugated bile acid hydrolase from *C. perfringens* (CBAH-1) crystal structure, are shown above the alignment. The α and β designations of *Lactobacillus johnsonii* refer to the two isoforms of the genes found in this bacterium.
with a wide range of substrate specificities, genetic regulation, and the occurrence of multiple isoforms in certain strains have created conflicting reports regarding the physiological benefit to the bacterium in hydrolyzing bile acid conjugates. Determining the mechanism(s) by which BSHs aid bacteria in the colonization of the mammalian intestine will be of great interest, especially with regard to bacterial pathogenesis.

### Taurine, hydrogen sulfide production, and colon cancer

The bile acid conjugates glycine and taurine serve as substrates in microbial metabolism. Unlike glycine, taurine contains a sulfonic acid moiety that is reduced and dissimilated to hydrogen sulfide after deconjugation (38, 39). Hydrogen sulfide is highly toxic and has been shown to increase colonocyte turnover (40). Activation and upregulation of the extracellular signal-regulated kinase 1 and 2 (ERK1/2) signaling pathway has been suggested as a possible mechanism for sulfide-induced colonocyte proliferation (41). Hydrogen sulfide also inhibits butyrate metabolism in colonocytes, a key nutrient and regulator of cell turnover in the gut (40). Levitt et al. (42) demonstrated that colonocytes have evolved a highly efficient mechanism to detoxify volatile reduced sulfides through oxidation to thiosulfate. Defects in this detoxification system are suggested to play a role in the pathogenesis of ulcerative colitis, a known risk factor for colon cancer (42, 43). Recently, sulfide was implicated in preventing apoptosis in the adenocarcinoma cell line HCT116 after exposure of cells to β-phenylethyl isothiocyanate, a phytochemical found in cruciferous vegetables, which has been shown to prevent colon carcinogenesis (44, 45).

A diet high in meat has been shown to significantly increase both the levels of taurine conjugation to bile acids (46, 47) and the production of hydrogen sulfide in the colon (48). A relationship exists between the generation of hydrogen sulfide in the colon and chronic GI illness, such as inflammatory bowel disease and colon cancer (5, 49). In human fecal slurries obtained from individuals consuming a “Western diet” (46, 47), the production of hydrogen sulfide was high and was shown to increase colonocyte turnover (40). Activation and upregulation of the extracellular signal-regulated kinase 1 and 2 (ERK1/2) signaling pathway has been suggested as a possible mechanism for sulfide-induced colonocyte proliferation (41). Hydrogen sulfide also inhibits butyrate metabolism in colonocytes, a key nutrient and regulator of cell turnover in the gut (40). Levitt et al. (42) demonstrated that colonocytes have evolved a highly efficient mechanism to detoxify volatile reduced sulfides through oxidation to thiosulfate. Defects in this detoxification system are suggested to play a role in the pathogenesis of ulcerative colitis, a known risk factor for colon cancer (42, 43). Recently, sulfide was implicated in preventing apoptosis in the adenocarcinoma cell line HCT116 after exposure of cells to β-phenylethyl isothiocyanate, a phytochemical found in cruciferous vegetables, which has been shown to prevent colon carcinogenesis (44, 45).  

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Although the extent to which taurine metabolism contributes to total colonic sulfide production has yet to be established, several key points have been made: 1) the extent of taurine conjugation in the bile acid pool is largely affected by diet; 2) the same dietary factors that increase taurine conjugation are hypothesized to increase colon cancer risk; 3) taurine metabolism by intestinal bacteria results in hydrogen sulfide generation; 4) sulfide generation is linked to the carcinogenesis process through enhanced cell proliferation, inhibition of butyrate metabolism, and activation of cell signaling pathways; and 5) sulfide generation may enhance DCA formation in the gut through stimulation of the microbial bile acid 7α-dehydroxylation pathway.

### MICROBIAL BILE ACID HYDROXYSTEROID DEHYDROGENASE(S)

#### Oxidation and epimerization

Oxidation and epimerization of the 3-, 7-, and 12-hydroxy groups of bile acids in the GI tract are carried out by hydroxysteroid dehydrogenase (HSDH) expressed by intestinal bacteria (Fig. 1). Epimerization of bile acid hydroxy groups is the reversible change in stereochemistry from α to β configuration (or vise versa) with the generation of a stable oxo-bile acid intermediate. Epimerization requires the concerted effort of two position-specific, stereochemically distinct HSDHs of intraspecies or interspecies origin. For example, the presence of both 7α- and 7β-HSDH in C. absonum allows epimerization by a single bacterium (51), whereas epimerization also can be achieved in cocultures of intestinal bacteria, one possessing 7α-HSDH and the other 7β-HSDH (52, 53).

