ABCG5 and ABCG8: more than a defense against xenosterols

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Abstract The elucidation of the molecular basis of the rare disease, sitosterolemia, has revolutionized our mechanistic understanding of how dietary sterols are excreted and how cholesterol is eliminated from the body. Two proteins, ABCG5 and ABCG8, encoded by the sitosterolemia locus, work as obligate dimers to pump sterols out of hepatocytes and enterocytes. ABCG5/ABCG8 are key in regulating whole-body sterol trafficking, by eliminating sterols via the biliary tree as well as the intestinal tract. Importantly, these transporters keep xenosterols from accumulating in the body. The sitosterolemia locus has been genetically associated with lipid levels and downstream atherosclerotic disease, as well as formation of gallstones and the risk of gallbladder cancer. While polymorphic variants raise or lower the risks of these phenotypes, loss of function of this locus leads to more dramatic phenotypes, such as premature atherosclerosis, platelet dysfunction, and thrombocytopenia, and, perhaps, increased endocrine disruption and liver dysfunction. Whether small amounts of xenosterol exposure over a lifetime cause pathology in normal humans with polymorphic variants at the sitosterolemia locus remains largely unexplored. The purpose of this review will be to summarize the current state of knowledge, but also highlight key conceptual and mechanistic issues that remain to be explored.

The description of the disease sitosterolemia by Bhatcharyya and Connor initiated a fundamental rewrite of how dietary sterols traffic and are eliminated by the body (1). And while sitosterolemia, named after the most abundant xenosterol detected in the plasma of the two affected sisters, was described in 1974, the disease is ancient, with mutant alleles that are several thousand years old (2). We know that the genetic locus whose dysfunction leads to sitosterolemia encodes two genes, ABCG5 and ABCG8, whose proteins function as obligate heterodimers (3, 4). ABCG5 and ABCG8 (also known as sterolins) are expressed only in hepatocytes, gallbladder epithelium, and enterocytes, and are responsible for excretion of sterols, with xenosterols preferred over cholesterol. Arguably, the naming of the disease has led to a bias in appreciating the fact that sterols keep a vast range of xenosterols, not just sitosterol, from accumulating in the body (5). A better name should be xenosterolemia to reflect appreciation of this biology. Although the spectrum of xenosterols our diets contain is extensive, we show a few sterol structures that highlight the differences between cholesterol and these xenosterols (Fig. 1). In general, the major differences reside in the R tail of the sterol structure, with change in shape that will likely impact their organization in membranes (6–8).

While the importance of sterol trafficking is well-recognized for its roles in the pathogenesis of endothelial dysfunction, foam-cell formation, and atherosclerosis, ensuring that xenosterols are kept out and why [an evolutionary mechanism, conserved from fish to man (9)], has garnered less attention. The loss of function in humans, as well as in animal models, shows that accumulation of

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xenosterols leads to dramatic phenotypes, such as macrothrombocytopenia and platelet dysfunction, liver disease, and cholesterol accumulation with xanthoma formation and atherosclerosis (10), and in mouse models (but not humans), infertility, immune dysfunction, and cardiomyopathy have been reported (11–13). This begs the question whether a lifetime of low level exposure to dietary bioactive xenosterols, whose levels of entry and retention may be altered by polymorphisms in \textit{ABCG5} and \textit{ABCG8}, may have biological consequences. This review will aim to highlight areas that, therefore, need further exploration.

\textbf{GENETIC VARIATION AT THE SITOSTELEOMIA LOCUS AND CLINICAL IMPLICATIONS}

\textit{ABCG5} and \textit{ABCG8} were key suspects in regulating cholesterol homeostasis in atherosclerotic CVD (ASCVD). This was supported by the clinical observations of severe, and sometimes fatal, atherosclerosis in humans with sitosterolemia (though see below). Support for this contention came from the genome-wide association studies, which showed a link not only between genetic variations at sitosterolemia locus (previously referred to as \textit{STSL}) and lipid levels, but also for coronary artery disease (14–16). And not surprisingly, because the \textit{ABCG5}/\textit{ABCG8} heterodimer pumps cholesterol into bile (17–21), this locus was shown to predispose to gallstone disease (22–28). What is surprising, and the mechanistic connection remains to be addressed, was the link between a presumed hypermorphic variant, D19H, in \textit{ABCG8} and gallbladder cancer risk (25, 29). This locus is one of the strongest known genetic risk factors for this very rare cancer (though the genetic risk remains small). One pathway of how this may occur is that this variant leads to increased gall stone formation, subsequent inflammation, and consequently a predisposition to gallbladder cancer; this explanation remains unsubstantiated. Cancer, by its nature, requires multiple genetic changes to occur and how this would be caused or accelerated by the sitosterolemia locus remains to be explored. It is also not clear whether the variant alters the kinds of xenosterols that are concentrated in the bile leading to the formation of gallstones, prolonged inflammation, generation of genotoxic agents, and then oncogenesis.

