Characterizing a model human gut microbiota composed of members of its two dominant bacterial phyla


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The adult human distal gut microbial community is typically dominated by 2 bacterial phyla (divisions), the Firmicutes and the Bacteroidetes. Little is known about the factors that govern the interactions between these members. Here, we examine the niches of representatives of both phyla in vivo. Finished genome sequences were generated from Eubacterium rectale and E. eligens, which belong to Clostridium Cluster XIVa, one of the most common gut Firmicute clades. Comparison of these and 25 other gut Firmicutes and Bacteroidetes indicated that the Firmicutes possess smaller genomes and a disproportionately smaller number of glycan-degrading enzymes. Germ-free mice were then colonized with E. rectale and/or a prominent human gut Bacteroidetes, Bacteroides thetaiotaomicron, followed by whole-genome transcriptional profiling, high-resolution proteomic analysis, and biochemical assays of microbial–microbial and microbial–host interactions. B. thetaiotaomicron adapts to E. rectale by up-regulating expression of a variety of polysaccharide utilization loci encoding numerous glycoside hydrolases, and by signaling the host to produce mucosal glycans that it, but not E. rectale, can access. E. rectale adapts to B. thetaiotaomicron by decreasing production of its glycan-degrading enzymes, increasing expression of selected amino acid and sugar transporters, and facilitating glycolysis by reducing levels of NADH, in part via generation of butyrate from acetate, which can access. These observations support the notion that gut microbes may live at the intersection of 2 forms of selective pressure: bottom-up selection, where fierce competition between members of a community that approaches a population density of 10^{11} to 10^{12} organisms per milliliter of colonic contents drives phyletic niche specialization and functional redundancy within members of its major bacterial phyla, and the importance of host glycans as a nutrient foundation that ensures ecosystem stability.

The adult human gut houses a bacterial community containing trillions of members comprising thousands of species-level (phylogenetic type) species. Culture-independent surveys of this community have revealed remarkable interindividual variabilities in these strain- and species-level phyotypes. Two bacterial phyla, the Firmicutes and the Bacteroidetes, commonly dominate this ecosystem (1), as they do in the guts of at least 60 mammalian species (2).

Comparative analysis of 5 previously sequenced human gut Bacteroidetes revealed that each genome contains a large repertoire of genes involved in acquisition and metabolism of polysaccharides. This repertoire includes (i) up to hundreds of glycoside hydrolases (GHs) and polysaccharide lyases (PLs); (ii) myriad paralogs of SusC and SusD, outer membrane proteins involved in recognition and import of specific carbohydrate structures (3); and (iii) a large array of environmental sensors and regulators (4). These genes are assembled in similarly organized, selectively regulated polysaccharide utilization loci (PULs) that encode functions necessary to detect, bind, degrade and import carbohydrate species encountered in the gut habitat—either from the diet or from host glycans associated with mucus and the surfaces of epithelial cells (5–7).

Studies of gnotobiotic mice colonized only with human gut-derived Bacteroides thetaiotaomicron have demonstrated that this organism can vary its pattern of expression of PULs as a function of diet, e.g., during the transition from mother’s milk to a polysaccharide-rich chow consumed when mice are weaned (5), or when adult mice are switched from a diet rich in plant polysaccharides to a diet devoid of these glycans and replete with simple sugars (under the latter conditions, the organism forages on host glycans) (6, 7).

Our previous functional genomic studies of the responses of B. thetaiotaomicron to cocolonization of the guts of gnotobiotic mice with Bifidobacterium longum, an Actinobacterium found in the intestines of adults and infants, or with Lactobacillus casei, a Firmicute present in a number of fermented diary products, have shown that B. thetaiotaomicron adapts to the presence of these other microbes by modifying expression of its PULs in ways that expand the breadth of its carbohydrate foraging activities (8).

