Simplified Intestinal Microbiota to Study Microbe-Diet-Host Interactions in a Mouse Model

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Graphical Abstract

Highlights

- Mice are colonized with ten bacterial strains to create a simple human microbiota model
- Dietary changes alter colonization patterns of the simplified intestinal microbiota (SIM)
- SIM-diet interactions affect some circulating metabolites in the host
- The SIM affects host metabolism in a diet-specific manner

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In Brief
Kovatcheva-Datchary et al. develop a mouse model colonized with a simplified intestinal microbiota (SIM) to investigate the microbe-microbe and host-microbe interactions in the mammalian gut, focusing on host metabolism. They combine dietary interventions and different omics approaches to show the potential of the SIM model to study the microbe-diet-host interplay.

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SUMMARY

The gut microbiota can modulate human metabolism through interactions with macronutrients. However, microbiota-diet-host interactions are difficult to study because bacteria interact in complex food webs in concert with the host, and many of the bacteria are not yet characterized. To reduce the complexity, we colonize mice with a simplified intestinal microbiota (SIM) composed of ten sequenced strains isolated from the human gut with complementing pathways to metabolize dietary fibers. We feed the SIM mice one of three diets (chow [fiber rich], high-fat/high-sucrose, or zero-fat/high-sucrose diets [both low in fiber]) and investigate (1) how dietary fiber, saturated fat, and sucrose affect the abundance and transcriptome of the SIM community, (2) the effect of microbe-diet interactions on circulating metabolites, and (3) how microbiota-diet interactions affect host metabolism. Our SIM model can be used in future studies to help clarify how microbiota-diet interactions contribute to metabolic diseases.

INTRODUCTION

The human gut is populated with a dense and complex community of microbes, collectively known as the gut microbiota (Dethlefsen et al., 2007). The dominating phyla in the human gut are Firmicutes, Bacteroidetes, Actinobacteria, Proteobacteria, and Verrucomicrobia, but other phyla, such as Fusobacteria, Cyanobacteria, Lentisphaerae, Spirochaetes, and TM7, are also present (Arunugam et al., 2011; Qin et al., 2010). The gut microbiota is associated with many essential functions for host physiology (Sommer and Backhed, 2013). For example, a key function of microbes in the gut is to process complex carbohydrates that cannot be digested by host enzymes (Sonnenburg and Backhed, 2016) into short-chain fatty acids (SCFAs), primarily acetate, propionate, and butyrate, and organic acids such as succinate and lactate (Cummings and Macfarlane, 1991). These metabolites are not only essential for the growth and cellular function of certain microbes in the gut (Cummings and Macfarlane, 1991; Koh et al., 2016; Macfarlane and Macfarlane, 2012). Furthermore, the gut microbiota produces other metabolites that are influenced by the diet, many of which are likely to play a role in host physiology (Koh et al., 2016; Zhao et al., 2018).

Although the production of SCFAs resulting from the bacterial fermentation of fiber-rich diets is generally associated with beneficial metabolic effects, the increased energy harvest has also been proposed to contribute to diet-induced obesity in mice (Koh et al., 2016). Furthermore, the gut microbiota produces other metabolites that are influenced by the diet, many of which are likely to play a role in host physiology (Koh et al., 2016; Makki et al., 2018; Zmora et al., 2019). Studies to delineate the interactions between gut bacteria, diet, and host metabolism are challenging because (1) of the complexity and high inter-individual variability of human gut microbiota, (2) bacteria interact in complex food webs in concert with the host, and (3) the microbiota consists mainly of non-sequenced members, thus limiting interpretation from metagenomic and metatranscriptomic analyses. One approach to overcome these issues is to use gnotobiotic animals colonized with a simplified intestinal microbial community consisting of well-characterized bacteria from humans. A recent example of such a model used rats colonized with eight bacterial species from the human gut to investigate how the microbiota composition changes in response to dietary challenges (Becker et al., 2011). Others have used mice colonized with a defined community of human bacteria to investigate microbiome–microbe interactions (McNulty et al., 2013; Rey et al., 2013) or the interactions between microbiota, dietary fiber, and the
RESULTS AND DISCUSSION

Colonization Pattern of the SIM Community in Mice on a Chow Diet

We introduced the ten selected bacteria (Table 1) into germ-free (GF) Swiss Webster mice and maintained them for four generations to generate stably colonized SIM mice. We showed that all ten bacteria of the SIM community colonized each region of the gut of SIM mice on a chow diet, although *Eubacterium hallii* was present only at a low density throughout (Figure 1A). Levels of SIM bacteria were higher in the distal part of the gut compared with the small intestine, with the highest levels in the feces of SIM mice (Figure 1A), consistent with the fact that the SIM bacteria were chosen because of their importance for carbohydrate fermentation.

A previous metagenome analysis has shown that all of the SIM bacteria, with the exception of *Roseburia inulinivorans*, are frequent colonizers of the human gut (Qin et al., 2010). We confirmed that the ten taxa of the SIM community were present in feces from two healthy human donors (Figures 1B and 1C). For comparison with SIM mice, we then colonized GF mice with unfractonated microbiota from each of the two human donors for 2 weeks. We showed that all ten SIM bacteria were present in the cecum of chow-fed mice colonized with feces from one of these donors (Figure 1B) and all except *Collinsella aerofaciens* were present in the cecum of chow-fed mice colonized with microbiota from the second donor (Figure 1C). This unsuccessful colonization of *C. aerofaciens* may be explained by the fact that the second donor had low fecal levels of *C. aerofaciens* and that this species must compete with other members of the human microbiota beyond the ten present in the SIM community. However, the overall profiles of the ten SIM bacteria were similar in the SIM mice and the humanized mice on a chow diet, indicating that the SIM community may be a suitable simplified model of the human gut microbiota.