The extent of the reversible oxidation and reduction of bile acid hydroxy groups by HSDH depends in part on the redox potential of the environment. Addition of oxygen to the culture medium increases the accumulation of oxo-bile acids (51). Generation of oxo-bile acids may be more favorable under the higher redox potentials found on the mucosal surface (4), whereas reduction of oxo-bile acids may be more favorable under the low redox potential (~200 to –300 mV) in the large intestinal lumen. Thus, although the redox potential of the colon is net reductive, microenvironments at the mucosa may provide oxidizing conditions favorable for certain microbial reactions. HSDHs differ in their reductive and oxidative pH optima, NAD(H) or NADP(H) requirements, molecular weight, and gene regulation (Table 2).

#### 3α- and 3β-HSDHs

3α/β-HSDHs specifically catalyze the reversible, stereospecific oxidation/reduction between 3-oxo-bile acids and 3α- or 3β-hydroxy bile acids. 3α-HSDHs have been detected in some of the most prevalent intestinal bacteria, including C. perfringens (54), Peptostreptococcus productus (55), and Eggerthella lenta (formerly Eubacterium lentum) (56, 57), as well as in intestinal bacteria present in lower numbers (~10^5/g wet weight of feces), including C. scindens (58) and C. hiranonis (59), and in nonintestinal bacteria, including Pseudomonas testosteroni (60, 61). 3β-HSDH activity has been described in species of Clostridium and Ruminococcus (62–64). It appears that intraspecies 3-epimerization favors the 3α-position. In fact, growing cultures of C. perfringens in the presence of 3-oxo-CDCA formed CDCA (84%) preferentially over iso-CDCA (16%) under anaerobic conditions (65).
Pyridine nucleotide cofactor requirements differ between \(3\alpha/\beta\)-HSDHs. \(3\alpha\)-HSDHs require NAD(H), with the exception of the enzyme purified from \(C.\) perfringens, which uses NADP(H), and that purified from \(C.\) scindens, which can use either NAD(H) or NADP(H) \((54, 56–58, 61)\). \(3\beta\)-HSDHs have been shown to preferentially require NADP(H), with the exception of \(C.\) innocuum, which uses NAD(H) \((62–64)\). Dihydroxy bile acids [DCA, CDCA, and ursodeoxycholic acid (UDCA)] are generally better substrates than trihydroxy bile acids (CA) \((62, 65)\).

\(3\alpha/\beta\)-HSDHs characterized to date are constitutively expressed with the exception of those from \(C.\) scindens and \(C.\) hiranonis, which are induced by the primary bile acids CA and CDCA. In fact, three copies of \(3\alpha\)-HSDH genes (the bile acid-inducible genes \(baiA1, baiA2,\) and \(baiA3\)) have been identified from \(C.\) scindens \((66, 67)\), and \(baiA1\) has been expressed in \(E.\) coli and characterized \((58)\). The \(baiA\) gene products are unique among \(3\alpha\)/\(\beta\)-HSDHs as a result of their high specificity toward CA-CoA and CDCA-CoA conjugates and relatively low activity toward free bile acids \((58)\).

### \(7\alpha\)- and \(7\beta\)-HSDHs

\(7\alpha/\beta\)-HSDHs catalyze the reversible, stereospecific oxidation/reduction of the \(7\alpha\) and \(7\beta\)-hydroxyl groups of bile acids. Although \(7\alpha/\beta\)-HSDHs are common among intestinal bacteria, the extent of \(7\alpha/\beta\)-dehydrogenation in the intestine, or in mixed fecal suspensions, is difficult to interpret because of the competing, and reversible, \(7\alpha/\beta\)-dehydroxylation of bile acids \((see\ below)\) \((68)\).

\(7\alpha/\beta\)-HSDHs are widespread among the bacteroides and clostridia as well as in \(E.\) coli and \(R.\) australis species \((Table\ 2)\) \((54, 64, 69–74)\). In addition, several intestinal clostridia express both \(7\alpha\)- and \(7\beta\)-HSDHs and have been shown to epimerize the \(7\alpha/\beta\)-hydroxy group \((75, 76–78)\). \(7\alpha/\beta\)-HSDHs have been partially purified from intestinal bacteria, including \(B.\) fragilis \((71, 79)\), \(B.\) thetaotaomicron \((74)\), \(C.\) scindens \((80)\), \(C.\) sordellii \((70)\), and \(E.\) coli \((73)\), as well as from the soil isolates \(X.\) maltophilia \((81)\), \(C.\) absonum \((82)\), and \(C.\) bifermentans \((83)\).

Interestingly, several intestinal bacteria with \(7\alpha/\beta\)-HSDH activity also possess BSH, including \(B.\) fragilis \((25)\), \(C.\) sordellii \((70)\), \(C.\) perfringens \((84)\), and \(C.\) innocuum \((63)\), as well as the soil isolate \(C.\) bifermentans \((83)\).