These observations support the notion that gut microbes may live at the intersection of 2 forms of selective pressure: bottom-up selection, where fierce competition between members of a community that approaches a population density of 10^{11} to 10^{12} organisms per milliliter of colonic contents drives phyletic to assume distinct functional roles (niches); and top-down selection, where the host selects for functional redundancy to ensure against the failure of bioreactor functions that could prove highly deleterious (9, 10).

The gene content, genomic arrangement and functional properties of PULs in sequenced gut Bacteroidetes illustrate the specialization and functional redundancy within members of this phylum. They also emphasize how the combined metabolic activities of members of the microbiota undoubtedly result in...
interactions that are both very dynamic and overwhelmingly complex (at least to the human observer), involving multiple potential pathways for the processing of substrates (including the order of substrate processing), varying patterns of physical partitioning of microbes relative to substrates within the ecosystem, plus various schemes for utilization of products of bacterial metabolism. Such a system likely provides multiple options for processing of a given metabolite, and for the types of bacteria that can be involved in these activities.

All of this means that the task of defining the interactions of members of the human gut microbiota is daunting, as is the task of identifying general principles that govern the operation of this system. In the present study, we have taken a reductionist approach to begin to define interactions between members of the Firmicutes and the Bacteroidetes that are commonly represented in the human gut microbiota. In the human colon, Clostridium cluster XIVa is 1 of 2 abundantly represented clusters of Firmicutes. Therefore, we have generated the initial 2 complete genome sequences for members of the genus \textit{Eubacterium} in Clostridium cluster XIVa (the human gut-derived \textit{E. rectale} strain ATCC 33656 and \textit{E. eligens} strain ATCC 27750) and compared them with the draft sequences of 25 other sequenced human gut bacteria belonging to the Firmicutes and the Bacteroidetes. The interactions between \textit{E. rectale} and \textit{B. thetaiotaomicron} were then characterized by performing whole-genome transcriptional profiling of each species after colonization of gnotobiotic mice with each organism alone, or in combination under 3 dietary conditions. Transcriptional data were verified by mass spectrometry of cecal proteins, plus biochemical assays of carbohydrate metabolism. Last, we examined colonization and interactions between these microbes from a host perspective; to do so, we performed whole-genome transcriptional analysis of colonic RNA prepared from mice that were gmo-free or colonized with one or both species. Our results illustrate how members of the dominant gut bacterial phyla are able to adapt their substrate utilization in response to one another and to host dietary changes, and how host physiology can be affected by changes in microbiota composition.

\textbf{Results and Discussion}

\textbf{Comparative Genomic Studies of Human Gut-Associated Firmicutes and Bacteroidetes.} We produced finished genome sequences for \textit{Eubacterium rectale}, which contains a single 3,449,685-bp chromosome encoding 3,627 predicted proteins, and \textit{Eubacterium eligens}, which contains a 2,144,190-bp chromosome specifying 3,627 predicted proteins, and \textit{Parabacteroides distasonis}, plus deep draft assemblies of the \textit{B. caccae}, \textit{B. ovatus}, \textit{B. uniformis}, \textit{B. stercoris} and \textit{P. merdae} genomes generated as part of the human gut microbiome initiative (HGM\textsuperscript{1} (http://genome.wustl.edu/hgm/HGM.frontpage.cgi), and (ii) 16 other human gut Firmicutes where deep draft assemblies were available through the HGM\textsuperscript{1} (see Fig. S1 for a phylogenetic tree). We classified the predicted proteins in these 2 genomes using Gene Ontology (GO) terms generated via Interproscan, and according to the scheme incorporated into the Carbohydrate Active Enzymes (CAZy) database [www.cazy.org (11)], and then applied a binomial test to identify functional categories of genes that are either over- or under-represented between the Firmicutes and Bacteroidetes phyla. This analysis, described in \textit{SI Results}, Figs. S2 and S3, and Table S2 and Table S3, emphasized among other things that the Firmicutes, including \textit{E. rectale} and \textit{E. eligens}, have significantly fewer polysaccharide-degrading enzymes and more ABC transporters and PTS systems than the Bacteroidetes (12). We subsequently chose \textit{E. rectale} and \textit{B. thetaiotaomicron} as representatives of these 2 phyla for further characterization of their niches in vivo, because of their prominence in culture-independent surveys of the distal human gut microbiota (13, 14), the pattern of representation of carbohydrate active enzymes in their glycometabolome and \textit{E. rectale}'s ability to generate butyrate as a major end product of fermentation (15, 16). These choices set the stage for an “arranged marriage” between a Firmicute and a Bacteroidetes, hosted by formerly germ-free mice.