Studies of these genetic variants have been association studies (30) and none have demonstrated a specific mechanistic pathway altered by these variants; we assume that cholesterol pumping into the bile or intestine would be altered. However, the liver has a predilection for excretion of xenosterols into bile (31), and the hypothesis that one of these dietary xenosterols is mechanistically involved has not been explored.

\textbf{CLINICAL MANIFESTATIONS}

The many clinical manifestations of sitosterolemia have been well-documented in a number of reviews (10, 32, 33). In summary, asymptomatic subjects (typically identified by cascade screening once a proband has been identified) can manifest no symptoms or signs to varying levels of atherostenoses. There is no question whether affected individuals can have severe ASCVD, sometimes with fatal presentations (34–38), but with increased knowledge and case identification, many affected subjects have now been identified, who in the untreated state, seem to have very little evidence of ASCVD. Two new studies have now highlighted this. In five subjects who were diagnosed after investigation of hyperlipidemia (ages 19–32 years), no clinical evidence of ASCVD was identified (39). More recently, and provocatively, cascade screening in the Hutterites (where all of the cases have the same mutations and share a similar lifestyle), many subjects were identified with no overt clinical manifestation of ASCVD or any blood dyscrasia (40). The latter study is also remarkable, as it may shed light on an old observation by Salen and his colleagues who reported that hepatic LDL-receptor expression was increased, in contrast to the paradoxical suppression of cholesterol biosynthesis enzymes (41, 42). We had shown that children with sitosterolemia can manifest very high levels of cholesterol and had previously been misdiagnosed as having pseudo-homozygous familial hypercholesterolemia (43, 44), yet it is not clear...
why they have such very high levels of plasma cholesterol, a manifest defect in hepatic clearance of LDL particles. Mymin et al. (40) reported a nonsignificant inverse association between age and plasma cholesterol; when subjects were grouped by pre- or postpuberty, both plasma cholesterol and sitosterol levels were significantly higher in prepubertal compared with postpubertal subjects. In one subject, in the absence of treatment, these sterol levels also fell progressively with time, though never normalized (40). This suggests that hepatic expression of lipoprotein receptors is dramatically altered by the progressive accumulation of xenosterols, and this process is influenced by puberty.

Macrothrombocytopenia

Macrothrombocytopenia is another phenotype that has been observed in humans with sitosterolemia (45–47). Remarkably, this phenotype can be the only manifestation of sitosterolemia. The first description of this entity, Mediterranean macrothrombocytopenia, was reported in Australia (48), and we suspect it was a cohort of subjects who had sitosterolemia; all of these subjects have been lost to follow-up to allow confirmation. However, Rees et al. (45) showed that not only were plant sterols elevated in their cohort in Europe, they also confirmed mutations at the sitosterolemia locus as causative, a finding that has been now reported from many other sites around the world. This platelet phenotype is also observed in the murine models of sitosterolemia (46, 49, 50). Mutations seen in macrothrombocytopenia are also seen in sitosterolemia without the platelet phenotype, suggesting that these manifestations may be dependent upon the type of xenosterol exposure. While there is no doubt that xenosterol accumulation leads to platelet dysfunction (50), biogenesis of the platelet is also abnormal. The mechanism of how xenosterolemia affects platelet biogenesis remains to be explored. Additionally, it is assumed that these effects are from sitosterol accumulation (the most abundant accumulating xenosterol), but more bioactive less abundant xenosterols could be responsible. Other possibilities are bulk xenosterols altering raft membrane structures, or as a result of displacing cholesterol from key active sites (51).

Liver disease

Although the first case of liver failure associated with sitosterolemia (52) was regarded as an outlier or coincidence (and was only interesting as liver transplantation cured the sitosterolemia), a second case with a fatal outcome has now been reported. Bazerbach et al. (53) reported a case of idiopathic cirrhosis, denied heart transplant from severe coronary artery disease, only to discover that the patient had sitosterolemia with the implication that both organ failures were caused by his one underlying condition. This raises the issues of the underlying mechanism for liver dysfunction and failure, and how many idiopathic cirrhosis cases are undiagnosed sitosterolemia. Mutational reviews of the two cases do not indicate any uniqueness when compared with others with sitosterolemia, and accumulation of misfolded proteins in the ER, akin to α1-antitrypsin deficiency, seems less likely (53). The Finnish case was heterozygous for mutations in ABCG8 (W361X and E423D) (52), but the US case was homozygous for premature stop codon in ABCG5 (R46X) (53). If so, then the hypothesis of toxic xenosterol accumulation is again raised. And if it is a toxic xenosterol species, it may be hard to detect as most sterol profiles have, to date, been limited to reporting sitosterol, campesterol, and stigmasterol. Other lesser known sterols are rarely reported. Expression of these mutants in vivo may be required to explore these possibilities.