\(7\alpha\)-HSDHs generally use NADP(H) as a cofactor, with the exception of \(E.\) coli \((85)\) and \(B.\) thetaotaomicron \((74)\). \(C.\) bifermentans, \(C.\) absonum, and \(B.\) fragilis \(7\alpha\)-HSDHs can use either NAD(H) or NADP(H) as a cofactor \((76, 79, 83)\). \(7\beta\)-HSDH enzymes characterized to date use NADP(H) as a cofactor \((53, 62, 75, 76)\). \(7\alpha/\beta\)-HSDH enzymes have

### Table 2. Characteristics of HSDHs from intestinal bacteria

<table>
<thead>
<tr>
<th>Organism</th>
<th>Stereospecificity</th>
<th>Cofactor</th>
<th>pH Optimum</th>
<th>Native Molecular Mass</th>
<th>Gene Regulation</th>
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</table>

BA, bile acids; CA, cholic acid; CDCA, chenodeoxycholic acid; CE, cell extract; HSDH, hydroxysteroid dehydrogenase; I, induced; LCA, lithocholic acid; NI, noninduced by bile acids; ND, not determined; P, purified; PP, partially purified; R, repressed; UDCA, ursodeoxycholic acid.
higher affinity for dihydroxy bile acids (CDCA and 7-oxo-LCA) than for trihydroxy bile acids (CA and 7-oxo-DCA). C. limosum cell extracts use both free and conjugated bile acids, whereas whole cells only oxidize free bile acids (75, 86). This is attributable to the intracellular location of the 7α-HSDH and the inability of the organism to take up a conjugated bile salt. The genes encoding 7α-HSDHs have been cloned from E. coli (73), C. scindens (69), and C. sordellii (70). Sequence similarity suggests that these enzymes belong to the short-chain polyol dehydrogenase family. Regulation of 7α/β-HSDH expression is generally growth phase-dependent and inducible by bile acid substrates (Table 2). C. scindens and E. coli constitutively express 7α-HSDHs and are noninducible (69, 85). Unexpectedly, the nonsubstrate DCA can induce 7α-HSDH expression in C. absonum and C. sordellii, although the reason for this induction remains unclear (70, 76). Macdonald, White, and Hylemon (82) observed the expression of five soluble and two membrane polypeptides upon exposure of C. absonum to DCA and CDCA in the culture medium, although the functions of these additional polypeptides have not been determined.

Crystal structures of the E. coli 7α-HSDH binary and ternary complexes have been solved and a mechanism for 7α-dehydrogenation proposed (Fig. 5) (87). Binding of 7α-hydroxy bile acids elicits major conformational changes at the substrate binding loop and C-terminal domain. A two-step mechanism is proposed in which Tyr159 acts as a catalytic base removing the C-7 hydroxy hydrogen. Hydrogen bonding by serine 146 (Ser146) is hypothesized to stabilize the intermediate. Regeneration of the catalyst occurs through the transfer of the acquired hydride from the phenolic group at Tyr159 to lysine 163 (Lys163) to position 4 of NAD+. Lys163 serves two important roles: anchoring NAD+ through bifurcated hydrogen bonding, and indirect hydride transfer from Tyr159 to position 4 of NAD+ (87). Site-directed mutagenesis confirmed the role of these amino acids in 7α-HSDH catalysis (88). Analysis of the 7-oxo-GLCA bile acid substrate/enzyme complex revealed tight binding of the sterol with loose association for the glycine conjugate. Binding of glycine and taurine conjugates of CDCA was not significantly different from that of the free bile acid (87).

12α- and 12β-HSDHs

12α/β-HSDHs have been detected mainly among members of the genus Clostridium. NADP-dependent 12α-HSDHs have been detected in C. leptum (89) in Clostridium group P (90), whereas NAD-dependent 12α-HSDH activity was reported in E. lenta (56) and C. perfringens (54). 12β-HSDHs have been detected in C. tertium, C. difficile, and C. paraputrificum (91, 92). 12α/β-HSDHs characterized to date are constitutively expressed and noninducible, with the exception of the 12β-HSDH from C. paraputrificum, which is induced by 12-oxo-bile acid substrates (92). 12α/β-HSDHs generally have higher affinity for dihydroxy bile acids (DCA) than for trihydroxy bile acids (CA and iso-CA) and for free versus conjugated bile acids. The 12α-HSDH from C. leptum is an exception, demonstrating higher affinity for CA conjugates than for free CA (89). 12α-HSDHs appear to be repressed by the addition of bile acid substrates (DCA > CDCA > CA) to the growth medium at 1 mM concentrations. It has been suggested that these enzyme activities should be repressed in bacteria colonizing the large intestine (56, 92), although 12-oxo-bile acids have been detected at low levels in the feces of healthy individuals (Fig. 3) (93, 94).

