Polyphenol-induced improvements in glucose metabolism are associated with bile acid signaling to intestinal farnesoid X receptor

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ABSTRACT

Introduction Bile acid (BA) biotransformation by gut bacteria impacts BA profile and signaling to nuclear receptors, such as the farnesoid X receptor (FXR) regulating glucose metabolism. Altered BA-FXR signaling was therefore investigated as a potential mechanism linking polyphenol-induced gut bacterial changes and improved glucose metabolism.

Research design and methods Diabetic db/db were fed low-fat diet (LFD) or LFD supplemented with a proanthocyanidin-rich extract of grape polyphenols (LFD-GP) for 4 weeks. Metabolic phenotypes, serum BAs, gut microbiota composition, and gene expression markers relevant to gut barrier and glucose metabolism were assessed. Gut organoids were used to investigate effects of individual BAs on ileal FXR activity.

Results Compared with LFD-fed controls, GP supplemented db/db mice showed improved glucose metabolism, decreased relative abundance of gut bacteria associated with production of secondary BAs (SBAs), and depleted serum levels of SBAs taurohyodeoxycholic acid (THDCA), ω-muricholic acid (ωMCA), and tauro-ω-muricholic acid (TωMCA). Serum levels of primary BAs (PBAs) increased, consistent with higher gene expression of PBA synthesis enzyme Cyp7a1. GP-induced BA changes associated with FXR inhibition as evidenced by reduced expression of FXR-responsive genes Shp, Fgf15, and Fabp6 in ileum tissue as well as hepatic Shp, which negatively regulates PBA synthesis. GP treatment did not affect expression of hepatic Fxr or expression of Abcb11, Slc51b, and Obp2a genes controlling BA transport. Ceramide biosynthesis genes Smpd3, Splic2, and Cers4 were decreased in liver and intestine suggesting lower tissue ceramides levels may contribute to improved glucose metabolism. THDCA, ωMCA, and TωMCA behaved as FXR agonists in ileal organoid experiments; therefore, their depletion in serum of GP-supplemented db/db and wild type (WT) mice was consistent with FXR inhibition.

Conclusion These data suggest that altering the gut microbiota, GPs modify BA-FXR signaling pathways to promote glucoregulation.

INTRODUCTION

Dietary polyphenols in plant-based foods contribute to improved glycemic control in mice and humans.1–3 Improved glucose metabolism in mice supplemented with berry/fruit extracts was related to a proanthocyanidin (PAC)-induced bloom of Akkermansia muciniphila,1,2,4 a microbe shown to attenuate symptoms of metabolic syndrome (MetS) and type-2 diabetes (T2D) in obese mice and humans.5,6 We hypothesized that metabolic improvements also result from changes in mouse and human gut microbiota.7,8 Improved glucose metabolism in mice supplemented with berry/fruit extracts was related to a proanthocyanidin (PAC)-induced bloom of Akkermansia muciniphila,1,2,4 a microbe shown to attenuate symptoms of metabolic syndrome (MetS) and type-2 diabetes (T2D) in obese mice and humans.5,6 We hypothesized that metabolic improvements also result from changes in gut microbiota.7,8

Significance of this study

What is already known about this subject?

► Dietary polyphenols, such as proanthocyanidins, alter the gut microbial community and are associated with improved metabolic resilience in humans and mice.

► Bile acids (BAs) signal to farnesoid X receptor (FXR), a nuclear transcription factor that regulates hepatic BA synthesis and glucose metabolism.

What are the new findings?

► Grape polyphenol (GP) supplementation decreased abundances of gut bacterial taxa associated with secondary BA (SBA) production concomitant with depletion of SBAs and increased primary BAs (PBAs) in murine serum.

► GP supplementation suppressed expression of FXR-regulated genes Fgf15, Fabp6, and Shp, an inhibitor of PBA synthesis, which was consistent with increased serum PBAs.

► GP-induced FXR inhibition was associated with decreased expression of genes required for biosynthesis of ceramides, which impair glucose homeostasis.

► The SBAs depleted in GP-treated mice were revealed as FXR agonists in ileal gut organoids.

How might these results change the focus of research or clinical practice?

► This study highlights BA-FXR signaling pathways as an important mechanism for further investigation in human intervention studies of dietary polyphenols and metabolic health.
host-derived bacterial metabolites regulating host energy metabolism.