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**Table 1. Phylogenetic and Metabolic Features of the Members of the SIM**

| SIM Bacterium                  | Phylum                  | Metabolic Function                  | Produced Metabolites       | References                        |
|-------------------------------|-------------------------|-------------------------------------|---------------------------|-----------------------------------|
| Akkermansia muciniphila: DSM 22959 | Bacteroidetes           | polysaccharide degradation          | acetate, propionate, formate | Derrien et al. (2004)             |
| Bacteroides thetaiotaomicron: ATCC 29148 | Actinobacteria         | polysaccharide degradation          | succinate, H₂S            | Martens et al. (2008); Xu et al. (2003) |
| Bifidobacterium adolescentis: L2-32 | Proteobacteria         | di- and oligosaccharide degradation | acetate, lactate          | Duranti et al. (2016); Marquet et al. (2009); Pokusaeva et al. (2011) |
| Collinsella aerofaciens: DSM 3979 | Actinobacteria         | di- and oligosaccharide degradation | acetate, lactate, H₂       | Flint et al. (2012); Kageyama and Benno (2000) |
| Desulfovibrio piger: DSM 749 | Proteobacteria         | sulfate reducer, lactate user       | acetate, H₂S              | Marquet et al. (2009)             |
| Eubacterium hallii: L2-7     | Firmicutes              | di- and monosaccharide degradation  | butyrate                  | Duncan et al. (2004b); Scott et al. (2014) |
| Eubacterium rectale: A1-86    | Firmicutes              | oligosaccharide degradation         | butyrate, lactate, formate, H₂ | Duncan and Flint (2008); Duncan et al. (2008) |
| Prevotella copri: DSM 18205  | Bacteroidetes           | polysaccharide degradation          | succinate, H₂             | Flint et al. (2012); Hayashi et al. (2007); Kovatcheva-Datchary et al. (2015) |
| Roseburia inulinivorans: A2-194 | Firmicutes              | oligosaccharide degradation         | butyrate, propionate      | Duncan et al. (2006); Scott et al. (2006) |
| Ruminococcus bromii: L2-63    | Firmicutes              | polysaccharide degradation          | acetate, formate, H₂      | Crost et al. (2018); Ze et al. (2012) |

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Colonic mucus barrier (Desai et al., 2016). However, none of these studies investigated how diet-induced changes in the gut microbiota affect host metabolism in parallel with altered microbial gene expression.

To investigate the effect of microbiota-diet interactions on host metabolism, we developed a gnotobiotic mouse model colonized with ten representatives of the human intestinal microbiota (simplified intestinal microbiota [SIM]). Our selection was based on the following criteria: (1) All members of the dominant phyla in the human gut should be represented. (2) The chosen strains must have been sequenced, thus allowing us to characterize how specific nutrients (fiber, saturated fat, and sucrose) affect the microbial gene expression. (3) The chosen strains must have well-characterized metabolic functions (to digest and/or ferment carbohydrates) and trophic interactions with a range of cross-feeding interactions for dietary conversions in the gut. The main metabolic function and metabolites produced by each SIM bacterial strain in *in vitro* studies are presented in Table 1. Because H₂ is produced during fermentation and accumulation of H₂ decreases the metabolic activity of microbes (Rey et al., 2013), we also included the H₂-consuming sulfate-reducing bacterium *Desulfovibrio piger*.
Dietary Changes Modify the Colonization Pattern of the Cecal SIM Community

A major selection criterion of the SIM bacteria was their capacity to metabolize carbohydrates and participate in anaerobic food webs in the gut. We therefore assessed how the abundance of SIM members was affected by changing the mouse diet from chow, which is low in fat (9% by weight) and high in dietary fiber (i.e., plant polysaccharides; 15% by weight), to one of two purified diets with low dietary fiber (primarily cellulose) but different macronutrient compositions. One group of SIM mice received a “Western” high-fat/high-sucrose (HF-HS) diet (fat 20% by weight, sucrose 18% by weight) for 2 weeks; a second group received a zero-fat/high-sucrose (ZF-HS) diet (sucrose 63% by weight) for 2 weeks (Table S1); a third group remained on chow.

The total bacterial load in the cecum of SIM mice was not affected by a change in the diet (Figure 2A). However, compared with SIM mice that remained on chow, SIM mice that switched to a diet low in dietary fiber had reduced cecal abundance of Prevotella copri and Bifidobacterium adolescentis (with either HF-HS or ZF-HS diet), Ruminococcus bromii (with HF-HS diet), and R. inulinivorans (with ZF-HS diet) as well as a trend toward reduced abundance of C. aerofaciens (with ZF-HS diet; p = 0.052) (Figure 2A). Thus, a reduction in dietary fiber reduced the abundance of these fiber-degrading bacteria. These results are consistent with earlier studies showing that P. copri, B. adolescentis, R. inulinivorans, R. bromii, and C. aerofaciens have increased abundance when the diet is rich in complex plant polysaccharides (Kovatcheva-Datchary et al., 2015; Ramirez-Farias et al., 2009; Scott et al., 2014; Walker et al., 2011; Vital et al., 2018). Furthermore, a recent study reported that bacteria from the phylogenetic order Bacteroidales (to which Prevotella species belong) are lost in mice after several generations of feeding with a Western diet (Sonnenburg et al., 2016). Similarly, a loss of Prevotella strains has been reported in the gut microbiome of non-Western individuals after immigration to a Western country (Vangay et al., 2018). In addition, two recent studies in humans showed that short-term dietary changes to reduce complex carbohydrates resulted in rapid reductions in abundance of Bifidobacterium, Ruminococcus, and Roseburia, bacteria that metabolize complex carbohydrates in the gut (David et al., 2014; Mardinoglu et al., 2018).

The cecal abundance of E. hallii was higher in SIM mice that switched from chow to an HF-HS or a ZF-HS diet (Figure 2A). In agreement, a dietary intervention study in humans showed that resistant starch (a prominent dietary fiber) reduced the abundance of E. hallii (Salonen et al., 2014). Furthermore,
in vitro studies have shown that *E. hallii* can use a broad range of substrates, including glucose, lactose, galactose, glycerol, and amino sugars (Bunesova et al., 2018; Duncan et al., 2004a; Engels et al., 2016; Scott et al., 2014). We observed a non-significant increase in the abundance of *Akkermansia muciniphila* and no change in the abundance of *Bacteroides thetaiotaomicron* or *Eubacterium rectale* in SIM mice that switched from chow to an HF-HS or a ZF-HS diet (Figure 2A), consistent with the fact that these bacteria are not exclusively dependent on dietary fiber and can degrade host mucin or simple sugars from the diet. *A. muciniphila* is an avid user of mucin glycans (Derrien et al., 2004). *B. thetaiotaomicron* is able to digest a broad range of dietary polysaccharides and adapts to different sources of carbohydrates, including mucin glycans, in the absence of dietary fibers (Sonnenburg et al., 2005). *E. rectale* can metabolize different carbohydrates (amylopectin, amylose, arabinoxylan, fructo-oligosaccharide, galacto-oligosaccharide, inulin, xylo-oligosaccharide) (Sheridan et al., 2016) but has also been shown to grow well on glucose (Cockburn et al., 2015; Ze et al., 2012).