Benefits of bile acid hydroxysteroid oxidoreductases to the bacterium

The oxidation of bile acid hydroxyl groups generates reducing equivalents for cellular biosynthetic reactions and possibly electron transport phosphorylation. Bile acid

![Fig. 5. Proposed catalytic mechanism of bile acid 7α-dehydrogenation based on the crystal structure and site-directed mutagenesis of active site amino acids of the 7α-dehydrogenation from E. coli. See text for a description of catalysis. Reprinted with permission from Tanaka et al. (87). Copyright © 1996 American Chemical Society.](https://drf.colorado.edu/)
dehydrogenation is hypothesized to generate energy in *Ba. thetaiotaomicron* (74). Sherod and Hylemon (74) suggested that reduced pyridine nucleotides generated from 7α-hydroxy oxidation serve to generate ATP via a cytochrome-linked electron transport chain in the presence of electron acceptors (i.e., fumarate).

Bile acids are potent antimicrobial agents provided that the proper concentration and proportion of hydrophobic bile acids (CDCA, LCA, and DCA) are present (95). Alteration of hydroxy group stereochemistry has a marked influence on the physiochemical properties of bile acids (96, 97). The epimerization of the 7α-hydroxy group of CDCA decreases the hydrophobicity and toxicity of the bile acid (97). Macdonald, White, and Hylemon (82) observed that *C. asbornum* grew on plates containing 1 mM UDCA, although it was unable to grow on plates containing 1 mM CDCA. Furthermore, when cultured in the presence of 7-oxo-bile acids, only low concentrations of CA and CDCA were formed, whereas the majority of 7-oxo-bile acids were reduced to 7-epicholic acid and UDCA, respectively, by log-phase *C. asbornum* (51). UDCA has also been shown to act as a repressor of 7α/β-HSDH production in *C. asbornum*, suggesting that UDCA is an end product (76). The enzyme also displays markedly higher affinity for CDCA than for CA, the former being more toxic. In summary, dehydrogenation may serve functions related to energy generation as well as attempts to maintain low concentrations of more hydrophobic bile acids in the bacterium’s microenvironment.

**Interplay between HSDH enzymes in human liver and intestinal bacteria**

The coevolution between host and gut flora is evident when observing the interplay between liver and bacterial biotransforming reactions. The liver synthesizes bile acids in which the hydroxy groups are in the α orientation. In the α-hydroxy orientation, one face of the molecule is hydrophobic and the other side is hydrophilic. This translates to efficient solubilization of lipid molecules through the formation of mixed micelles capable of efficient emulsiﬁcation while remaining soluble in aqueous environments. Generation of β-hydroxy bile acids by microbial enzymes alters the efficiency of micelle formation as a result of hydrophilic groups on both faces of the sterol molecule. The differences observed between the composition of bile acids in serum and bile are a result of the continual interplay between liver and bacterial enzymes. Exposure of bile salts to intestinal bacteria results in ~50% of bile acids requiring reconjugation and low levels of bile acids returned to the liver in the 3β-hydroxy orientation (98). Without a means of epimerizing bile acid hydroxy groups in the human liver, β-hydroxy bile acids would accumulate in the bile acid pool. Interestingly, the liver seems to “allow” the accumulation of UDCA (7β-hydroxy) in the biliary pool, as in the case of therpeutic administration of CDCA (99). The protective effects of UDCA observed in clinical studies (100) as well as cell culture studies (97, 101) may provide an evolutionary explanation for this phenomenon.

**THE BIOCHEMISTRY AND MOLECULAR BIOLOGY OF BILE ACID 7α/β-DHEDHYROXYLATION**

**Introduction**

Secondary bile acids (DCA and LCA) predominate in human feces (Fig. 3). Therefore, 7α-dehydroxylation is the most quantitatively important bacterial bile salt biotransformation in the human colon. The rapid rate of conversion of primary to secondary bile acids is surprising given current estimates that this metabolic pathway is found in ~0.0001% of total colonic flora (102–104). Human intestinal bacteria capable of bile acid 7α-dehydroxylation have been isolated (104, 105), and 16S rDNA phylogenetic analysis has led to their classification to the genus *Clostridium* (106–108).

Unlike bile acid oxidation and epimerization, 7α/β-dehydroxylation appears restricted to free bile acids. Removal of glycine/taurine bile acid conjugates via BSH enzymes is thus a prerequisite for 7α/β-dehydroxylation by intestinal bacteria (109–112). Some intestinal bacteria are capable of both 7α/β-dehydroxylation activities (113), whereas 7β-dehydroxylation activity is absent in other intestinal 7α-dehydroxylating bacteria (113). Epimerization of UDCA (7β-hydroxy) to CDCA (7α-hydroxy) via 7β-HSDHs produced by members of the gut flora results in subsequent 7α-dehydroxylation. In this regard, it appears that the presence of 7β-dehydroxylation activity is more of a luxury than a necessity.