\textbf{Functional Genomic Analyses of the Minimal Human Gut Microbiota. Creating a ‘’minimal human gut microbiota” in gnotobiotic mice.} Young adult male germ-free mice belonging to the NMR1 inbred strain were colonized with \textit{B. thetaiotaomicron} or \textit{E. rectale} alone (monoaosociations) or cocolonized with both species (biosaociation). Ten to fourteen days after inoculation by gavage, both species colonized the cecum of recipient mice, fed a standard chow diet rich in complex plant polysaccharides, to high levels \((n = 4–5\) mice per treatment group in each of 3 independent experiments; Fig. S4A). Moreover, cecal levels of colonization for both organisms were not significantly different between mono- and biosaociated animals (Fig. S4). \textit{B. thetaiotaomicron}’s response to \textit{E. rectale}. A custom, multispecies, human gut microbiome Affymetrix GeneChip was designed (\textit{SI Methods}), and used to compare the transcriptional profile of each bacterial species when it was the sole inhabitant of the cecum, and when it coexisted together with the other species. A significant number of \textit{B. thetaiotaomicron} genes located in PULs exhibited differences in their expression upon \textit{E. rectale} colonization \([55\text{ of } 106; P < 10^{-15}\text{ (cumulative hypergeometric test); see \textit{SI Methods} for the statistical criteria for defining significantly different levels of gene expression}]. Of these 55 genes, 51 (93\%) were up-regulated (Fig. S4B; see Table S4A for a complete list of differentially regulated \textit{B. thetaiotaomicron} genes).

As noted in the Introduction, 2 previous studies from our lab examined changes in \textit{B. thetaiotaomicron}’s transcriptome in the ceca of monosaociated gnotobiotic mice when they were switched from a diet rich in plant polysaccharides to a glucose-sucrose chow (6), or in sucking mice consuming mother’s milk as they transitioned to a standard chow diet (5). In both situations, in the absence of dietary plant polysaccharides, \textit{B. thetaiotaomicron} adaptively forages on host glycans. The genes up-regulated in \textit{B. thetaiotaomicron} upon cocolonization with \textit{E. rectale} have a significant overlap with those noted in these 2 previous datasets \((P < 10^{-14}\text{, cumulative hypergeometric test}; Fig. S4C). In addition, they include several of the genes up-regulated during growth on minimal medium containing porcine mucosal glycans as the sole carbon source (7). For example, in cocolonized mice and in vitro, \textit{B. thetaiotaomicron} up-regulates several genes (BT3787-BT3792; BT3774-BT3777) (Fig. S4D) used in degrading α-mannosidic linkages, a component of host N-glycans and the diet. (Note that \textit{E. rectale} is unable to grow in defined medium containing α-mannan or mannose as the sole carbon sources; Table S3). \textit{B. thetaiotaomicron} also up-regulates expression of its starch utilization system (Sus) PUL in the presence of \textit{E. rectale} (BT3698–3704) (Fig. S4D). This well-characterized PUL is essential for degradation of starch molecules containing \(\geq 6\) glucose units (17).

Thus, it appears that \textit{B. thetaiotaomicron} adapts to the presence of \textit{E. rectale} by up-regulating expression of a variety of PULs so that it can broadcast its niche and degrade an increased variety of glycan substrates, including those derived from the host that \textit{E. rectale} is unable to access. There are a number of reasons why the capacity to access host glycans likely represents an important trait underpinning microbiota function and stability; (i) glycans in the mucus gel are abundant and are a consistently represented source of nutrients; (ii) mucus could serve as a microhabitat for Bacteroidetes spp. to embed in (and adhere to via SusD paralogs), thereby avoiding washout from the
ecosystem; and (iii) the products of polysaccharide digestion/fermentation generated by Bacteroidetes spp. could be shared with other members of the microbiota that are also embedded in mucus (7).