The abundance of *D. piger* was also not affected by a change in diet (Figure 2A). The growth and survival of *D. piger* is dependent on the availability of sulfate (Rey et al., 2013), which in the mammalian gut is sourced not only from the diet but also from sulfated glycans in host mucin. Although *D. piger* lacks sulfatase activity, *A. muciniphila* and *B. thetaiotaomicron* are capable of removing sulfate from sulfated glycans when metabolizing host glycans and thus can contribute to the availability of sulfate required by *D. piger* (Rey et al., 2013).

There were no significant differences in individual bacterial species between mice on an HF-HS or a ZF-HS diet (Figure 2A), indicating that a reduction in plant polysaccharides has a greater influence on the abundance of the SIM members than variations in either fat or sucrose.

**Dietary Changes Affect the Fermentative Capacity of the Cecal SIM Community**

We confirmed the functional capacity of the SIM community to metabolize carbohydrates by showing that levels of SCFAs and organic acids were dramatically higher in cecal samples from SIM mice compared with GF mice on a chow diet (Figure 2B). To investigate the functional response of the SIM microbiota to dietary changes, we also compared SCFA levels in cecal samples from SIM mice that switched to an HF-HS diet or a ZF-HS diet with those that remained on chow. We showed that a switch to a diet low in plant polysaccharides resulted in reduced cecal concentrations of the SCFAs acetate and propionate as well as of the organic acids lactate and succinate (Figure 2B), in parallel with the reduced abundance of *P. copri*, *B. adolescentis*, *R. bromii*, and *R. inulinivorans* observed in response to this diet change (Figure 2A).

Butyrate was not significantly affected when SIM mice switched to a ZF-HS or an HF-HS diet (Figure 2B), consistent with the unchanged/increased abundance of the known butyrate producers *E. rectale* and *E. hallii* in response to these diets change (Figure 2A). Previous *in vitro* findings have shown that *E. hallii* can produce butyrate from glucose, lactate, acetate, and mucin-derived monosaccharides (Bunesova et al., 2018; Duncan et al., 2004b; Scott et al., 2014; Shetty et al., 2017).
Dietary Changes Modify the Transcriptome of the Cecal SIM Community

To further investigate the functional response of the SIM community to dietary changes, we performed RNA sequencing on cecal microbiota from the SIM mice that remained on chow or switched to an HF-HS or a ZF-HS diet for 2 weeks. We obtained an average depth of 15 million paired-end reads per sample, which was sufficient to cover the genomes of each SIM bacterium (Figure 2C). We identified differences (adjusted \( p < 0.05 \)) in 5,765 and 3,134 genes in cecal samples from the mice that switched to an HF-HS and a ZF-HS diet, respectively, compared with SIM mice that remained on chow (Table S2). By contrast, only 166 genes were significantly differently expressed when comparing samples from the mice on an HF-HS versus a ZF-HS diet (Table S2). These results demonstrate that gene expression in the SIM cecal microbiota is influenced to a greater extent by a reduction in plant polysaccharides than variations in either fat or sucrose.

We compared differences in gene expression at the species level in SIM mice fed an HF-HS or a ZF-HS diet with SIM mice fed chow and observed that (1) \( E. \) hallii, \( A. \) muciniphila, \( B. \) thetaiotaomicron, and \( D. \) piger had mostly upregulated genes, consistent with increased or no change in abundance of these species in the absence of plant polysaccharides, and (2) \( P. \) copri had exclusively downregulated genes, consistent with its reduced abundance in the absence of fiber (Figure 2C). In addition, \( R. \) inulinivorans, \( R. \) bromii, \( B. \) adolescentis, \( E. \) rectale, and \( C. \) aerofaciens had both upregulated and downregulated genes (Figure 2C), consistent with the fact that these SIM bacteria are capable of engaging in different trophic interactions (e.g., by competition and cooperation) to meet their nutrient requirements. These species can be primary degraders of dietary carbohydrates (Flint et al., 2012; Pokusaeva et al., 2011; Scott et al., 2011; Sheridan et al., 2016; Ze et al., 2012) and are also involved in metabolic cross-feeding interactions to maintain gut homeostasis (Belenguer et al., 2006; Crost et al., 2018; Flint et al., 2015; Marquet et al., 2009).

To explore functional features, we next used KEEG orthology (KO) and the protein domain database Pfam) to annotate genes that were significantly altered in cecal samples from mice fed HF-HS or ZF-HS compared with mice fed chow for each member of the SIM microbiota (Table S2). A switch to reduced plant polysaccharides in the diet resulted in (1) decreases in KOs associated with the fermentation of plant polysaccharides (e.g., alpha-amylase, beta-glucosidase, endoglucanase, and xylan 1,4-beta-xylolysidase), specifically in \( P. \) copri, \( E. \) rectale, and \( B. \) adolescentis; (2) increases in KOs associated with the conversion of simple sugars (such as ribose, ribulose, xylulose, fructose, mannose, sucrose, and glucose), amino sugars, pyruvate, and SCFAs (mostly butyrate and propionate), particularly in \( B. \) thetaiotaomicron, \( A. \) muciniphila, \( E. \) hallii, and \( R. \) inulinivorans; and (3) increases in KOs associated with the metabolism of mucin, particularly in \( A. \) muciniphila related to the degradation of mucin O-glycans and in \( E. \) hallii related to the metabolism of mucin monosaccharides (primarily galactose, glucose, and N-acetylglucosamine) (Table S2). For most of the genes that were significantly altered in SIM mice that switched to reduced plant polysaccharides in the diet, we observed similar annotations with both Pfam domains and KOs (Table S2). The most frequent Pfam annotations in genes that were significantly altered included those that were related to amino acid metabolism and to the transport of degraded carbohydrates (Table S2).

Dietary Changes Affect Carbohydrate-Active Enzyme Genes in the Cecal SIM Community

To investigate how dietary changes affect the contribution of the individual bacteria to the conversion of dietary carbohydrates, we next analyzed changes in SIM transcripts encoding carbohydrate-active enzymes (CAZymes) in cecal samples from mice fed an HF-HS or a ZF-HS diet compared with mice fed chow using the meta server dbCAN2, which integrates three tools for CAZyme genome annotation (Zhang et al., 2018). We observed similar results with all three tools (Table S3). In total, we identified diet-induced alterations in genes encoding 192 unique CAZymes belonging to 24 different families, including (1) enzymes involved in the assembly of carbohydrates such as glycosyl transferases (Lairson et al., 2008); (2) enzymes involved in carbohydrate breakdown including auxiliary activity enzymes, carbohydrate esterases, glycoside hydrolases, and polysaccharide lyases; and (3) accessory modules such as carbohydrate-binding modules (El Kaoutari et al., 2013) (Table S3).