Transintestinal cholesterol excretion in humans

Until recently, it was assumed that an elevation of plant sterols in the blood was diagnostic of sitosterolemia. Even in cases of liver dysfunction and failure, plant sterols remain low and the only clinical scenario where they are elevated is during administration of total parenteral nutrition, which contains large amounts of plant sterols (54, 55). This has now been challenged by a report of a case of primary biliary cirrhosis (PBC), where plant sterols were dramatically elevated (56). It should be noted that plant sterols are not normally elevated in PBC, thus this is a unique case. While the speculation was that the liver sterolin function was affected by this autoimmune process (56), it should be noted that ABCG5/ABCG8 should remain functionally active in the intestine, and they are key to transintestinal cholesterol excretion (TICE). Thus, TICE should have resulted in maintaining lower plant sterol levels [akin to the restoration of intestinal sterolin function by transgenes in the Abcg8 knockout mice (12), or in liver-only knockouts (57)], but it did not. In PBC, the secretion of bile into the biliary tree is dramatically reduced and perhaps in this case there may have been very little bile secreted; this would mean that for TICE to be effective, loss of all bile in the intestine may also impair TICE. Green and colleagues have shown that the phospholipid content, rather than bile salts, was a major regulator of TICE (58), opening the avenue for therapeutic use of dietary phospholipids in cholestatic diseases to improve lipids via TICE and ABCG5/ABCG8. And in passing, it should be noted that TICE should be TISE, because sterols are excreted and continue to bias interpretation in a cholesterol-centric manner.

REGULATION OF ABCG5/G8

The secretion of cholesterol into bile is coupled to the secretion of phospholipids and bile acids that forms the mixed micelle acceptors for cholesterol within the aqueous lumen of the canaliculus. Mouse models of ABCG5/G8 deficiency suggest that the complex accounts for 70–90% of biliary cholesterol secretion (17, 20, 21), and in mouse models with a gain or loss of function, there is an approximate gene dosage effect on biliary sterol excretion (17, 18, 20, 59). Factors that increase or decrease the coupling of biliary cholesterol secretion to bile acids and phospholipids are likely to be ABCG5/G8 dependent and reflect changes in either ABCG5/G8 abundance or activity, although it
should be noted that ABCG5/G8-independent biliary cholesterol secretion has been reported (60, 61).

### Transcriptional regulation of ABCG5/G8

Expression of sterolins is restricted to the liver, small intestine, and gallbladder epithelium, but the mechanisms responsible for this limited tissue distribution have not been elucidated (3, 62, 63). Within these tissues, ABCG5 and ABCG8 share a common bidirectional promoter of 374 base pairs that separates their initiation codons (3, 62, 64). Regulatory elements within the intergenic region have been identified for hepatocyte nuclear factor 4a (NR2A1), liver receptor homolog-1 (NR5A2), nuclear factor kappa-light-chain-enhancer of activated B cells (NFκB), and Forkhead box protein O1 (64–68). Liver X receptor (LXR) response elements have been mapped to two conserved regions roughly 100 kbp distal to the intergenic promoter, shown to bind both LXRα (NR1H3) and LXRβ (NR1H2) and synergistically activate the ABCG5/G8 promoter in combination with retinoid X receptor α (NR2B1), GATA4, hepatocyte nuclear factor 4a, and liver receptor homolog-1 (69). These findings corroborated earlier data in animal models demonstrating that ABCG5/G8 mRNAs were responsive to dietary cholesterol as well as LXR and retinoid X receptor agonists in a LXRω-dependent fashion (70). Farnesoid X receptor (FXR; NR1H4) receptor agonists have also been shown to regulate ABCG5/G8 mRNAs in cultured hepatocytes, suggesting a direct mechanism of action. However, FXR response elements have yet to be mapped for ABCG5/G8 (70, 71). Thus, while a number of transcriptional factors are implicated in regulating the sitosterolemia locus, there remains a paucity of direct evidence of their roles.