Bile acids (BAs) are signaling molecules linking the gut microbiota to host energy metabolism. Primary BAs (PBAs) synthesized in the liver are conjugated with taurine (mice) or glycine (humans).7 Bacterial-derived bile salt hydroxylases deconjugate taurine or glycine from the sterol core of PBAs followed by bacterial transformations, such as 7α/β-dehydroxylation, dehydrogenation, and epimerization, which generate secondary BAs (SBAs).7 PBAs and SBAs signal to key regulators of energy metabolism, such as nuclear transcription factor farnesoid X receptor (FXR) and Takeda G protein-coupled receptor 5 (TGR5).7 Fxr−/− mice were protected from high-fat diet (HFD)-induced obesity and had altered BA and gut microbiota (higher Bacteroidetes, less Firmicutes) profiles compared with wild type (WT) mice.8 Transfer of cecal microbiota from HFD-fed Fxr−/− mice to germ free WT mice resulted in less adiposity and improved glucose metabolism compared with mice that received microbiota from HFD-fed WT mice, suggesting the altered gut microbial and BA profiles in FXR deficient mice contributed to metabolic improvements.8 Studies using tissue-specific FXR knockout mice showed that intestinal FXR activity was required to mediate HFD-induced metabolic dysfunctions.9 10 Pharmacological inhibition of intestinal FXR using tempol, antibiotics, metformin, or glycine-β-muricholic acid (Gly-MCA) led to gut microbial remodeling and improved glucose and lipid homeostasis.9 10 12

In addition to the A. muciniphila bloom, mice fed HFD-supplemented with PAC-rich grape polyphenols (GPs) showed other profound bacterial community changes raising the possibility that altered BA profile and signaling could contribute to observed improvement in glucose homeostasis.1 4 The present study provides compelling evidence in support of this hypothesis.

**RESEARCH DESIGN AND METHODS**

**Diets**

A complex of 10% grape polyphenols and soy protein isolate (GP-SPI) was prepared as previously described.14 16 Nutritional profiles for SPI and GP-SPI are provided in online supplementary table 4. Isocaloric, ingredient-matched diets (Research Diets, New Brunswick, New Jersey, USA) used in this study were previously described.4 Mice were fed low-fat diet (LFD) containing 10% SPI, LFD formulated with 10% GP-SPI delivering 1% GP (LFD-GP), HFD containing 10% SPI or HFD formulated with 10% GP-SPI delivering 1% GP (HFD-GP). Online supplementary table 2 provides diet formulation details.

**Animals**

Four-week-old db/db mice (B6.BKS(D)-Leprb/J; stock no. 000697; Jackson Laboratory, Bar Harbor, Maine, USA) were single-housed on a 12-hour light/dark cycle (07:00 to 19:00 hours light) with ad libitum access to LFD and water in a controlled-temperature room (24°C±1°C) for 1 week for acclimation. Mice were randomized to receive LFD (n=7) or LFD-GP (n=7) for 28 days. Metabolic phenotyping included food intake, body weight, body composition (EchoMRI 3-1 system; Echo Medical Systems, Houston, Texas, USA), and oral glucose tolerance tests, as previously described.14 On day 29, mice were euthanized by CO₂ inhalation, followed by cardiac puncture and collection of tissues, as previously described.14 Fecal and cecal samples were collected for microbial community profiling. Wild type C57BL/6J mice (n=10) aged 5 weeks were acclimated on LFD for 1 week and randomly divided into two groups and fed HFD or HFD-GP (n=5 per group) for 10 weeks after which mice were euthanized by CO₂ inhalation.

**Tissue gene expression analysis**

RNA was extracted from ileum, jejunum, colon, and liver (30–60 mg) with RNeasy Plus Universal Mini Kit (QIAGEN) followed by RNA cleanup (Machery-Nagel, RNA purification kit). RNA (5 μg) was reverse transcribed to cDNA and qPCR was performed using TaqMan primers (online supplementary table 4) as previously described.4 Data were analyzed using 2⁻ΔΔCT method using hydroxymethylbilane synthase (HMBS) as housekeeping gene.

qPCR of A. muciniphila in fecal and cecal samples was performed as previously described.4 Briefly, gDNA extracted from fecal/cecal samples was diluted to 2.5 ng/μL for quantification of A. muciniphila abundance relative to total bacteria and archaea by qPCR using A. muciniphila (AM1, AM2) and universal primer (U341F, U515R) sets.14

**16S rRNA gene sequencing**

Genomic DNA was extracted from 84 fecal and 14 cecal samples collected from db/db mice. Illumina protocols were used to prepare V4 amplicons of the 16S rRNA gene for sequencing on a MiSeq system, resulting in >1.099×10⁷ reads. Denoising and clustering were conducted using DADA2 algorithm to differentiate sequences into amplicon sequence variants (ASVs) for downstream analysis using QIME 2.0.14 Details are available in online supplementary materials.