A subset of these CAZymes showed the same pattern of expression changes in SIM mice that switched to either an HF-HS or a ZF-HS diet compared with SIM mice that remained on chow (Figure 3). For example, we observed reduced gene expression of \( P. \) copri-affiliated CAZymes potentially involved in the conversion of plant polysaccharides (including cellulose, hemicellulose, beta-glucans, beta-mannan, pectin, and starch) in mice that switched to either of the fiber-reduced diets (Figure 3), in parallel with the reduced abundance of \( P. \) copri in these mice (Figure 2A) and consistent with studies showing that \( Prevotella \) abundance is associated with fiber intake in humans (David et al., 2014; Wu et al., 2011). In addition, we observed increased gene expression of \( A. \) muciniphila-affiliated CAZymes potentially involved in the metabolism of host glycans and mucin in mice that switched to either of the fiber-reduced diets (Figure 3). A recent study in mice colonized with a synthetic human gut microbiota reported enhanced mucin degradation in response to a fiber-deprived diet (Desai et al., 2016). In this earlier study, \( A. \) muciniphila and \( Bacteroides \) caccae but not \( B. \) thetaiotaomicron contributed to the degradation of host glycans and mucin. However, it is likely that \( B. \) thetaiotaomicron is outcompeted by \( B. \) caccae if both are present; \( B. \) caccae had a higher colonization rate in mice colonized with selected human bacteria and fed a fiber-free diet (Desai et al., 2016) and has earlier been shown to have a higher growth rate on mucin and host glycans in vitro (Desai et al., 2016; McNulty et al., 2013).

A number of the diet-induced changes in CAZyme gene expression were specific to the dietary change (Figure 3). For example, in mice that switched to an HF-HS diet, we observed increased gene expression of CAZymes associated with conversion of plant polysaccharides, degradation of mucin, and host glycans, and synthesis of lipopolysaccharide and affiliated with \( E. \) hallii, \( A. \) muciniphila, \( R. \) inulinivorans, and \( B. \) adolescentis.
In contrast, only a few transcripts were increased specifically in mice that switched to a ZF-HS diet (Figure 3); these transcripts mostly belong to glycosyl transferase families, which are involved in carbohydrate assembly rather than degradation (Lairson et al., 2008). Furthermore, we observed reduced gene expression of *E. rectale*-affiliated CAZymes associated with conversion of plant polysaccharides mainly in SIM mice that switched to a ZF-HS diet (glycoside hydrolase 94 was the only *E. rectale*-affiliated CAZyme that showed reduced gene expression when SIM mice switched to either an HF-HS or a ZF-HS diet) (Figure 3).

Together these findings show the flexibility of the SIM community to efficiently metabolize the carbohydrates from the diet. They also indicate that the diet-induced reduction in the fermentative capacity of the SIM community is not a result of high levels of fat in the diet but rather is caused by the reduction in carbohydrates that can be metabolized by the bacteria.

### Dietary Changes Affect SIM-Produced Metabolites in the Portal Vein

To investigate the effect of microbe-diet interactions on circulating metabolites, we performed metabolomics on plasma samples collected from the portal vein of the SIM mice that switched to an HF-HS or a ZF-HS diet for 2 weeks and from those that remained on chow. To discriminate between host and microbially derived plasma metabolites, we also analyzed plasma samples from GF mice on the three different diets. In total, 548 metabolites were measured (Table S4); of these, metabolites whose levels were significantly altered by the microbiota (75) or by diet change (89) are shown in Figures 4A and 4B, respectively. We identified 17 metabolites that were both microbially regulated and modulated by dietary changes in plasma samples from SIM mice (shown in red text in Figures 4A and 4B). These metabolites were mostly linked to lipid and amino acid metabolism.

Specifically, the 17 microbially regulated metabolites that were affected by reducing the consumption of dietary fiber were (1) the unsaturated fatty acids oleate and 10-nanodecanoate (increased in SIM mice on HF-HS or ZF-HS versus chow); (2) the acylcarnitines 2-methylbutyrylcarnitine, 2-methylbutyrylglycine, isobutyrylcarnitine, propionylcarnitine, and propionylglycine; the purine metabolite xanthosine; succinate; and ferulic acid 4-sulfate (decreased in SIM mice on HF-HS or ZF-HS versus chow); and (3) kynurenate, anthranilate, and xanthurenate, key metabolites in the kynurenine pathway of

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(Figure 3).
tryptophan catabolism (decreased in SIM mice on HF–HS versus chow and increased in SIM mice on ZF–HS versus chow) (Figure 4B; Table S4).

The reduction of succinate in the portal vein of SIM mice on an HF–HS or a ZF–HS diet versus chow was consistent with reduced cecal levels of succinate (Figure 2B). *P. copri* is an efficient succinate producer (Franke and Deppenmeier, 2018; Kovatcheva-Datchary et al., 2015), and we have previously demonstrated that succinate can improve metabolism by stimulating intestinal gluconeogenesis (De Vadder et al., 2016). Ferulic acid 4-sulfate, which was also reduced in the portal vein of SIM mice on an HF–HS or a ZF–HS diet versus chow, is obtained after conjugation of ferulic acid during its uptake in the intestinal epithelium and passage through the liver (Bourne and Rice-Evans, 1998). The beneficial effect of ferulic acid and ferulic acid 4-sulfate in diabetes, cardiovascular diseases, improved lipid metabolism, and blood pressure lowering has been well recognized (Alam et al., 2016; Van Rymenant et al., 2017; Vinayagam et al., 2016). Thus, dietary fiber-induced increases in succinate and ferulic acid 4-sulfate may be beneficial for host metabolism.