**Elucidating the bile acid 7α/β-dehydroxylation pathway**

Samuelsson (114) administered [6α-3H,6β-3H,8β-3H] [24-14C]CA to bile duct-cannulated rabbits and rats. Analysis of the products recovered after exposure to intestinal bacteria revealed a differential loss of the 6β-3H during 7α-dehydroxylation of CA. Previous work showed complete retention of the 7β-3H in [7β-3H][24-14C]CA during 7α-dehydroxylation in the rat intestine (115, 116). These data led Samuelsson (114) to propose a mechanism for CA 7α-dehydroxylation involving two steps: diaxial trans-elimination of the 7α-hydroxy group and 6β-hydrogen atom, followed by reduction through trans-hydrogenation of the 6β and 7α positions of the cholen-6-oic acid intermediate forming DCA. Björkhem et al. (117) showed the formation of a 3-dehydro-4-cholenoic acid intermediate after the differential loss of the 5βH in vitro and in vivo using [3β-3H][24-14C]CA and [5β-3H][24-14C]CA. Hylemon et al. (118) subsequently observed the accumulation of multiple bile acid intermediates in cell extracts of *C. scindens* induced by CA (Fig. 6). These radiolabeled CA intermediates were identified by mass spectrometry, then chemically synthesized and added to cell extracts of CA-induced *C. scindens*. Each 24[14C]CA intermediate was converted to 24[14C]DCA in cell extracts prepared from CA-induced cultures of *C. scindens*. These observations suggested that the 7α-dehydroxylation mechanism was more complex than the two-step mechanism proposed by Samuelsson (114). Furthermore, these data demonstrated that bile acid 7α-dehydroxylation was a multistep pathway in *C. scindens* and suggested the presence of multiple bai genes.
The induction of 7α-dehydroxylation activity in *C. scindens* by unconjugated C24 primary bile acids resulted in the appearance of several new polypeptides, as observed by one- and two-dimensional SDS-PAGE (119, 120). Purification and N-terminal sequencing of these *bai* polypeptides facilitated the cloning of *bai* genes through the design of degenerate probes (121–123). Northern blot analysis indicated the presence of a large CA-inducible (>10 kb) mRNA transcript and a smaller transcript (≤1.5 kb) in *C. scindens* (121, 124). These studies led to the discovery of a *bai* regulon encoding at least 10 open reading frames (Fig. 7). Individual *bai* genes have been subcloned into *E. coli* and the functions of many of them determined (58, 67, 123, 125–130; P. B. Hylemon, unpublished data). The proposed bile acid 7α/β-dehydroxylation pathway in *C. scindens* is shown in Fig. 8. A *bai* operon has also been characterized from *C. hiranonis* (59), although the discussion below of the 7α/β-dehydroxylation pathway will center on *C. scindens*, from which the functions of the gene products have been determined.

**bai genes: a regulon for 7α/β-dehydroxylation**

The transport of unconjugated primary bile acids into *C. scindens* is facilitated by the *baiG* gene product, which belongs to a major pump/facilitator superfamily of protein transporters (126). The *baiG* gene has been cloned into *E. coli* and shown to encode a 50 kDa H⁺-dependent bile acid transporter (126). BaiG facilitates the transport of unconjugated CA and CDCA but not of the secondary bile acids DCA and LCA (126). Computer-aided modeling suggests that the *baiG* polypeptide contains 14 membrane-spanning domains (126).

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**Fig. 6.** Accumulation of [24-14C]CA intermediates during 7α-dehydroxylation in cell extracts of *C. scindens* VPI 12708. Cell extracts were prepared from either CA-induced (I) or control (C) cells. This figure was modified from Hylemon et al. (118). Copyright © 1992 American Society for Biochemistry and Molecular Biology, Inc.

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**Fig. 7.** Gene organization of the bile acid-inducible (*bai*) 7α/β-dehydroxylation operons characterized in *C. scindens* VPI 12708. P indicates the promoter region.
After transport, ligation to CoA is the first step in activating CA and CDCA for 7α-dehydroxylation, as several subsequent enzymatic steps are specific for CoA conjugates. The **baiB** gene was shown to encode a 58 kDa bile acid CoA ligase (125). CoA ligation is ATP-, CoA-, and Mg2⁺-dependent and also requires a free carboxyl group on C24 bile acids (125). CoA ligation may function both to sterically hinder the constitutive 7α-HSDHs, committing the bile acid to 7α-dehydroxylation, and to trap the bile acid inside the cell.

The 3α-hydroxy group is oxidized after CoA ligation. Oxidation of the 3-hydroxy group inhibits 7α-hydroxy group dehydrogenation, favoring 7-dehydroxylation over constitutively expressed 7α-HSDHs in *C. scindens* (58, 80). The **baiA** gene products encode 27 kDa polypeptides that have significant similarity with the short-chain alcohol/polyol dehydrogenase gene family (58, 124). Amino acid multiple sequence alignment and comparison between **baiA** gene products and other members of the short-chain alcohol dehydrogenase family revealed a possible NAD(P) binding site and catalytic active site (58). Three **baiA** genes have been cloned from *C. scindens*; the **baiA1** and **baiA3** genes are monocistronic, whereas the **baiA2** gene is part of the polycistronic **bai** operon (66, 67, 124). The **baiA** genes from *C. scindens* were cloned in *E. coli* and shown to encode 3α-HSDHs (58). The enzymes only recognize bile acid CoA conjugates and can use either NAD⁺ or NADP⁺ as electron acceptors (58). Interestingly, the different 3α-HSDHs of *C. scindens* share 92% amino acid sequence identity with one another, suggesting gene duplication. The physiological importance of multiple **baiA** genes remains unclear.