E. rectale’s response to B. thetaiotaomicron. E. rectale’s response to B. thetaiotaomicron in the mouse cecum stands in marked contrast to B. thetaiotaomicron’s response to E. rectale. Carbohydrate metabolism genes, particularly GHs, are significantly overrepresented among the E. rectale genes that are downregulated in the presence of B. thetaiotaomicron compared with monoassociation; i.e., 12 of E. rectale’s predicted 51 GHs have significantly reduced expression while only 2 are up-regulated (Fig. S4 E and F; see Table S4B for a complete list of E. rectale genes regulated by the presence of B. thetaiotaomicron). The 2 up-regulated GH genes (EUBREC.1072, a 6-P-β-glucosidase and EUBREC.3687, a cellobiose phosphorylase) are predicted to break down cellobiose. Three simple sugar transport systems with predicted specificity for cellobiose, galactoside, and arabinose/lactose (EUBREC.3689, EUBREC.0479, and EUBREC.1075–6, respectively) are among the most strongly up-regulated genes (Fig. S4G and Table S4B). Phosphoenolpyruvate carboxykinase (Pck EUBREC.2002) is also induced with cocolonization (Table S4B, GeneChip data were verified by qRT-PCR assays in 2 independent experiments involving 3–4 mice per treatment group; Fig. 1B). This enzyme catalyzes an energy conserving reaction that produces oxaloacetate from phosphoenolpyruvate. In a subsequent transaminase reaction, oxaloacetate can be converted to aspartate, linking this branching of the glycolytic pathway with amino acid biosynthesis (Fig. 1A).

Additional data support the notion that E. rectale is better able to access nutrients in the presence of B. thetaiotaomicron. For example, a number of peptide and amino acid transporters in E. rectale are up-regulated, as are the central carbon and nitrogen regulatory genes CodY (EUBREC.1812), glutamate synthase (EUBREC.1829) and glutamine synthetase (EUBREC.2543) (Fig. 1B and Fig. S4H; note that these genes are also up-regulated during growth in tryptone glucose medium; Table S4C).

Changes in E. rectale’s fermentative pathways. E. rectale possesses genes (EUBEC733–737; EUBEC1017) for the production of butyrate that show high similarity to genes from other Clostridia. This pathway involves condensation of 2 molecules of acetylCoA to form butyrate and is accompanied by oxidation of NADH to NAD⁺ (Fig. 1). Transcriptional and high-resolution proteomic analyses (see below) disclosed that the enzymes involved in
found that the NAD+ in the glycolytic pathway. Consistent with these observations, we observed that 
E. rectale thetaiotaomicron (Fig. 1). E. rectale of these key metabolic genes when turned can be reoxidized via hydrogenases, or via the membrane-

force (Fig. 1). Reduced ferredoxin, in energy conservation, via a bifurcation of electrons from NADH (20). Several observations indicate that E. rectale utilizes B. thetaiotaomicron-derived acetate to generate increased amounts of butyrate in the ceca of our gnotobiotic mice. First, E. rectale up-regulates a phosphate acetyltransferase (EUBREC.1443; EC 2.3.1.8)—1 of 2 enzymes involved in the interconversion of acetyl-CoA and acetate (Fig. 1B). Second, cecal acetate levels are significantly lower in cocolonized mice compared with B. thetaiotaomicron monoassociated animals (Fig. 2). Third, although cecal butyrate levels are similar in E. rectale mono- and biassociated animals (Fig. 2), expression of mouse Mct-1, encoding a monocarboxylate transporter whose inducer and preferred substrate is butyrate (19), is significantly higher in the distal gut of mice containing both E. rectale and B. thetaiotaomicron versus E. rectale alone (P < 0.05; Fig. 2). The cecal concentrations of butyrate we observed are similar to those known to up-regulate Mct-1 in colonic epithelial cell lines (19). Higher levels of acetate (i.e., those encountered in B. thetaiotaomicron monoassociated mice) were insufficient to induce any change in Mct-1 expression compared with germ-free controls (Fig. 2).