To investigate if we could identify links between any of these 17 portal vein metabolites (which were both microbiobly regulated and modulated by dietary changes) and SIM-regulated genes, we searched for SIM- and diet-regulated operons in the metatranscriptome and investigated whether their resulting genes could contribute to SIM- and diet-induced regulation of the portal vein metabolites. We identified one operon (of *E. rectale* consisting of EUBREC_1041 and EUBREC_1042) that was significantly reduced in the cecal transcriptome of SIM mice on a ZF–HS diet versus chow (Table S2). EUBREC_1041 encodes acyl-CoA thioesterase (Mahowald et al., 2009) and EUBREC_1042 encodes arabinofuranosidase (also termed xylan 1,4-beta-xylosidase), which can release ferulic acid from dietary fibers (Duncan et al., 2016; Nishizawa et al., 1998). We next searched the CAZyme dataset for enzyme families that could contribute to microbiota activity associated with ferulic acid release. We observed that transcripts of the carbohydrate esterase family CE1, which includes ferulic acid esterase activity (Dodd and Cann, 2009), were decreased in cecal samples from mice on a ZF–HS diet versus chow, with a major contribution of *E. rectale* (Figure 3). Thus, we propose that *E. rectale*, which is abundant in the human microbiome, may contribute to ferulic acid release from diets rich in plant polysaccharides in the mammalian gut.

**SIM Affects Host Metabolism in a Diet-Specific Fashion**

To investigate whether the SIM system also could be used to study microbe-diet effects on host metabolism, we compared body weight, adiposity, steatosis, and glucose metabolism in SIM versus GF mice on each of the three diets. In mice on chow, we observed that SIM was sufficient to increase body weight and epididymal adipose weight, induce hepatic steatosis, and increase blood glucose levels (Figures 5A–5E). Thus, SIM has effects on host metabolism similar to those seen in mice colonized with a normal mouse microbiota (Bäckhed et al., 2004; Caesar et al., 2012) and may induce adiposity, resulting in impaired glucose tolerance by enhancing energy harvest (i.e., increased SCFA production from dietary fibers) in the host.

In mice on an HF–HS diet, we again demonstrated that SIM increased blood glucose levels (Figure 5E). However, this response was not paralleled by an increase in body weight (Figure 5A), in contrast to our earlier study comparing HF–HS diet-fed conventionally raised and GF mice (Bäckhed et al., 2007). This discrepancy may be explained by the fact that the present diet intervention was shorter than in the previous study (2 versus 8 weeks) and that another mouse strain was used (Swiss Webster versus C57BL/6). However, we observed increased adiposity and steatosis in SIM versus GF mice on an HF–HS diet (Figures 5B–5D), and both of these responses are known to be associated with impaired glucose tolerance (Cani et al., 2007).

In mice on the most extreme diet, ZF–HS, we observed that SIM had its main effect on hepatic steatosis, resulting in a 2.8 ± 0.5 fold increase in liver fat compared with GF mice on the same diet (Figure 5D). SIM promoted a small increase in body weight and epididymal white adipose weight (Figures 5A and 5C), without affecting overall adiposity determined by magnetic resonance imaging (Figure 5B) in mice on a ZF–HS diet. Surprisingly, we did not observe significant changes in blood glucose levels in SIM mice compared with GF mice on the ZF–HS diet (Figure 5E), despite the difference in hepatic steatosis. One explanation can be that the ZF–HS diet has direct effects on glucose metabolism in GF mice, independently of steatosis, as the glucose excursion curve was elevated in GF mice on this diet compared with GF mice fed chow or an HF–HS diet (Figure 5E).

**Prospectus**

Studies of the gut microbiota are complex and often limited by a lack of sequenced strains and poor gene annotations. To overcome these limitations, we colonized GF mice with a set of relatively well-characterized bacteria that represent the main phyla in the human gut and used the model to investigate how SIM-diet interactions affect the transcriptome, metabolome, and host metabolism. The SIM community has many of the features of the complete microbiota in modulating host metabolism in a diet-dependent fashion and, as such, the SIM model may facilitate our understanding of how the microbes interact with macronutrients to affect host metabolism. Because we use bacterial strains isolated from humans, our findings may be extrapolated to humans and potentially provide guidance for human...
intervention studies. However, further experiments that consider the complexity of human diets and the variety of dietary fibers are required to determine how well data from our simplified system can be translated into human physiology.

STAR METHODS

Detailed methods are provided in the online version of this paper and include the following:

- KEY RESOURCES TABLE
- CONTACT FOR REAGENT AND RESOURCE SHARING
- EXPERIMENTAL MODEL AND SUBJECT DETAILS
  - Colonization of mice with the SIM bacteria
  - Colonization of mice with human feces
  - Dietary experiments with SIM mice
  - Bacteria strains
- METHOD DETAILS

SUPPLEMENTAL INFORMATION

Supplemental Information can be found with this article online at https://doi.org/10.1016/j.celrep.2019.02.090.

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AUTHOR CONTRIBUTIONS

P.K.-D. and F.B. conceived and designed the project. P.K.-D., A.H., and A.W. performed experiments. I.N. participated in the functional analysis of data. P.K.-D., S.S., I.N., A.W., R.P., J.N., and F.B. analyzed and interpreted the data. P.K.-D., S.S., R.P., and F.B. wrote the paper. All authors commented on the manuscript.

DECLARATION OF INTERESTS

J.N. and F.B. are founders and shareholders of Metabogen AB.