**Serum biochemistry**

Serum for BA analysis was collected by cardiac puncture. BAs were analyzed on a Waters’s Alliance e2695 HPLC system (Waters Milford, Massachusetts, USA) coupled to a Waters’s Acquity QDA mass spectrometer equipped with an electrospray interphase (ESI; Waters Milford, Massachusetts, USA) and quantified by external calibration curves using >95% pure standards. Details are provided in online supplementary materials. Serum leptin, polypeptide YY (PYY), interleukin-6 (IL-6) and insulin were determined using a MILLIPLEX MAP Mouse Metabolic Hormone Magnetic Bead kit (Millipore) with a MagPix instrument (Luminex), as previously described.4

**Organoid experiments**

Intestinal crypts were isolated from WT C57BL/6J mouse ileum, according to established methods.16 Crypts were counted and added to Matrigel (five crypts per μL).
(Corning 356231, growth factor reduced) and 25 µL was added per well (in 48-well plates) and allowed to polymerize (37°C for 15–20 min), followed by addition of 1 mL of complete growth medium (CGM, see online supplementary materials). CGM (without Y-27632 dihydrochloride monohydrate) was replaced every 2–3 days. Organoids were passed every 5–7 days (1:3 ratio). Mature organoids (4 days postpassage) were treated in triplicate (n=3 wells) with 5% methanol (vehicle), chenodeoxycholic acid (CDCA) (100 µM) alone, or CDCA plus another BA (200 µM) in CGM for 17 hours. RNA was extracted for qPCR analysis using TaqMan primers, as described above.4

Statistical analyses
Analyses were conducted using Prism 8.0.2 (GraphPad Software, La Jolla, California, USA). Significant differences (two groups) were assessed with a two-tailed, unpaired Student’s t test with Welch correction for unequal variance when needed or by one-way or two-way analysis of variance (>2 groups) followed by Sidak’s or Tukey’s multiple comparison test. Statistical analysis of alpha and beta-diversity metrics was calculated using QIIME 2 (details in online supplementary materials). ADONIS and permutation analysis were conducted using R Studio V.3.4.2 (R Studio Software, Boston, Massachusetts, USA) and Python 3.7.1.

RESULTS
GP s improve glucose metabolism in db/db mice independent of obesity
Leptin receptor-deficient db/db mice develop obesity, gut barrier dysfunction, and hyperglycemia independent of HFD feeding.17 Compared with LFD-fed controls, db/db mice fed LFD containing PAC-rich GPs (LFD-GP) for 4 weeks showed significantly improved oral glucose tolerance (figure 1A).4 Area under the curve remained stable over time for the LFD-GP group but increased in the LFD group (figure 1A). Mice fed LFD-GP initially exhibited a transient decrease in food intake presumably due to taste, but at later time points LFD and LFD-GP groups consumed similar amounts of food (4.99±0.69 and 4.29±0.69 g/day/mouse, respectively, p>0.05, online supplementary figure 1A). The LFD-GP group consumed 42.9±6.9 mg of GPs per day. Both groups had similar body weight gain, body composition, and liver weights (online supplementary figure 1B–D).

GPs promote a bloom in A. muciniphila without improving markers of metabolic endotoxemia
Compared with controls, GP-supplemented db/db mice had decreased α-diversity as evidenced by richness, Shannon index, and Faith’s phylogenetic diversity index (figure 2A). Principal coordinate analysis showed that GPs significantly altered fecal community structure within 2 days (online supplementary figure 2). As previously observed,4 4 GPs promoted increased cecal mass (online supplementary figure 1E), a phenotype common to antibiotic-treated and germ-free mice and consistent with reported antibacterial properties of PACs.18 19 GP-supplemented mice had an increased relative abundance of phylum Verrucomicrobia at the expense of Firmicutes in fecal (days 6–27) and ceca samples (day 27) (figure 2B,C, online supplementary figure 3A,B). Ceca of GP-supplemented mice had higher relative abundance of Bacteroidetes (25%>LFD) although, except for day 13, fecal Bacteroidetes remained similar between groups (figure 2B, online supplementary figure 3C). GP supplementation did not consistently alter levels of Proteobacteria or Actinobacteria (figure 2B, online supplementary figure 3D,E). Consistent with increased Verrucomicrobia, quantitative qPCR analyses confirmed that GPs promoted a bloom in Akkermansia muciniphila (figure 2C) at the expense of other taxa (figure 2D).