The **baiCD** gene from *C. scindens* has been cloned and expressed in *E. coli* and was recently demonstrated to encode a steroid oxidoreductase specific for the CoA conjugates of 3-dehydro-4-cholenoic acid and 3-dehydro-4-chenodeoxycholenoic acid (Fig. 8) (P. B. Hylemon et al., unpublished data). The ~70 kDa enzyme requires FMN and NAD⁺ or NADP⁺ for activity and shows stereospecificity toward 7α-hydroxy bile acids. The **baiCD** polypeptide shows considerable amino acid sequence identity with the Old Yellow Enzyme family, a putative NADH oxidase from *Li. monocytogenes*, several 2,4-dienoyl CoA reductases, and **baiH** from *C. scindens* (121, 129). The **baiH** gene en-
codes a 72 kDa polypeptide containing 661 amino acids (121). The BaiH protein exists as a homotrimer with NADH:flavin oxidoreductase activity (121). The baiH gene has been subcloned into E. coli, purified, and shown to contain 1 mol of FAD, 2 mol of iron, and 1 mol of copper per mole polypeptide subunit (129). The baiH gene product from C. scindens was recently determined to encode a steroid oxidoreductase specific for CoA conjugates of 3-dehydro-4-ursodeoxycholenoic acid and 3-dehydro-4-epicholenoic acid (Fig. 8) (P. B. Hylemon et al., unpublished data). Oxidation of CA-CoA, CDCA-CoA, and UDCA-CoA to their respective 3-dehydro-4-bile acid CoA conjugates appears to make the bile acid chemically labile for 7α/β-dehydration.

The baiF gene encodes a 47.5 kDa polypeptide containing 426 amino acids that was shown to have bile acid CoA hydrolase activity (122, 130). However, baiF is hypothesized to encode a CoA transferase because of energy conservation (Fig. 8) and homology to the type III family of CoA transferases (131). The first few cycles of 7α-dehydroxylation would require ATP hydrolysis (Fig. 8), although ATP-independent recycling of the thioesterase intermediates via transfer of CoA from 3-dehydro-4-cholenoic acid to CA, 3-dehydro-4-chenodeoxycholenoic acid to CDCA, or 3-dehydro-4-ursodeoxycholenoic acid to UDCA by the baiF would significantly conserve energy. However, this hypothesis remains to be tested.

7α-Dehydration of 3-dehydro-4-cholenoic acid and 3-dehydro-4-chenodeoxycholenoic acid results in the generation of a conjugated double bond in rings A and B, forming stable 3-dehydro-4,6-deoxycholenoic acid and 3-dehydro-4,6-lithocholenoic acid intermediates, respectively. The 7α-dehydration step results in the largest calculated energy change (~9.4 kcal/mol) of any reaction in this pathway, and the reverse reaction was not detected in vitro (128). The 19.5 kDa bile acid 7α-dehydratase is encoded by the baiE gene (128). This enzyme showed no activity with 3-dehydro-4-ursodeoxycholenoic acid (128). The baiE gene product was modeled on the crystal structure of the protein homologs (secondary structure) ketosteroid isomerase and scytalone dehydratase (Fig. 9) (K. Woodford et al., unpublished data). The putative active site/binding pocket is shown in Figure 10. A catalytic mechanism has been proposed for the 7α-dehydration step based on the conservation in secondary structure and site-directed mutagenesis of the key active site amino acids. Tyr30 acts as a general acid withdrawing electron density from the 3-oxo group. This shift in electron density is hypothesized to make the 6β-hydrogen labile for removal by the general base histidine 83 (His83) (assisted by Asp35). His83 is thought to donate its hydrogen to the 7α-hydroxy forming the water-leaving group that is stabilized by Asp106. Site-directed mutagenesis based on the proposed mechanism supports the important role of the putative active site amino acids in enzyme catalysis. It is hypothesized that the baiE gene encodes a bile acid 7β-dehydratase, because of amino acid sequence homologies between the baiE and baiI gene products.

Genes involved in the reductive arm of the 7α/β-dehydroxylation pathway have not been isolated. These genes should encode oxidoreductases catalyzing the reduction of 3-dehydro-4,6-deoxycholenoic acid to 3-dehydro-4-deoxycholenoic acid to 3-dehydro-deoxycholic acid to DCA as well as a bile acid exporter to remove secondary bile acid end products from the bacterium (Fig. 8). Genes encoding putative transcriptional regulators have been detected upstream of the bile acid-inducible promoter region (Table 3) (D. H. Mallonee and P. B. Hylemon, unpublished data). Additional studies will be required to determine the mechanism of induc-
tion/repression of this pathway and to identify additional bai genes.