In vivo, sterolin mRNAs and proteins can be modulated by a number of experimental conditions. Disruptions of bile acid metabolism alter hepatic ABCG5/G8 expression, presumably due to changes in bile acid signaling through FXR (72–74). Similarly, intestinal expression of ABCG5/G8 is modulated by bile acids (75, 76). Tissue-specific deletion of hepatic insulin receptors revealed a role for insulin signaling in the regulation of ABCG5/G8 through Forkhead box protein O1 (67). Alterations in ABCG5/G8 mRNAs have also been observed in experimental models of type 1 and type 2 diabetes, as well as in human diabetic subjects (77–81). Additional factors shown to alter hepatic, intestinal, or gallbladder ABCG5/G8 expression include thyroid hormone signaling, dietary calcium, iron depletion, constitutive androstane receptor agonists, and gallbladder disease (82–86). However, the molecular mechanisms underlying these effects are unclear and the extent to which they reflect novel regulatory nodes for ABCG5/G8 or are secondary to disruptions in established regulators is not known.

CpG islands have been identified near the transcriptional start sites for ABCG5 and ABCG8 and were hypomethylated in the liver and hypermethylated in kidney and cerebrum (87). This same study revealed differential histone H3 acetylation between liver and these non-ABCG5/G8-expressing tissues, suggesting that epigenetic mechanisms may control tissue-specific expression of the complex. A highly intriguing mechanism by which p53 promotes gene expression is by recognition of a structural-specific motif called triplex DNA (88). The ABCG5 proximal promoter contains a predicted T·A·T triplex. Expression of ABCG5 mRNA increased following transient transfection of p53 in cultured cells, as well as treatment with p53-stabilizing agents in p53-positive cells, but not in p53-negative cells (88). However, expression of ABCG8 was not examined, the cultured cells were not of hepatic, intestinal, or gallbladder epithelial origin, and the extent to which this mechanism influences ABCG5/G8 expression or sterol metabolism is not known. Interestingly, the insulin receptor was also identified as a T·A·T triplex-responsive gene. Whether this is coincidental or mechanistically links alterations in insulin signaling and ABCG5/G8 expression is unclear.

### Regulation of complex formation and trafficking

ABCG5 and ABCG8 are required to form a functional sterol transport complex. This marriage is formed as the complex folds in the ER and is dependent on the presence of N-linked glycans and the calnexin/calreticulin chaperone system (89). Using a series of chimeras comprised of the N- and C-terminal domains of ABCG1, ABCG2, ABCG5, and ABCG8, evidence for an ER-retrieval signal within the N-terminal cytosolic domains of ABCG5 and ABCG8 that prevents the monomers from exiting the ER prematurely was identified (90). The heterodimer complex formation has now been supported by a high-resolution crystal structure (91) that will allow for future structure-function studies to be conducted (Fig. 2), graciously provided by Dr. Jyh-Yeung Lee, University of Ottawa). Although the N-terminal structures of ABCG5 and ABCG8 could not be established within the crystal structure (91), many of the nonsynonymous polymorphic changes, as well as the missense mutations can be mapped onto a structure (Fig. 2). The missense mutations (R419 and R550 in ABCG5 and L572, G574, L596, R573, L501, and L405 in ABCG8) are very close to the putative sterol-binding pocket [see also the putative sterol-binding pocket for a model of ABCG4 (92) based upon the ABCG5/ABCG8 crystal structure]. Interestingly, the polymorphic residues, I523 in ABCG5 and G575 in ABCG8, would also be close to this site and form good future candidates to explore their effect on sterol-specificity of binding. In polarized cultured hepatocytes, recombinant ABCG5/G8 was shown to reside at the canalicular membrane (93). Within the small intestine, endogenous mouse, as well as a human, ABCG8 transgenes were localized almost exclusively to the apical surface (19). However, biochemical fractionation and immunolocalization approaches suggest a broader intracellular distribution of ABCG5 and ABCG8 (94). More recently, a recombinant fluorescently tagged ABCG5/G8 complex was shown to reside in an intracellular pool that is recruited to the canalicular surface in response to a lithogenic diet (95). In isolated rat liver canalicular membranes, ABCG5 was shown to reside in Triton-soluble Lubrol-resistant microdomains.
Cholesterol and xenosterol trafficking

Similar results were obtained using bile acids as detergents in which ABCG5, ABCB4, and ABCB11 are co-localized, suggesting the complete biliary lipid secretion apparatus resides within microdomains at the canalicular surface (97).