Reduced abundance of A. muciniphila and metabolic endotoxemia (characterized by gut dysbiosis, compromised gut barrier integrity, lipopolysaccharide (LPS) leakage, and intestinal inflammation) was associated with impaired glucose metabolism in obese mice and humans.5 6 20 Oral administration of A. muciniphila in obese, diabetic mice and humans resulted in improved glucose homeostasis and attenuated metabolic endotoxemia through improved gut barrier integrity.5 6 20 GP-induced improvement in glucose metabolism in HFD-fed mice has therefore been considered a consequence of the A. muciniphila bloom leading to reduced inflammation and increased gut barrier integrity.1 2 4 Reduced metabolic endotoxemia could not, however, explain the improved glucose tolerance in GP-supplemented db/db mice despite increased A. muciniphila (~15%) in feces (day 27) and cecum (figure 2C). Relative to control, GP-supplementation did not change intestinal gene expression of markers of inflammation (Tnf, Il-6, iNos), gut barrier integrity (Tjp1, Ocln, Muc2), peripheral lipid deposition (Fiaf) or glucose transport (Glut2) (online supplementary figure 4A–C). GP-supplemented db/db mice had less Muc3 expression in jejunum and ileum (online supplementary figure 4A,B), suggesting lower mucus secretion. There were no differences in serum insulin, Il-6, or glucoregulatory hormones PYY and leptin (online supplementary table 3). These data suggested that other mechanism(s) were driving improved glucose metabolism.

GPs reduced gut bacterial taxa associated with production of SBAs
Although total bacterial and archaeal abundance was not significantly different between LFD and LFD-GP groups (figure 2E), GPs induced profound gut microbial changes (online supplementary figure 5) that would be expected to alter BA diversity, abundance, and signaling. Targeted liquid chromatography-mass spectrometry (LC-MS) analysis revealed that GP-supplemented db/db mice had higher serum concentrations of
PBAs driven by increased cholic acid (CA) and taurocholic acid (TCA) (figure 3A and C, online supplementary figure 6A). Concentrations of β-muricholic acid (βMCA), tauro-βMCA (TβMCA), tauro-αMCA (TαMCA), taurochenodeoxycholic acid (TCDDA) were similar between groups; αMCA was not detected in LFD-GP group (figure 3A, online supplementary figure 6A). Increased PBA pool correlated with an overall reduction in SBAs (figure 3C) where ωMCA, TωMCA, and taurohydroxycholic acid (THDCA) were undetectable in db/db mice fed LFD-GP, although deoxycholic acid (DCA) was increased (figure 3A). Total serum BAs were similar between groups (figure 3B).

The GP-induced depletion of SBAs was unrelated to the leptin receptor mutation in db/db mice. Compared with HFD-fed controls, wild type C57BL6/J mice fed HFD supplemented with 1% GP (HFD-GP) for 10 weeks also showed serum depletion of SBAs THDCA, ωMCA, and TωMCA, reduced overall levels of serum SBAs, and no difference in concentration of total serum BAs (online supplementary figures 6B and 7). Unlike db/db mice, GP-supplemented WT mice had decreased taurodeoxycholic acid (TDCA) and no significant difference in CA or overall PBA pool, although TCA concentration was increased (online supplementary figures 6B and 7).

In db/db mice, multiple correlation analyses were performed to associate GP-induced changes in serum BAs (figure 3) to changes in fecal/cecal gut bacteria (online supplementary figure 5). The GP-induced increase of CA, TCA (PBAs), and DCA (SBA) was positively and significantly associated with increased abundance of Akkermansia, Blautia, Clostridium (ASV-59), and S24-7 (figure 3E,F). Blautia and Clostridium possess...
7α-dehydroxylating activity; therefore, taxa within these genera may be responsible for DCA production via dehydroxylation of CA.21 GP-treated mice had lower levels of taxa within the *Clostridiales* order (ASV-50 and 51), *Ruminococcaceae* and *Lachnospiraceae* families (ie, ASV-56, 57, 61, 65, 70, 75, and 76), and *Clostridium* genus (ASV-74) (online supplementary figure 5), which are reported to possess 7α-dehydroxylating activity required for conversion of PBAs to SBAs.21–24 In agreement with evidence from mice and humans, the GP-induced decrease in these bacterial taxa encoding 7α-dehydroxylation activity was highly correlated to the GP-associated depletion of SBAs ωMCA and TωMCA (figure 3E,F).21 25 Finally, positive and significant associations were found between reduced αMCA, TωMCA, and/or ωMCA and reductions in taxa belonging to *RF39, Anaeroplasma, Ruminococcus, Butyricicoccus, Dorea, Dehalobacterium, Christensenellaceae, Lactococcus, Streptococcus*, and *Oscillospira* (figure 3E,F).