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## STAR METHODS

### KEY RESOURCES TABLE

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| **Bacterial and Virus Strains** | | |
| Akkermansia muciniphila | Prof. Willem M. de Vos | DSM 22959 |
| Bacteroides thetaiotaomicron | ATCC | 29148 |
| Bifidobacterium adolescentis | Dr. Karen Scott | L2-32 |
| Collinsella aerofaciens | Dr. Karen Scott | DSM 3979 |
| Desulfovibrio piger | Dr. Karen Scott | DSM 749 |
| Eubacterium hallii | Dr. Karen Scott | L2-7 |
| Eubacterium rectale | Dr. Karen Scott | A1-86 |
| Prevotella copri | DSM | 18205 |
| Roseburia inulinivorans | Dr. Karen Scott | A2-194 |
| Ruminococcus bromii | Dr. Karen Scott | L2-63 |
| **Chemicals, Peptides, and Recombinant Proteins** | | |
| Bacto casitone | BD | Cat#225910 |
| Yeast extract | Oxoid | Cat#LP0021 |
| Sodium bicarbonate | Merck | Cat#1.06329.0500 |
| Glucose/microbiology | Merck | Cat#346351 |
| Cellobiose | Sigma-Aldrich | Cat#22150 |
| Starch | Sigma-Aldrich | Cat#59765 |
| Di-potassium hydrogen phosphate | Sigma-Aldrich | Cat#P3786 |
| Di-hydrogen potassium phosphate | Merck | Cat#1.04873.0250 |
| Di-ammonium sulfate | Sigma-Aldrich | Cat#A4418 |
| Magnesium sulfate heptahydrate | Sigma-Aldrich | Cat#M7634 |
| Calcium chloride dihydrate | Merck | Cat#1.02382.0500 |
| Acetic acid | Sigma-Aldrich | Cat#A6283 |
| Propionic acid | Sigma-Aldrich | Cat#P1386 |
| n-Valeric acid | Sigma-Aldrich | Cat#240370 |
| Isovaleric acid | Sigma-Aldrich | Cat#129542 |
| Isobutyric acid | Sigma-Aldrich | Cat#1754 |
| Hemin | Sigma-Aldrich | Cat#51280 |
| Thiamine-HCl | Sigma-Aldrich | Cat#T1270 |
| Riboflavin | Sigma-Aldrich | Cat#R9504 |
| Biotin | Sigma-Aldrich | Cat#B4639 |
| Cobalamin | Sigma-Aldrich | Cat#V2876 |
| 4-Aminobenzoic acid | Sigma-Aldrich | Cat#A9878 |
| Folic acid | Sigma-Aldrich | Cat#F8758 |
| Pyridoxamine dihydrochloride | Sigma-Aldrich | Cat#P9158 |
| Cysteine | Sigma-Aldrich | Cat#C7477 |
| Resazurin | Sigma-Aldrich | Cat#R7017 |
| Rumen fluid | Prof. Hauke Smidt | N/A |
| Trypticase peptone | BD | Cat#211921 |
| Proteose peptone | Sigma-Aldrich | Cat#82450 |
| Beef extract | BD | Cat#212303 |
| Tween 80 | Sigma-Aldrich | Cat#P4780 |
| Vitamin K1 | Sigma-Aldrich | Cat#V3501 |

(Continued on next page)
| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Sodium sulfide nonahydrate | Sigma-Aldrich | Cat#208043 |
| Sodium butyrate – 13C4 | Sigma-Aldrich | Cat#488380 |
| Sodium Acetate –1-13C, d3 | Sigma-Aldrich | Cat#298042 |
| Propionic acid-d6 | Sigma-Aldrich | Cat#490644 |
| Succinic acid 13C4 99% | Loradan Fine Chemicals AB | Cat#CLM-1571 |
| Sodium lactate 13C4 98% | Loradan Fine Chemicals AB | Cat#CLM-1579 |
| Succinic acid (98%) | Loradan Fine Chemicals AB | Cat#15-0400 |
| Butyric acid | Sigma-Aldrich | Cat#19215 |
| Sodium acetate | Sigma-Aldrich | Cat#S8750 |
| Sodium propionate | Sigma-Aldrich | Cat#P1880 |
| Sodium DL-lactate | Sigma-Aldrich | Cat#71720 |
| N-tert-Butyldimethylsilyl-N-methyltrifluoroacetamide | Sigma-Aldrich | Cat#19915 |
| Diethyl ether | Sigma-Aldrich | Cat#309958 |
| Autoclavable Mouse Breeder Diet 5021 (Chow diet) | LabDiet | Cat#0006539 |
| Adjusted Fat Diet (HF-HS diet) | Envigo | TD.09683 |
| 62% Sucrose diet (ZF-HS diet) | Envigo | TD.03314 |
| EDTA | Sigma-Aldrich | Cat#EDS |
| TRIS | Sigma-Aldrich | Cat#76066-1KG |
| Ammonium acetate | Sigma-Aldrich | Cat#09688-1KG |
| Isopropyl | Sigma-Aldrich | Cat#9516-500ML |
| Macaloid | Laguna Clay | Cat#MBENMAC |
| DEPC Treated water | Sigma-Aldrich | Cat#95284 |
| RNase A Solution | QIAGEN | Cat#158922 |
| RNase I | Roche | Cat#4716728001 |
| UltraPure Phenol:Water (3.75:1; v/v) | Invitrogen | Cat#15594-047 |
| UltraPure Phenol:Chloroform:Isoamyl Alcohol (25:24:1, v/v) | Invitrogen | Cat#15593-049 |
| Chloroform:Isoamyl Alcohol (24:1, v/v) | Sigma-Aldrich | Cat#25666-100ml |

Critical Commercial Assays

| 1x SYBR Green Master Mix | Thermo Fisher Scientific | Cat#4309155 |
| RNeasy Mini Kit | QIAGEN | Cat#74106 |
| QIAMP DNA mini Kit | QIAGEN | Cat#51360 |
| RibonZero Magnetic Kit | Nordic Biolabs | Cat#MRZB1242 |
| Agilent RNA 6000 Pico Kit | Agilent Technologies | Cat#5067-1513 |
| ScriptSeq v2 RNA-Seq Library Preparation Kit | Nordic Biolabs | Cat#SSV2112 |
| FallSafe Enzyme Mix | Nordic Biolabs | Cat#FSE51100 |
| ScriptSeq Index PCR Primer | Nordic Biolabs | Cat#SSIP1234 |
| Agencourt AMPure XP | Beckman Coulter | Cat#A63880 |

Deposited Data

| RNA-Seq data | This paper | SRA: PRJEB22735 |
| PFAM | (Sonnhammer et al., 1997) | [https://pfam.xfam.org/](https://pfam.xfam.org/) |
| dbCAN2 | (Zhang et al., 2018) | [http://cys.bios.niu.edu/dbCAN2/](http://cys.bios.niu.edu/dbCAN2/) |
| Akkermansia muciniphila genome build | Ensembl bacteria | [ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_19_collection/akkermansia_muciniphila_atcc_baa_835](ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_19_collection/akkermansia_muciniphila_atcc_baa_835) |
| Bacteroides thetaiotaomicron genome build | Ensembl bacteria | [ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_0_collection/bacteroides_thetaiotaomicron_vpi_5482](ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_0_collection/bacteroides_thetaiotaomicron_vpi_5482) |

(Continued on next page)
### REAGENT or RESOURCE SOURCE IDENTIFIER

| REAGENT or RESOURCE | SOURCE | IDENTIFIER |
|---------------------|--------|------------|
| Bifidobacterium adolescentis genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_25_collection/bifidobacterium_adolescentis_atcc_15703/ |
| Collinsella aerofaciens genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_4_collection/collinsella_aerofaciens_atcc_25986/ |
| Desulfovibrio piger genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_12_collection/desulfovibrio piger_atcc_29098/ |
| Eubacterium hallii genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_2_collection/eubacterium_hallii_dsm_3353/ |
| Eubacterium rectale genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_24_collection/eubacterium_rectale_atcc_33656/ |
| Prevotella copri genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_20_collection/prevotella_copri_dsm_18205/ |
| Roseburia inulinivorans genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_20_collection/roseburia_inulinivorans_dsm_16841/ |
| Ruminococcus bromii genome build | Ensembl bacteria | ftp://ftp.ensemblgenomes.org/pub/release-38/bacteria/fasta/bacteria_21_collection/ruminococcus_bromii_l2_63/ |