Regulation of ABCG5/G8 activity

The mechanism by which ABCG5/G8 promotes (chole)sterol efflux is not fully understood. It was originally suggested that ABCG5/G8 function as a floppase, promoting the movement of cholesterol to the exofacial leaflet of the canalicular membrane based on the observation that there was less cholesterol in isolated canalicular membranes of ABCG8-deficient mice (98). In cell-based models, coexpression of ABCG5 and ABCG8 will promote cholesterol efflux to bile-acid micelles, but not other traditional acceptors, such as ApoAI, HDL, or cyclodextrins, indicating that the molecular mechanism by which ABCG5/G8 promotes sterol efflux at the apical membrane is distinct from other cholesterol efflux pumps, such as ABCA1 and ABCG1 (99, 100). Using purified native mouse ABCG5/G8, sterol transport between donor and acceptor liposomes suggests a 1:1 stoichiometry for ATP hydrolysis and cholesterol transport, and bile acids could stimulate this ATP hydrolysis (101). Among the tested bile acids, cholate was the most potent at stimulating ATPase activity, whereas its taurine or glycine conjugates were approximately 50% less effective. Using various lipid extracts and defined preparations, the presence of cholesterol stimulated ATPase activity to varying degrees. Thus, the presence of both bile acids and cholesterol appear to act cooperatively to activate the pump and promote cholesterol secretion. This is an attractive model for the coupling of bile acid and cholesterol secretion and supported by the publication of the ABCG5 and ABCG8 crystal structures that show a sterol-binding pocket, which may undergo structural changes upon ATP binding to make this more available at the luminal surface for phospholipid-bile acid acceptors to allow for excretion (91). What remains to be established is whether this mechanism is constantly active or is regulated by translocation of these heterodimers from an intracellular location to the apical surface. The crystal structure should also allow for exploration of how missense mutations disrupt function, as well as how the nonsynonymous polymorphic variants seen in humans may alter function.

Under a number of conditions, cholesterol and bile secretion can be uncoupled, resulting in reduced cholesterol excretion or the supersaturation of bile (102). The extent to which these conditions are due to alterations in ABCG5/G8 activity awaits further investigation.

MOUSE MODELS USED TO ELUCIDATE THE IMPACT OF STEROLINS IN PHYSIOLOGY AND DISEASE

The genes encoding ABCG5 and ABCG8 have been disrupted, as well as overexpressed, in mice in order to perform detailed physiological studies of sterolin function. Mice lacking functional sterolins in both liver and intestine have been created by standard gene targeting of Abcg5 (21), Abcg8 (20), or both Abcg5 and Abcg8 (17). Recently, mice were created with liver- and intestine-specific knockouts of Abcg5 and Abcg8 (57). The integration of the human ABCG5 and ABCG8 genes into the genome of mice resulted in mouse models with increased sterolin function in both liver and intestine (18) or liver only (103). In addition, adenoviral vectors have been employed in mice to increase or restore hepatic ABCG5 and ABCG8 function (19, 104, 105). These various mouse models have been employed to study the impact of ABCG5/G8 on sterol absorption and secretion, sitosterolemia, reverse cholesterol transport (RCT), TICE, metabolic disease, and atherosclerosis.

Xenosterol exclusion from the body

Like patients with sitosterolemia, mice lacking ABCG5/G8 function are unable to preferentially excrete xenosterols into bile or the intestinal lumen. When fed a diet that contains phytosterols, mice with homozygous disruption of Abcg5, Abcg8, or both Abcg5 and Abcg8 consistently developed sitosterolemia (12, 17, 20, 21, 46, 49, 72, 106). Plasma and tissue phytosterol levels in mice with intestinal- or liver-specific knockout of Abcg5 and Abcg8 were significantly less than those of mice with whole-body ABCG5/G8 deficiency.
The rescue of intestinal ABCG5/G8 function in Abcg8<sup>−/−</sup> mice also significantly reduced, but did not normalize, plasma phytosterol concentrations (12). It can be concluded that intestinal ABCG5/G8 excretes the majority of xenosterols back into the lumen, acting as a first-pass gate. Hepatic ABCG5/G8 then pumps into the bile those xenosterols that had evaded the intestinal checkpoint. In the absence of ABCG5/G8 function, xenosterols can accumulate at high levels in mice causing several pathological conditions. Infertility was observed in male and female Abcg5<sup>−/−</sup> and Abcg8<sup>−/−</sup> mice consuming a phytosterol-rich chow diet, but when phytosterol accumulation in the body was blocked with a meat-based (xenosterol free) diet, or the sterol absorption inhibitor, ezetimibe, fertility was restored (11, 12, 107). Several studies have shown that mice lacking functional ABCG5/G8 developed macrothrombocytopenia, a disorder characterized by significantly reduced platelet levels and increased platelet size. Abcg5<sup>−/−</sup>, Abcg8<sup>−/−</sup>, and Abcg5<sup>−/−</sup>Abcg8<sup>−/−</sup> mice also displayed hemolytic anemia, prolonged bleeding times, and decreased platelet activation and aggregation (11, 49, 50). The macrothrombocytopenia and abnormal bleeding was the result of phytosterol accumulation in platelet membranes resulting in dysfunctional platelets (50). Mice deficient in ABCG5 and/or ABCG8 also developed cardiomyopathy characterized by phytosterol accumulation, histiocytic infiltration, multifocal fibrosis, and tissue calcification (11, 13, 49, 50). Interestingly, the development of cardiomyopathy is dependent both on the presence of high levels of xenosterols, as well as intact B and T cell function (13).