**GP-associated BA changes promote inhibition of FXR signaling and upregulation of classical BA synthesis pathway**

To investigate the consequences of GP-induced serum BA changes, gene expression of FXR, TGR5 and their downstream targets were analyzed in tissues. GP supplementation did not change ileal *Fxr* gene expression; however, FXR transcriptional activity was decreased as expression of its target genes, fibroblast growth factor 15 (*Fgf15*), small heterodimer partner (*Shp*), and ileal BA binding protein (I-BABP, gene *Fabp6*) were suppressed in ileum (figure 4A). Intestinal FXR signaling negatively regulates hepatic PBA synthesis through interaction of Fgf15 with hepatic fibrolast growth factor receptor 4/βKlotho receptor complex or by regulating...
Figure 3  GPs increased PBAs and reduced SBAs in serum in association with depletion of bacterial ASVs related to SBA production. (A) LC-MS and pure standards were used to determine the mean concentration of individual PBAs and SBAs (mean±SD) in serum samples (n=7 samples) collected from individual mice fed LFD (white bars) or LFD-GP (purple bars). (B) Total serum BA concentration (mean±SD) was determined based on sum of individual PBA and SBA concentrations (shown in panel A) quantified for each mouse fed LFD (n=7) or LFD-GP (n=7). (C) Serum PBA and SBA concentrations (mean±SD) in LFD versus LFD-GP diet groups were calculated by summing the individual PBAs or SBAs shown in panel A. (D) Using data from panels B and C, pie charts illustrate pooled PBAs (green) and pooled SBA (gray) as a percentage of total serum BA concentration quantified for LFD and LFD-GP groups. For panels A–C, significant difference was determined using unpaired, two-tailed t-test followed by Welch’s correction: *p<0.05, **p<0.01. Heatmap representation of the Spearman’s r correlation coefficient between serum BA profile and significantly changed bacterial ASVs (at genera or family level of taxonomy) in GP-treated mice (relative to control diet group) from (E) day-27 fecal samples or (F) day-29 cecal samples. Shades of red indicate serum BA and bacterial taxa are positively correlated (0 to +1) while shades of blue indicate a negative correlation (0 to −1). Significant positive or negative correlations are shown: *p<0.05, **p<0.01, ***p<0.001. ASV, amplicon sequence variant; CA, cholic acid; DCA, deoxycholic acid; GP, grape polyphenol; LC-MS, liquid chromatography-mass spectrometry; LFD, low-fat diet; MCA, muricholic acid; PBA, primary bile acid; SBA, secondary bile acid; TCA, taurocholic acid; TCDCA, taurochenodeoxycholic acid; TDCA, tauro-deoxycholic acid; THDCA, taurohydroxydeoxycholic acid; TmMCA, tauro-α-MCA; TβMCA, tauro-β-MCA; TωMCA, tauro-ω-MCA; TUDCA, tauro-ursodeoxycholic acid.
the hepatic expression and gene-repressive function of Shp via liver receptor homologue-1 and hepatocyte nuclear factor-4α. 26-29

In agreement with reduced ileal Fgf15, hepatic Shp expression was significantly decreased (figure 4B). Fgf15 and Shp negatively regulate BA synthesis; therefore, their reduced expression was consistent with hepatic upregulation of cytochrome P450 family 7 subfamily A member 1 (Cyp7a1), the rate-limiting enzyme in the classical BA synthesis pathway, and a trending increase in downstream enzyme Cyp8b1 (p=0.06; figure 4B).

Figure 4  Expression of genes involved in FXR signaling, ceramide synthesis, and glucose metabolism in response to GP-supplementation. Scatter plot of relative mRNA levels of indicated genes expressed in (A) ileum and (B) liver tissues collected from individual mice fed LFD (open squares) or LFD-GP (closed squares). Group mean±SD (n=7 samples/group) is illustrated by horizontal and vertical lines. Data represent qPCR of technical duplicates analyzed by 2−ΔΔCT method. Between group difference was determined by unpaired, two-tailed t-test with or without Welch’s correction for unequal variance: *p<0.05, **p<0.01, ***p<0.001. FXR, farnesoid X receptor; GP, grape polyphenol; LFD, low-fat diet.
supplementary figure 4E), involved in the alternative BA synthesis pathway, were unaffected by GP treatment. To determine if reduced hepatic Shp expression altered hepatic FXR activity, we examined FXR target genes *Acb* encoding bile salt export pump (*Bsep*), *Slc51b* encoding organic solute transporter-β (*Ostβ*), and *Obp2a* encoding lipocalin 13 (*Lpn13*), an acute phase protein. Compared with control, GP treatment did not affect hepatic expression of *Fxr*, *Acb*11, *Slc51b*, or *Obp2a* (figure 4B). These data suggest that GP-mediated effects on FXR activity are restricted to intestine resulting in suppression of ileal *Fgf15* and *Shp* transcription and increased Cyp7a1 activity and PBA synthesis (figure 3).

FXR activity regulates *Tgr5* expression; therefore, we measured *Tgr5* mRNA levels and downstream targets. On activation by BAs, TGR5 signals the release of incretin glucagon-like peptide-1 (Glp-1) to promote euglycemia. Glp-1 is produced when proglucagon protein (*Gcg*) is cleaved by prohormone convertase 1/3 (PC1/3), encoded by *Pcsk1*. GP-supplemented mice showed reduced ileal expression of *Tgr5* and no changes in *Gcg* or *Pcsk1* (figure 4A). GPAs did not change expression of *Fxr*, *Fgf15*, or *Tgr5* in colon tissue. GPs induced an increase in colonic *Gcg* however, *Pcsk1* was unchanged indicating Glp-1 levels were unaffected (online supplementary figure 4D). These data further support the idea that GPs inhibit ileal FXR.