The benefits of \( \text{7a}/\beta \)-dehydroxylation to the bacterium

The ability to use bile acids as electron acceptors is an important niche for \( \text{7a} \)-dehydroxylating bacteria in the human colon. The \( \text{7a}/\beta \)-dehydroxylation pathway requires multiple oxidative and reductive steps with a net 2 electron reduction (Fig. 8). The hypothesized energy benefits of this pathway assume, however, that the \( \text{baiF} \) gene encodes a CoA transferase and the toxic end products, the secondary bile acid, are removed from the microenvironment in vivo (precipitation and binding to insoluble fiber). The generation of secondary bile acids may also function to exclude bacteria sensitive to these hydrophobic molecules.

SECONDARY BILE ACIDS AND DISEASE

In humans, DCA accumulates in the bile acid pool to high levels in some individuals. An increase in DCA in the bile acid pool is associated with a decrease in CDCA (Fig. 11). Unlike rodents, the human liver cannot \( \text{7a} \)-hydroxylate DCA, forming CA. Hence, under normal physiological conditions, there is no metabolic pathway for removing DCA from the bile acid pool in humans. The amount of DCA in the bile acid pool is a function of at least three variables: 1) the rate of formation and absorption of DCA through the colon (input) (132); 2) colonic transit time (133); and 3) colonic pH (134).

High levels of DCA in blood, bile, and feces have been correlated with an increased risk of cholesterol gallstone disease and colon cancer, two major diseases of Western society (5, 135). High levels of CA \( \text{7a} \)-dehydroxylating fecal bacteria have been correlated with increased amounts of DCA in bile of a subset of cholesterol gallstone patients (132). Treatment of these cholesterol gallstone patients (high DCA group) with antibiotics significantly decreased the levels of fecal CA \( \text{7a} \)-dehydroxylating bacteria, DCA in bile, and the cholesterol saturation index in bile (132). Early studies by Low-Beer and Nutter (135) reported that treating control individuals with metronidazole, an antibiotic effective against anaerobic bacteria, significantly decreased the cholesterol saturation index of bile. Moreover, excess DCA in bile has been reported to decrease the nucleation time for cholesterol crystallization (136, 137). In total, these results suggest a possible link between intestinal bacteria, DCA, and the risk of cholesterol gallstone disease in some patients.

DCA and LCA have been linked to colon carcinogenesis in a number of laboratory animal models and human epidemiological studies (for reviews, see 5, 138). Most animal studies conclude that DCA is a promoter of carcinogenesis (139–142). However, some researchers argue

\[
\begin{array}{|c|c|c|c|c|}
\hline
\text{bai} & \text{Molecular Mass} & \text{Catalytic Activity/Function} & \text{Gene Family} & \text{Reference} \\
\hline
\text{baiA} & 27 & \text{3a-HSDH} & \text{Short-chain alcohol/polyol dehydrogenase} & 58 \\
\text{baiB} & 58 & \text{Bile acid CoA ligase} & \text{AMP binding} & 125 \\
\text{baiCD} & 70 & \text{3-Dehydro-4-CDC/CA} & \text{Pyridine nucleotide-disulfide oxidoreductase; NADH:flavin oxidoreductase} & -^a \\
\text{baiH} & 72 & \text{3-Dehydro-4-UDCA/7-epiCA} & \text{Pyridine nucleotide-disulfide oxidoreductase; NADH:flavin oxidoreductase} & 129 \\
\text{baiE} & 19.5 & \text{7a-Dehydrogenase} & \text{COG4875} & 128 \\
\text{baiI} & 22 & \text{7\beta-Dehydrogenase} & \text{COG4876} & 168 \\
\text{baiF} & 47.5 & \text{Bile acid CoA hydrolase} & \text{Type III CoA transferase} & 130 \\
\text{baiG} & 50 & \text{H\textsubscript{+}-dependent bile acid transporter} & \text{Major facilitator superfamily} & 126 \\
\text{baiA} & 46 & \text{Transcriptional regulation} & \text{AraC/XylS} & -^a \\
\text{baiB} & 22 & \text{Transcriptional regulation} & \text{RpoB; permeases of the major facilitator superfamily} & -^a \\
\hline
\end{array}
\]

\( ^a \) P. B. Hylemon et al., unpublished data.

\( ^b \) Hypothesized function.