**Cholesterol absorption**

Action of ABCG5/G8 in the enterocytes is believed to reduce cholesterol absorption. Fractional cholesterol absorption (FCA) decreased and fecal neutral sterol excretion (FNSE) increased in transgenic mice with both intestine and liver (INTLIV-Tg) overexpression of human ABCG5/G8 (18, 108, 109). In contrast, FCA and FNSE were similar for wild-type controls and transgenic mice with liver-specific overexpression of human ABCG5/G8 (103). Because both transgenic lines had increased biliary cholesterol secretion, the difference in FCA and FNSE between the models could have been due to the human and mouse ABCG5/G8 pumping greater amounts of biliary and dietary cholesterol out of the enterocytes and back into the intestinal lumen of the INTLIV-Tg mice. However, when considering that the INTLIV-Tg mice versus the wild-type controls had 6- to 7-fold more cholesterol in bile, the absolute amount of cholesterol absorbed by the INTLIV-Tg mice versus the wild-type controls would have been significantly greater even with the 50% reduction in FCA. It is also possible that the increased FNSE was due to the bile of the INTLIV-Tg mice becoming supersaturated with cholesterol, thus reducing luminal cholesterol bioavailability to the enterocytes.

If overexpression of ABCG5/G8 decreases FCA and increases FNSE, then it would be anticipated that the absence of ABCG5/G8 would have the opposite effects. FNSE was reduced, but FCA, when measured using the fecal dual isotope or plasma dual isotope methods, was unchanged in chow-fed Abcg5<sup>−/−</sup>, Abcg8<sup>−/−</sup>, and Abcg5<sup>−/−</sup>Abcg8<sup>−/−</sup> mice (17, 21, 72, 110, 111). Cholesterol absorption, as assessed by the movement of luminal-infused cholesterol into lymph, was found to be decreased in two studies using Abcg5<sup>−/−</sup>Abcg8<sup>−/−</sup> mice and increased in another study using Abcg8<sup>−/−</sup> mice (112–114). Compared with wild-type controls, mice with intestinal-specific ABCG5/G8 deficiency had no changes in FCA or FNSE (57). These studies indicate that the absence of intestinal ABCG5/G8 does not lead to increased cholesterol absorption and, consequently, the reduced FNSE observed in whole-body ABCG5/G8-deficient mice could be driven by dramatically decreased biliary cholesterol secretion. Because most of the above studies were conducted in mice fed a chow diet (low in fat and cholesterol, but high in phytosterols), it will be important to determine the impact of liver and/or intestinal ABCG5/G8 deficiency on cholesterol absorption in mice consuming a Western-type diet (WTD) with a higher fat and cholesterol content. It will also be essential to assess whether cholesterol absorption in these models is affected by the inclusion in the diet of animal- or plant-derived fats that contain cholesterol and phytosterols, respectively.

**Biliary cholesterol secretion**

ABCG5/G8 are critical to the secretion of cholesterol into the bile. Abcg5<sup>−/−</sup>, Abcg8<sup>−/−</sup>, whole-body, and liver-specific Abcg5<sup>−/−</sup>Abcg8<sup>−/−</sup> mice fed a chow diet displayed a 70–90% reduction in biliary cholesterol concentration or secretion (17, 20, 21, 57). In contrast, transgenic overexpression of human ABCG5/G8 in liver and intestine increased biliary cholesterol concentrations by 5-fold and 2- to 4-fold in chow-fed male and female mice, respectively (18). Biliary cholesterol concentration and secretion by hepatic overexpression of human ABCG5/G8 was unchanged on a chow diet, but was increased 1.5- to 2-fold on a high-cholesterol diet (103). In the absence of the biliary phospholipid transporter, Abcb4, biliary cholesterol concentration in mice was very low and was not increased by the hepatic expression of a human ABCG5/G8 transgene, indicating that biliary phospholipid secretion is required for ABCG5/G8-mediated cholesterol secretion (108). In addition, studies in which wild-type or Abcg8<sup>−/−</sup> mice were infused with hydrophilic and hydrophobic bile salts indicated that ABCG5/G8 drives biliary cholesterol secretion by flopping cholesterol from the inner to outer leaflet of the canalicular membrane (98). When fed a lithogenic diet, Abcg5<sup>−/−</sup> and Abcg5<sup>−/−</sup>Abcg8<sup>−/−</sup> mice, compared with control C57BL/6J mice, had decreased incidence of gallstone development (115).