**GP-mediated FXR inhibition is associated with downregulation of ceramide synthesis genes and improved markers of hepatic energy metabolism**

Tissue accumulation of ceramides is linked to insulin resistance and diabetes, which can be ameliorated by pharmacological or genetic inhibition of ceramide biosynthesis. FXR activity positively upregulates genes required for ceramide synthesis in ileum, which leads to impaired glucose metabolism and hepatic steatosis in mouse models of MetS/T2D. Synthesis and accumulation of ceramides in liver contributes to hepatic insulin resistance, steato-hepatisis, and metabolic disease. We found that GP-induced FXR inhibition was associated with lower expression of de novo ceramide synthesis genes, specifically sphingomyelin phosphodiesterase 3 (*Smpd3*) in ileum (figure 4A) and serine palmitoyltransferase long-chain base subunit 2 (*Sptlc2*) and ceramide synthase 4 (*Cers4*) in liver (figure 4B). Consistent with lower expression of ceramide biosynthesis genes, GP-supplemented mice showed improvements in markers of hepatic energy metabolism, evidenced by lower hepatic expression of: (1) carbohydrate response element binding protein (*Chrebp*), a transcription factor that activates key enzymes of de novo lipogenesis; (2) glucose-6-phosphatase (*G6Pase*), which catalyzes the final step in hepatic glucose production; and (3) *Idh3a*, a subunit of the IDH3 isocitrate dehydrogenase heterotetramer complex that regulates fatty acid metabolism and whose inhibition is associated with hepatic glycogen synthesis (figure 4B). GP-supplemented mice had reduced expression of *Lbp*, encoding LPS-binding protein, suggesting decreased liver inflammation and insulin resistance. Hepatic gene expression of gluconeogenesis enzyme phosphoenolpyruvate carboxykinase (*Pck1*) and C/EBP homologous protein (*Chop*), which are normally upregulated during hepatic endoplasmic reticulum stress, were similar between groups (online supplementary figure 4E).

**SBAs depleted in GP-supplemented mice are FXR agonists that promote expression of ceramide synthesis genes**

The majority of secreted BAs (>95%) are reabsorbed in ileum and returned to the liver via the portal vein. Cecum, colon, and feces have similar BA profiles while the serum BA profile is most closely related to that of ileum and portal vein as a minor fraction of reabsorbed BAs enter systemic circulation. We sought to investigate how individual BAs altered by GP supplementation might affect ileal FXR signaling. Gut organoids were cultured from ileal crypts isolated from WT mice (figure 5A) and treated with individual PBAs or SBAs that were differentially detected in serum of GP-supplemented mice (figure 5B). To validate the system, organoids were treated with CDCA, a potent FXR agonist, or CDCA in combination with the FXR antagonist TβMCA. As expected, CDCA increased the expression of *Fxr*, *Fgf15*, and *Shp* compared with vehicle-treated organoids and this effect was attenuated by addition of TβMCA (figure 5B). When organoids were treated with individual BA alone (ie, 100 µM in the absence of CDCA activator), FXR activity was unaffected as determined with CDCA activator), FXR activity was unaffected, as previously reported, when organoids were treated with CDCA, a potent FXR agonist, or CDCA in combination with the FXR antagonist TβMCA. As expected, CDCA increased the expression of *Fxr*, *Fgf15*, and *Shp mRNA expression remained similar to untreated organoids (online supplementary figure 8). We therefore investigated individual BAs in the presence of FXR activator, CDCA, as previously reported, to reveal agonistic or antagonistic effects on FXR signaling. Organoid cultures treated with CDCA in combination with ωMCA, ToMCA, THDCA, or DCA increased CDCA-induced expression of *Fxr*, *Fgf15*, and *Shp*, indicating that these SBAs are FXR agonists (figure 5B). In agreement with FXR agonistic activity, organoids treated with CDCA in combination with ToMCA resulted in increased expression of *Smpd3*, *Cers4*, and Splice2 ceramide synthesis genes. Cotreatment with CDCA and ωMCA, ToMCA, THDCA, or DCA only upregulated *Splice2* (figure 5B). A PBA reported to be a weak FXR agonist and detected at higher concentration in GP-supplemented db/db (but not WT mice), increased CDCA-induced activation of *Fxr*, *Fgf15* and *Shp* and increased expression of *Cers4* and *Splice2*. TCA, a PBA detected at higher concentration in GP-supplemented db/db and WT mice, attenuated CDCA-induced gene expression of *Fxr* and *Fgf15* but not *Shp* and reduced CDCA-induced gene expression of *Cers4* (figure 5B). These organoid data suggest that GPs led to the depletion of FXR activators (ωMCA, ToMCA and THDCA) and increase of an FXR antagonist (TCA). Consistent with in vivo data, the net effect
Figure 5  Ileal organoids treated with BAs revealed agonistic or antagonistic effects on FXR and ceramide pathway genes