The last step in E. rectale’s butyrate production pathway is catalyzed by the butyrylCoA dehydrogenase/electron transfer flavoprotein (Bcd/Etf) complex (EUBREC.0735–0737; EC 1.3.99.2), and offers a recently discovered additional pathway for the interconversion of acetyl-CoA and acetate (Fig. 1). B. thetaiotaomicron encounters M. smithii in the ceca of gnotobiotic mice, there is increased production of acetate by B. thetaiotaomicron, no diversion to butyrate and no induction of Mct-1 (21), increased serum acetate levels, and increased adiposity compared with B. thetaiotaomicron mono-associated controls. In contrast, serum acetate levels and host adiposity (as measured by fat pad to body weight ratios) are not significantly different between B. thetaiotaomicron-rectale cocolonized animals (n = 4–5 animals/group; n = 3 independent experiments; data not shown).

Colonic transcriptional changes evoked by E. rectale-B. thetaiotaomicron cocolonization. We subsequently used Affymetrix Mouse 430 2 GeneChips to compare patterns of gene expression in the proximal colons of mice that were either germ-free, monoassociated with E.rectale or B. thetaiotaomicron, or cocolonized with both organisms (n = 4 mice per group; total of 16 GeneChip datasets). In contrast to the small number of genes whose expression was significantly changed (≥1.5-fold, FDR <1%) after colonization with either bacterium alone relative to germ-free controls (Table S7 A and B), cocolonization produced significant alterations in the expression of 508 host genes (Table S7C). Expression of many of these genes also changed with monoassociation with either organism, and in the same direction as seen after cocolonization, but in most cases the changes evoked by B. thetaiotaomicron or E. rectale alone did not achieve statistical significance. Unsupervised hierarchical clustering of average expression intensity values derived from each of the 4 sets of GeneChips/group, revealed that the E.rectale monoassociation and E.rectale-B. thetaiotaomicron biassociation profiles clustered separate from the germ-free and B. thetaiotaomicron monoassociation datasets (Fig. S5).

Ingenuity Pathway Analysis (www.ingenuity.com) disclosed that the list of 508 host genes affected by cocolonization was significantly enriched in functions related to cellular growth and proliferation (112 genes; Table S8A), and cell death (130 genes) (Table S8B). A number of components of the canonical wnt/b catenin pathway, which is known to be critically involved in controlling self-renewal of the colonic epithelium, were present in this list (Akt3, Axin2, Csnk1D, Dkk3, Frzb, Fzd2, Gja1, Mdm2, Ppp2r5e, Sfrp2, Tgfβ3, Tgfb1, and Tgfb2). Many of the changes observed in biassociated mice are likely to be related to the production of butyrate are among the most highly expressed in cecal contents recovered from mono- and biassociated mice containing E. rectale (Table S4B and Table S6A).

In vitro studies have shown that in the presence of carbohydrates, E. rectale consumes large amounts of acetate for butyrate production (18). Several observations indicate that E. rectale utilizes B. thetaiotaomicron-derived acetate to generate increased amounts of butyrate in the ceca of our gnotobiotic mice. First, E. rectale up-regulates a phosphate acetyltransferase (EUBREC.1443; EC 2.3.1.8)—1 of 2 enzymes involved in the interconversion of acetyl-CoA and acetate (Fig. 1B). Second, cecal acetate levels are significantly lower in cocolonized mice compared with B. thetaiotaomicron monoassociated animals (Fig. 2). Third, although cecal butyrate levels are similar in E. rectale mono- and biassociated animals (Fig. 2), expression of mouse Mct-1, encoding a monocarboxylate transporter whose inducer and preferred substrate is butyrate (19), is significantly higher in the distal gut of mice containing both E. rectale and B. thetaiotaomicron versus E. rectale alone (P < 0.05; Fig. 2). The cecal concentrations of butyrate we observed are similar to those known to up-regulate Mct-1 in colonic epithelial cell lines (19). Higher levels of acetate (i.e., those encountered in B. thetaiotaomicron monoassociated mice) were insufficient to induce any change in Mct-1 expression compared with germ-free controls (Fig. 2).