### Experimental Models: Organisms/Strains

| Female Swiss Webster | Own breeding | N/A |
| Male Swiss Webster | Own breeding | N/A |

### Oligonucleotides

| Table S5 | N/A | N/A |

### Software and Algorithms

| HMMER | (Eddy, 1998) | http://hmmer.org/ |
| STAR | (Dobin et al., 2013) | https://github.com/alexdobin/STAR |
| Samtools | (Li et al., 2009) | http://samtools.sourceforge.net/ |
| HTSeq | (Anders et al., 2015) | http://htseq.readthedocs.io |
| edgeR | (Robinson et al., 2010) | https://bioconductor.org/packages/release/bioc/html/edgeR.html |
| Prism (version 6) | GraphPad | N/A |

### Other

| CFX96 Real-Time PCR Detection System | Bio-Rad | C1000 |
| Gas Chromatograph-Mass Spectrometer | Agilent Technologies | 7890A + 5975C series |
| pH/Ion meter | Mettler Toledo | S220 SevenCompact |
| Coy chamber | COY Laboratory Products | http://coylab.com/products/anaerobic-chambers/vinyl-anaerobic-chambers/ |
| Glucose meter | HemoCue AB | HemoCue Glucose 201+ |
| Centrifuge | Eppendorf | 5430R |
| Centrifuge | Thermo Scientific | Heraeus Megafuge 16R. (TX-400 Swinging Bucket Rotor) |
| INTELLI-MIXER with rack | LabTeamet | EL-RM-2L (EL-16mm test tube) |
| EchoMRI Body Composition Analyzers for Live Small Animals | Echo Medical Systems | http://www.echomri.com |
| PCR machine | Eppendorf | Vapo.Protect |
| Fast-Prep-24 Classic | MP Biomedicals | Cat#116004500 |

(Continued on next page)
CONTACT FOR REAGENT AND RESOURCE SHARING

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Fredrik Bäckhed (fredrik.backhed@wlab.gu.se).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Colonization of mice with the SIM bacteria

All the mouse experiments were performed using protocols approved by the University of Gothenburg Animal Studies Committee.

For all experiments, mice were maintained in plastic gnotobiotic research isolators housed in a climate-controlled room (22°C ± 2°C) and subjected to a 12 h light/dark cycle (7:00 a.m.–7:00 p.m.) with free access to autoclaved water and food. Mice were housed in groups (n = 4-5 mice/group) according to their dietary treatment.

To colonize mice with the SIM bacteria, adult female GF Swiss Webster mice were orally gavaged twice after a 4 h fast: first, with 0.2 mL of 10^8 CFU *B. thetaiotaomicron* to obtain a more reduced gut environment, and three days later with 0.2 mL of mix of 10^8 CFU from each of the rest of the SIM strains. Mice were then bred with male GF Swiss Webster mice to generate SIM mice. Mice were bred for four generations. To characterize the colonization pattern of SIM bacteria through the gut, intestinal segments, cecal content and feces were harvested from 8-week-old male SIM mice, immediately snap-frozen in liquid nitrogen and stored at −80°C until further processed.

Colonization of mice with human feces

To generate mice colonized with human feces, fecal samples from one woman and one man were separately added to PBS buffer supplemented with reducing solution (Na_2S and cysteine dissolved in NaHCO_3 buffer) and transferred by oral gavage to 10-12-week-old female and male GF Swiss Webster (5 per group), respectively. The mice were colonized for 14 days and fed regular chow diet. Cecum was harvested, immediately snap-frozen in liquid nitrogen and stored at −80°C until further processed. The selected adult human donors were healthy, did not have any special dietary requirements, and had not taken any medication in the 4 months before sample donation. The participants gave informed consent and the study was approved by the Ethics Committee at Gothenburg University.

Dietary experiments with SIM mice

Adult male SIM mice aged 12-16 weeks were randomized into 3 groups (5-7 mice/group) to receive: chow (5021 rodent diet, LabDiet; fat 9% wt/wt); HF-HS (fat 20% wt/wt; sucrose 18% wt/wt, 96132 Teklad Custom Research Diet) or ZF-HS (03314 Teklad Custom Research Diet; sucrose 62% wt/wt) (Table S1). Mice were fed their respective diet ad libitum for 2 weeks and maintained in separate plastic gnotobiotic research isolators. At the end of the study, blood was collected from the portal vein under deep isoflurane-induced anesthesia following a 4 h fast, and intestinal segments with contents were harvested. All tissues were immediately snap-frozen in liquid nitrogen and stored at −80°C until further processed.

Bacteria strains

*E. hallii* L2-7 (DSM 17630), *E. rectale* A1-86 (DSM 17629), *B. adolescentis* L2-32, *C. aerofaciens* DSM 3979, *D. piger* DSM 749, *R. inulinivorans* A2-194 (DSM 16841), and *R. bromii* L2-63 were obtained from Dr Karen Scott, the Rowett Institute of Nutrition and Health, Aberdeen, UK. *B. thetaiotaomicron* ATCC 29148 and *P. copri* DSM 18205 were obtained from ATCC and DSMZ,
respectively. *A. muciniphila* DSM 22959 was obtained from Professor Willem de Vos, Laboratory of Microbiology, Wageningen University, the Netherlands. All strains in this study originate from human feces.

*E. hallii*, *E. rectale*, *B. adolescentis*, *B. thetaiotaomicron*, *C. aerofaciens*, *R. inulinivorans* were maintained in Hungate-like tube (Ochs Laborbedarf, Germany) under CO₂ in YCFA medium (Ze et al., 2012). *R. bromii* was maintained in Hungate-like tubes in M2GSC media supplemented with rumen fluid (Miyazaki et al., 1997). *D. piger* was maintained in Hungate-like tubes in CO₂ in YCFA medium supplemented with 30 mM lactate. *P. copri* was grown in PYG medium (Media 104 in the DSMZ catalog), and *A. muciniphila* was grown in a basal mucin-based media (Derrien et al., 2004). All bacteria were cultured anaerobically in 15 mL Hungate tubes at 37°C in a Coy chamber.