(A) Ileal crypts were isolated and cultured in Matrigel medium (detailed in methods). Spheroids (representative day 1 photo) matured into organoids (representative day 4 photo). (B) Scatter plot of relative mRNA levels of indicated genes expressed in organoids after 17 hours of treatment with: vehicle (5% methanol, open black diamonds); 100 µM CDCA (closed red circles); a combination of 100 µM CDCA and known FXR inhibitor TJMCA (200 µM); a combination of 100 µM CDCA and indicated PBA (200 µM, open red circles), a combination of 100 µM CDCA and indicated SBA (200 µM, open blue squares). Data shown were combined from two independent experiments, and for each experiment three wells containing mature organoids were treated with indicated BAs. Data represent qPCR of technical duplicates analyzed by 2−ΔΔCT method. Group mean±SD (n=6 wells total per treatment group) is illustrated by horizontal and vertical lines. Difference compared with CDCA (FXR agonist) was determined by one-way ANOVA followed by Dunnett’s test: *p<0.05, **p<0.01, ***p<0.001. ANOVA, analysis of variance; BA, bile acid; CA, cholic acid; CDCA, chenodeoxycholic acid; DCA, deoxycholic acid; FXR, farnesoid X receptor; MCA, muricholic acid; SBA, secondary bile acid; TCA, taurocholic acid; THCDCA, taurohyodeoxycholic acid; TJMCA, tauro-β-MCA; ToMCA, tauro-ω-MCA; WT, wild type.
was a BA profile that suppressed FXR signaling and de novo ceramide synthesis.

**DISCUSSION**

These data suggest PAC-rich GPs alter a gut microbiota-BA-FXR signaling axis, providing a novel mechanism for glucose regulation. The db/db mouse model is characterized by morbid obesity, gut dysbiosis, intestinal barrier dysfunction, gut inflammation, and hyperglycemia allowing investigation of whether a GP-induced bloom in *A. muciniphila* could ameliorate these metabolic dysfunctions.\(^7\)\(^44\) We showed that GP-induced improvement in glucose metabolism in *db/db* mice was unrelated to an *A. muciniphila*-driven reduction of metabolic endotoxemia. These data are consistent with our previous findings in HFD-fed mice supplemented with PAC-rich GPs for 2 weeks where the *A. muciniphila* bloom and improved glucose metabolism co-occurred, but there were limited positive changes in intestinal gene expression markers associated with metabolic endotoxemia and no significant difference in serum LPS.\(^6\) Similarly, oral administration of the antidiabetic drug metformin increased *A. muciniphila* abundance but metabolic benefits were driven by other bacterial species.\(^11\)\(^45\)\(^46\) Increased *A. muciniphila* has been associated with improvements in markers of metabolic endotoxemia.\(^1\)\(^2\)\(^20\) The absence of this association in the present study may be due to differences in mouse genotypes (WT, *ob/ob*, *db/db*) combined with differences in treatment type (probiotic versus prebiotic) and/or duration of dietary supplementation.

In murine models of HFD-induced obesity and accelerated aging, oral administration of *A. muciniphila* correlated with increased intestinal mucus thickness suggesting improved gut barrier integrity.\(^20\)\(^47\) In the latter study, gene expression of Muc2 and Muc3 were unchanged while Muc1 was downregulated.\(^47\) Rats supplemented with wild blueberry PACs showed increased colonic mucus thickness but no difference in mucin gene expression, which is consistent with mucus being regulated mainly at the post-translational level.\(^48\)\(^49\) In agreement, GP supplementation has consistently shown no increase in Muc2 or Muc3 gene expression; however, mucus thickness was not directly investigated in the present study.\(^14\) *A. muciniphila* has generally been associated with decreased gut inflammation and improved health outcomes. However, increasing *A. muciniphila* abundance is not desirable in all contexts as in other studies it was correlated to mucus depletion, inflammation, and increased susceptibility to enteric pathogen colonization.\(^50\)\(^52\)

Concentration of total quantified serum BAs did not differ between groups, but GP-treated *db/db* mice had increased PBA levels (CA and TCA) in serum and decreased SBAs (THDCA, oMCA and ToMCA). Administration of the CA was reported to improve energy metabolism in diet-induced obese (DIO) mice.\(^53\)\(^54\) Increased levels of circulating PBA correlated with *A. muciniphila* in association with improved glucose and lipid metabolism in the bile diversion mouse model and improved endothelial function in Apoe\(^-/-\) mice fed inulin-type fructans.\(^25\)\(^54\) Although our results indicated a positive association between serum PBA and *Akermania* abundance, studies investigating cause-effect relationships are needed. BAs modify the gut microbiota, which in turn dictates BA diversity. The gut microbial alterations we observed likely resulted from combined actions of GPs and BAs. GPs may also influence the gut microbiota and FXR activity by sequestration of BAs, as this activity was recently reported for PACs extracted from persimmon fruit.\(^35\)