**Macrophage RCT**

Macrophage RCT (mRCT) is an anti-atherogenic pathway through which excess cholesterol is effluxed from plaque macrophages, transported on lipoproteins to the liver, secreted into bile, and excreted from the body. The primary method used to assess mRCT quantifies the movement of <sup>3</sup>H-cholesterol from intraperitoneally injected macrophage foam cells to feces over 48 h (116). C57BL/6J
mice with adenoviral-mediated overexpression of murine ABCG5/G8 (AdABC5/G8) displayed increased fecal macrophase-derived \(^3\)H-neutral sterol (i.e., cholesterol), consistent with an increase in biliary cholesterol secretion. However, because the sum of \(^3\)H-neutral sterol and \(^3\)H-bile acid in feces was similar for AdABC5/G8 and AdNull controls, it was concluded that increased hepatic ABCG5/G8 function alone does not impact macrophase-to-feces RCT. Studies in sterolin-deficient mice found that, in spite of significant reductions in biliary cholesterol secretion, mice lacking ABCG5/G8 function had no deficits in mRCT (105, 117, 118). However, treatment with LXR agonist increased \(^3\)H-neutral sterol excretion from wild-type mice, but not Abcg5\(^{-/-}\),Abcg8\(^{-/-}\) mice, leading to the conclusion that stimulation of mRCT by LXR activation requires ABCG5/G8 (117).

**TICE**

TICE is the elimination of plasma cholesterol into the intestinal lumen by enterocytes. Several studies have indicated that intestinal ABCG5/G8 play an important role in TICE. Cholesterol fluxes, measured with tracer techniques, showed that the contribution of TICE to fractional neutral sterol excretion (FNSE) was 25% in Abcg5\(^{+/+}\) mice, but 15% in Abcg5\(^{-/-}\) mice (110). Blocking cholesterol absorption with ezetimibe in wild-type mice elevated FNSE by 3.8-fold and TICE was responsible for 74% of this increase (111). Treatment of wild-type mice with the FXR agonist, PX20606, stimulated FNSE and TICE to a level similar to that observed with ezetimibe (119). In the absence of ABCG8, ezetimibe or PX20606 treatment blunted the increase in FNSE and TICE (111, 119). To gauge the importance of intestinal ABCG5/G8 in RCT, mice with intestinal-specific deficiency of ABCG5/G8 were injected with \(^3\)H-cholesterol. FNSE and TICE (111, 119). To gauge the importance of intestinal ABCG5/G8 in RCT, mice with intestinal-specific deficiency of ABCG5/G8 were injected with \(^3\)H-cholesterol in RCT, mice with intestinal-specific deficiency of ABCG5/G8 were injected with \(^3\)H-cholesterol and the levels of \(^3\)H-cholesterol in bile and feces were measured. In spite of similar biliary cholesterol and FCA, the intestinal-specific ABCG5/G8 knockouts compared with wild-type control mice had decreased fecal \(^3\)H-cholesterol excretion, likely due to a reduction in TICE (57). In contrast to the above studies, no difference in TICE was observed when measured in wild-type or sterolin mice by intestinal perfusion (120). The overall results indicate that ABCG5/G8 play a major role in TICE.

**Atherosclerosis**

Atherosclerosis development has been assessed in transgenic mice with both intestine and liver (IntLiv-Tg) and liver only (Liv-Tg) expression of human ABCG5/G8. Compared with controls lacking only the LDL receptor (Ldlr\(^{-/-}\)), IntLiv-Tg;Ldlr\(^{-/-}\) mice fed a WTD for 6 months had significantly less aortic atherosclerosis (109). The combination of increased biliary cholesterol secretion and decreased cholesterol absorption in the IntLiv-Tg;Ldlr\(^{-/-}\) mice resulted in a significant reduction in the plasma cholesterol. In contrast, the Liv-Tg on either the LDLR or ApoE knockout background did not have reduced atherosclerosis development (103). Liv-Tg;ApoE\(^{-/-}\) mice fed chow and Liv-Tg;Ldlr\(^{-/-}\) mice fed WTD had significantly increased biliary cholesterol secretion, but compared with controls, no changes in cholesterol absorption or plasma cholesterol concentrations were noted. It was concluded that atherosclerosis was unaltered in IntLiv-Tg mice because the increased biliary cholesterol was being absorbed back into the body. When fed a WTD plus the cholesterol absorption inhibitor, ezetimibe, Liv-Tg;Ldlr\(^{-/-}\) compared to Ldlr\(^{-/-}\) mice displayed significant reductions in total plasma cholesterol and atherosclerosis development (121).