**METHOD DETAILS**

**Body composition and glucose tolerance test**

Body composition (including liver fat) was determined using magnetic resonance imaging (EchoMRI, Houston, TX). For the glucose tolerance test, mice were fasted for 4 h before intraperitoneal injection of 30% glucose solution (1.5 mg/g body weight). Blood glucose levels from tail vein were measured at baseline, and after 15, 30, 60, 90 and 120 min.

Genomic DNA extraction and 16S rRNA quantitative PCR

After isolation of genomic DNA from intestinal segments, cecal content and feces using repeated bead beating (Salonen et al., 2010), 16S rRNA quantitative PCR was performed with a CFX96 Real-Time System (Bio-Rad Laboratories). Samples were analyzed in a 25 μL reaction mix consisting of 12.5 μL 1xSYBR Green Master Mix buffer (Thermo Scientific, Waltham, Massachusetts, USA), 0.2 μM of each primer and 5 μL of template DNA, water or genomic DNA extracted from feces or cecal content. Standard curves of 16S rRNA PCR product of *E. hallii*, *E. rectale*, *B. adolescentis*, *C. aerofaciens*, *D. piger*, *R. inulinivorans*, *R. bromii*, *B. thetaiotaomicron*, *P. copri* and *A. muciniphila* were created using serial 10-fold dilution of the purified PCR product. qPCR was performed as reported previously: for *E. hallii*, *E. rectale*, *B. adolescentis*, *R. inulinivorans*, *R. bromii* and total bacteria (Ramirez-Farias et al., 2009); for *C. aerofaciens* (Kageyama et al., 2000); for *D. piger* (File et al., 2004); for *P. copri* (Matsuki et al., 2002); for *B. thetaiotaomicron* (Samuel and Gordon, 2006); and for *A. muciniphila* (Collado et al., 2007) (Table S5). All reactions were performed in duplicate in one run and in duplicate PCR runs. The data are expressed as log of 16S rRNA copy per g of luminal content or feces.

Quantification of SCFAs and organic acids

SCFAs and organic acids in the cecum were measured by gas chromatography. For the extractions, 100-250 mg of frozen fecal content was transferred to a glass tube (16x125 mm) fitted with a screw cap and 100 μL of stock solution of internal standard (1 M [1-13C] acetate, 0.2 M [1-13C] propionate and [13C] butyrate, 0.5 M [13C] lactate and 40 mM [13C4] succinic acid) was added. Prior to extraction, samples were freeze-dried at −50°C for 3 h (yield 40–98 mg dry weight). The extraction and quantification of SCFAs and organic acids was performed as previously reported (Ryan et al., 2014).

**Microbial RNA extraction**

Approximately 100 mg frozen fecal content was re-suspended in 500 μL ice-cold TE buffer (Tris–HCl pH 7.6, EDTA pH 8.0). Total RNA was extracted from the re-suspended cell pellet according to the Macaloid-based RNA isolation protocol (Egert et al., 2007) with the use of Phase Lock Gel heavy (5 Prime, Hamburg) (Murphy and Hellwig, 1996) during phase separation. The aqueous phase was purified using the RNAeasy mini kit (QIAGEN, USA), including an on-column DNaseI (Roche, Germany) treatment as described previously (Egert et al., 2007). Total RNA was eluted in 30 μL ice-cold TE buffer and the RNA quantity and quality were assessed using a NanoDrop ND-1000 spectrophotometer (Nanodrop Technologies, Wilmington, USA).

**Metatranscriptome analysis of the SIM community**

The metatranscriptome datasets were generated by Illumina sequencing of 15 cDNA libraries derived from mRNA enriched samples of the cecal SIM microbiota. The mRNA enrichment was performed by the removal of 16S and 23S rRNA using sequence-based capture probes attached to magnetic beads [RibozeroTM Magnetic Kit (Bacteria), Epicenter, Illumina] using the manufacturer’s protocols. The enriched mRNA was quantified spectrophotometrically (NanoDrop) and its quality was assessed using an Agilent 2100 Bioanalyzer (Agilent Technologies). Thirty Illumina sequencing libraries were constructed from double-stranded cDNA prepared after RNA fragmentation according to the manufacturer’s protocols (ScriptSeq v2 RNA-Seq Library preparation Kit, Epicenter, Illumina). Each sequencing library was barcoded at the Genomics Core Facility of the Gothenburg University using Illumina Hisseq2000, which generates on average ~15 million reads for each sample. The reads were trimmed from adaptor sequences and quality (Phred quality score > 30), then simultaneously aligned to the corresponding bacterial genome of SIM using STAR 2.3.1u (Dobin et al., 2013). The resulting bam-files were indexed and sorted using Samtools 0.1.18 (Li et al., 2009). Gene counts were calculated using HTSeq-count 0.5.4p3 (Anders et al., 2015) based on the transcript annotation and globally normalized using a standard negative binomial approach through the R-package edgeR (Robinson et al., 2010). Circos was used to plot the genome coverage and the comparative analyses of differentially expressed genes (Krzywinski et al., 2009).
The genes were functionally annotated using KEGG orthology (KO), CAZymes and Pfam. We annotated CAZymes based on the HMMER, DIAMOND and Hotpep tools of dbCAN2 meta-server (Zhang et al., 2018) using the following: E-value < 1e-15, coverage > 0.35 for HMMER; E-value < 1e-102, hits per query (-k) = 1 for DIAMOND; and frequency > 6.0, hits > 2.6 for Hotpep. We annotated Pfam domains using HMM profiles of Pfam Database (version 31) by HMMER (default E-value, 10) (Eddy, 1998).

**Metabolomic analysis of plasma from SIM mice**

Aliquots of portal vein plasma from SIM mice fed chow, HF-HS or ZF-HS diets were analyzed by untargeted liquid chromatography - mass spectrometry (LC-MS) and gas chromatography - mass spectrometry (GC-MS) at Metabolon (Durham, USA).

**QUANTIFICATION AND STATISTICAL ANALYSIS**

Values are presented as means ± SEM. For graph plotting and statistical analysis we used GraphPad Prism (version 6, GraphPad Software, San Diego, CA, USA) unless otherwise indicated. Statistical comparison of two groups was performed by Student’s t test, comparisons of three or more groups were performed by one-way analysis of variance (ANOVA) and corrected for multiple comparison with Tukey post-tests.

**DATA AND SOFTWARE AVAILABILITY**

The RNA-seq data reported in this paper have been deposited in the ENA sequence read archive (http://www.ebi.ac.uk/ena/data/view) under accession number PRJEB22735.