GP-treated mice had higher hepatic *Cyp7a1* expression compared with controls suggesting increased PBA synthesis contributed to elevated serum CA and TCA rather than increased ileal BA absorption. Hepatic expression of *Cyp7a1* is negatively controlled by intestinal and hepatic FXR activation.\(^28\) Gene expression of hepatic BA transporters (*Abcd1* and *Slc51b*) and acute phase protein Lcn13 (*Obp2a*) was similar between groups suggesting GPs did not alter hepatic FXR activity. Intestinal FXR activity was decreased as evidenced by reduced expression of *Fgf15*, *Shp*, and *Tgr5* (figure 4A). Increased *Cyp7a1* expression may be mediated by inhibition of intestinal FXR, likely driven by GP-induced changes in gut microbiota and BA diversity. Mice overexpressing *Cyp7a1* were protected from DIO, glucose intolerance, insulin resistance and dyslipidemia in association with reduced adiposity and increased energy expenditure via activation of uncoupling protein-1 (UCP-1) in brown adipose tissue.\(^55\)\(^56\) Inhibition of ileal FXR represents a key pathway connecting the gut microbiota and BAs with glucose metabolism. Metformin reduced abundance of *Bacteroides fragilis* in subjects with T2D resulting in increased circulating levels of glycuronosodeoxycholic acid, which inhibited intestinal FXR leading to improved serum glucose levels.\(^11\) Previous studies in DIO mice showed that inhibition of intestinal FXR by caffeic acid phenethyl ester, tempol, or antibiotics improved glycemia by decreasing synthesis of intestinal ceramides and their subsequent migration via lymph to the liver where they promote oxidative stress, insulin resistance, and gluconeogenesis.\(^10\)\(^36\) GP supplementation resulted in downregulation of *Smpd3* in the intestine and *Splt2* and *Cer4* in liver concomitant with diminished mRNA levels of genes involved in hepatic lipogenesis (*Chrebp*), gluconeogenesis (*G6pase*), β-oxidation (*Idh3a*), and inflammation (*Idh3a*). A recent study using loss and gain of function models showed that decreasing hepatic ceramides improved glucose tolerance, insulin resistance, and reduced hepatic steatosis in mice.\(^37\) As GP-treated mice had lower expression of ceramide synthesis genes, improvements in glucose metabolism may be linked to lower hepatic ceramide levels; however, studies using intestine-specific FXR KO mice and gnotobiotic/antibiotic-treated mice and ceramide quantification are needed for confirmation.
Different types or mixtures of dietary polyphenols likely have differential effects on the gut microbiota, BA profile, and FXR signaling. In Caco-2 cells, PAGs inhibited FXR signaling and p-coumaric acid, a microbial-derived PAC metabolite, inhibited FXR in a yeast 2-hybrid assay. PACs were reported to directly modulate FXR signaling and attenuate hyperglycemia but it remains unclear whether this effect was due to FXR activation or suppression. FXR signaling has been reported to both induce and prevent metabolic impairments, underlining its complex role in host energy metabolism. Organoid experiments suggested that SBAs THDCA, αMCA, ToMCA and PBA αMCA are FXR agonists (figure 5). GP-induced depletion of these agonistic BAs in vivo is therefore consistent with the observed downregulation of Fxr-responsive genes (Fig15, Shp, Tgr5) in ileum tissue. Moreover, GP supplementation increased levels of serum TCA, an FXR antagonist in our organoid system. TCA was also reported as an FXR agonist, however, this discrepancy does not appear to be due to our experimental conditions as the activities of well-established FXR agonist CDCA and antagonist TβMCA were highly reproducible in our system. GP supplementation induced a state of FXR inhibition despite increased concentrations of DCA and CA, which was reported to be a weak FXR agonist. These data suggest that GP-induced FXR inhibition is determined by the overall balance of agonistic and inhibitory BAs. While our data suggest that GP-induced depletion of agonistic SBAs resulted in FXR inhibition, the exact mechanism(s) involved warrant further investigation.

In summary, PAC-rich GPs reduced gut bacteria associated with production of SBAs identified as FXR agonists leading to FXR inhibition and improved glucose metabolism, potentially via reduced ceramide biosynthesis. These findings suggest a BA-FXR signaling axis as a putative novel mechanism to explain glucoregulatory effects of dietary PACs.

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