Fig. 11. Relationship between the percentage of CDCA and DCA in bile of patients at McGuire VA Hospital (Richmond, VA) (P. B. Hylemon et al., unpublished data).
that bile acids may cause DNA damage and act as carcinogens in humans (138). Higher levels of DCA are found in the blood of colon cancer patients compared with control patients (98, 143). Moreover, DCA is a logical candidate for promoting colon carcinogenesis for the following reasons: 1) it is found in fecal water in high concentrations (>100 μM) (138); 2) it can cross biological membranes via passive diffusion; and 3) it can activate mammalian cell signaling pathways that are known to be involved in promoting carcinogenesis. In this regard, cell signaling pathways activated by DCA in mammalian epithelial cells include protein kinase C (144), ERK1/2 via the epidermal growth factor receptor (101, 145, 146), β-catenin (147), and Jun-N-terminal kinase 1 and 2 (JNK1/2) (148).

Secondary bile acids have been shown to cause apoptosis in colonic epithelial cells, and high concentrations of DCA and LCA in stool may promote carcinogenesis by exerting selective pressure for the emergence of epithelial cell mutants that are resistant to apoptosis (e.g., via loss of p53) (138). LCA has been found to be an excellent activator of the vitamin D receptor (149, 150). Activation of this receptor in intestinal epithelial cells activates genes that metabolize LCA (150). This may be a protective mechanism that evolved to limit LCA toxicity to intestinal epithelial cells.

**DISCUSSION**

Bile salt metabolism is a widespread and fundamental property of the gastrointestinal microflora encompassing the most commonly isolated species of intestinal bacteria, including but not limited to the genera *Bacteroides*, *Clostridium*, *Lactobacillus*, *Bifidobacterium*, *Eubacterium*, and *Escherichia*. However, our current understanding of the intestinal microbiology of bile salt modifications is limited to cultivated species. Discovering novel genes in the gut microbiome (collective genomes of the gastrointestinal flora) encoding BSHs, HSDHs, and enzymes involved in the 7α/β-dehydroxylation pathway through molecular techniques will facilitate a much greater understanding of the diversity and complexity of these reactions in the human colon. Techniques such as PCR-denaturing gradient gel electrophoresis can be used to measure the diversity of organisms based on specific phylogenetic markers, such as 16S rDNA, functional genes, and potentially bile salt-modifying enzymes (151, 152). Measuring the true diversity of bile salt-modifying bacteria is crucial in studying the relationship between the levels and activities of these bacteria and disease risk.

Determining the conditions in which secondary bile acids are formed in significant quantities and retained in the enterohepatic circulation of certain individuals is suggested to be important in the etiology of cholesterol gallstone disease and colon cancer. Because secondary bile acids are formed exclusively through bacterial enzymatic reactions, the study of microbes capable of bile acid 7α/β-dehydroxylation is important in understanding these chronic GI illnesses. The goal of such research is to find ways to block the source of secondary bile acid production. Long-term use of antibiotics to prevent 7α-dehydroxylation of bile acids would be impractical. The design of pharmaceuticals to block the 7α-dehydroxylation pathway is a possibility. However, this approach requires targeting microbial enzymes and runs the risk of eventual drug resistance, as well as antibiotics. Alternatively, reducing secondary bile acid production may be achieved by administering specialized CA-accumulating probiotic bacteria (153). Studies with bifidobacteria and lactobacilli isolated from human feces have shown their ability to assimilate CA spontaneously in vitro (153, 154). The mechanism for CA uptake in lactic acid bacteria appears to be diffusion of a hydrophobic weak acid through the membrane via the transmembrane proton motive force (153, 154). The higher intracellular pH causes the bile acids to become trapped as a result of ionization.

The use of specific bacteria as “drugs” to treat chronic GI illnesses caused in part by other intestinal bacteria certainly has potential, although rigorous studies are needed to demonstrate the effectiveness of such therapies. Overall, clinical trials using probiotics to solve specific health disorders have met with mixed results. Discrepancies between studies are attributable in part to different methodologies, choice of probiotic strains, colony-forming units administered per day, and patient characteristics. One particular variable that would have to be addressed when determining bile acid assimilation by lactic acid bacteria in clinical trials is viability in vivo after passage through the gastric juice and bile. A bacterium must be alive to create a membrane potential capable of accumulating bile acids. Targeted delivery of probiotic bacteria to the intestine in microencapsulated form has shown promise in improving viability in the presence of gastric conditions and bile (155, 156). Another issue is whether the introduction of billions of probiotic bacteria to the small bowel will have an effect on bile acid input into the colon. Significant bile salt hydrolysis proximal to the terminal ileum reduces the efficiency of bile salt uptake through high-affinity transport, allowing enhanced excretion of bile acids in feces. This principle is behind attempts to decrease serum cholesterol using probiotics with bile salt hydrolytic activity (157–159). Thus, the choice of probiotic delivery mechanisms is important in addressing issues of bacterial viability and in preventing increased bile salt hydrolysis proximal to the terminal ileum. The potential impact of probiotic bacteria in reducing secondary bile acid formation in the human colon by sequestering bile acids would be greatly reduced if these same bacteria cause an increase in bile acid input into the large intestine greater than their capacity for sequestering bile acids.

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