To assess the impact of ABCG5/G8 deficiency and consequently increased phytosterol levels on atherosclerosis development, whole-body ABCG5/G8-deficient mice expressing or lacking LDL receptor were studied (109). Feeding either a chow diet or WTD for 7 months caused a significant increase in plasma phytosterol concentration in Abcg5\(^{-/-}\),Abcg8\(^{-/-}\) and Abcg5\(^{-/-}\),Abcg8\(^{+/+}\);Ldlr\(^{-/-}\) mice compared with wild-type controls. Regardless of the genotype, atherosclerosis did not develop in mice fed chow. Consumption of the WTD by the Abcg5\(^{-/-}\),Abcg8\(^{-/-}\);Ldlr\(^{-/-}\) and Abcg5\(^{+/+}\),Abcg8\(^{+/+}\);Ldlr\(^{-/-}\) mice resulted in similar increases in total plasma sterols and atherosclerosis formation. Based upon these results, it was concluded that there was no association between phytosterol levels and atherosclerosis development in mice. The lack of difference in atherosclerosis formation between the Abcg5\(^{-/-}\),Abcg8\(^{-/-}\);Ldlr\(^{-/-}\) and Abcg5\(^{+/+}\),Abcg8\(^{+/+}\);Ldlr\(^{-/-}\) could have been the result of LXR activation by phytosterols offsetting the decreased biliary cholesterol secretion (122).

**Metabolic disease**

ABCG5/G8 function also appears to limit the development of metabolic disease. Sterolin-deficient mice fed a phytosterol-rich chow diet developed hypertriglyceridemia due to increases in intestinal triglyceride (TG) absorption and hepatic TG secretion and reductions in TG-rich lipoprotein clearance and lipoprotein lipase activity (123). When fed a phytosterol-free high-fat diet, these mice had increased hepatic TG and cholesterol accumulation leading to liver inflammation, loss of glycemic control, and insulin resistance (124). Obese db/db mice have reduced hepatic ABCG5/G8 protein and decreased biliary cholesterol (80). Adenoviral-mediated overexpression of ABCG5/G8 in db/db mice increased biliary cholesterol secretion and restored hepatic insulin sensitivity resulting in reduced fasting glucose and TGs and improved glucose tolerance (104). However, the relationship between biliary and intestinal sterol transport and the underlying mechanisms that contribute to metabolic disturbances remain unclear.

**CONCLUSIONS**

The human diet is very varied and exposes us to more than 60 different xenosterols, from shellfish, mushrooms, yeasts, seaweeds, plants, etc., and we need to have robust mechanisms to prevent toxic xenosterols from accumulation. The sitosterolemia locus is an evolutionarily conserved locus that has allowed organisms with a liver and biliary system to perform this function, from fish to man.
(9) Sitosterol may represent the most abundant of these dietary xenosterols, but this may have led to an error of scientific bias; all attention has focused on this sterol without critical questioning of whether other more potent and toxic sterols (stigmasterol,avenosterol, fucosterol, and methyl-cholesterol to name a few) are major endocrine disruptors, harmful to megakaryocyte biology, or even alter macrophage and immune function (125). In sitosterolemia (and we propose that the term xenosterolemia be used to increase this awareness), feeding a diet rich in shellfish sterols led to accumulation of these exotic sterols within the body (5). Blocking entry of these xenosterols by the drug ezetimibe has been shown to ameliorate this condition (126, 127). Control subjects who have normal ABCG5/ABCG8 function show trace plasma levels of these xenosterols, indicating that we have a daily exposure of these xenosterols to our tissues (5). Daily dietary intakes of these xenosterols are collectively greater than our daily dietary intake of cholesterol. ABCG5/G8 function as cholesterol regulators and in this role, because >99% of the mammalian body has cholesterol as its main sterol, cholesterol (not xenosterol) trafficking remains as a key center piece. However, while the full spectrum of loss of sterolin function in humans may continue to evolve for this rare disease, the mammalian animal models have indicated that several biological steps can be disrupted by these xenosterols and that these processes are dependent on the continued presence of these xenosterols in their diets. We believe that it is time to contemplate mechanistic paths away from the cholesterol-centric view of pathophysiology, especially where diets are a key component, and consider the role of xenosterols. If so, our scientific rigor in measuring and reporting the sterol spectrum may be a first step in defining the problem. 

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