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The pathway for acetate metabolism observed in this simplified model human gut community composed of B. thetaiotaomicron and E. rectale differs markedly from what is seen in mice that harbor B. thetaiotaomicron and the principal human gut methanogenic archaeon, Methanobrevibacter smithii. When B. thetaiotaomicron encounters M. smithii in the ceca of gnotobiotic mice, there is increased production of acetate by B. thetaiotaomicron, no diversion to butyrate and no induction of Mct-1 (21), increased serum acetate levels, and increased adiposity compared with B. thetaiotaomicron mono-associated controls. In contrast, serum acetate levels and host adiposity (as measured by fat pad to body weight ratios) are not significantly different between B. thetaiotaomicron-rectale cocolonized animals (n = 4–5 animals/group; n = 3 independent experiments; data not shown).

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increased influx of butyrate, generated by *E. rectale*, into colonic cells (Fig. 1A). Butyrate, a histone deacetylase inhibitor that evokes pronounced transcriptional changes in different types of cultured epithelial cell lines (22–25), is the preferred energy substrate for colonic enterocytes (26). While transcriptional changes caused by butyrate differ depending upon the cell lineage, state of cellular differentiation, and cellular energy status (23, 24, 27, 28), in vitro and in vivo studies have shown that it affects expression of genes involved in proliferation, differentiation, and apoptosis (25, 28).

As mentioned above, as part of its adaptation to the presence of *E. rectale*, *B. thetaiotaomicron* up-regulates a number of genes involved in the harvest of host glycans. Included among these *B. thetaiotaomicron* genes are components of a fucose utilization operon linked to production of a bacterial signal that induces synthesis of intestinal mucosal fucosylated glycans, and microbial catabolism of fucose from O-glycans (29). GeneChip profiling of colonic gene expression disclosed that cocolonization results in increased expression of Fut2 (α-1,2 fucosyltransferase), Fut4 (α-1,3-fucosyltransferase), plus 10 other genes involved in the synthesis of mucosal glycans (glycosphingolipids and O-glycans) (Table S8C). Thus, by increasing host production of glycans, *B. thetaiotaomicron* can benefit itself, and through its metabolic products, *E. rectale*.

### E. rectale’s Colonization Levels and Production of Butyrate Are Affected by Host Diet

In a final series of experiments, we assessed how *E. rectale* and *B. thetaiotaomicron* were affected by changes in host diet. Groups of age- and gender-matched cocolonized mice were fed 1 of 3 diets that varied primarily in their carbohydrate and fat content: (i) the standard low-fat, plant polysaccharide-rich diet used for the experiments described above (abbreviated “LF/PP” for low-fat/plant polysaccharide), (ii) a high-fat, “high-sugar” Western-type diet (abbreviated HF/HS) that contained sucrose, maltodextrin, and corn starch, plus complex polysaccharides (primarily cellulose) that were not digestible by *B. thetaiotaomicron* or *E. rectale*, and (iii) a control diet that was similar to (ii) except that the fat content was 4-fold lower (“LF/HS” for low-fat, high-sugar; *n* = 5 mice per group). Whereas *B. thetaiotaomicron*’s colonization levels were similar in all 3 diets, colonization of *E. rectale* was significantly reduced (5-fold) in mice fed either the LF/HS or HF/HS diets (*P < 0.01*, heteroscedastic *t* test).

Whole-genome transcriptional profiling of both bacterial species showed that relative to the standard polysaccharide-rich chow diet (LF/PP), both the Western style HF/HS diet and its LF/HS control produced a significant up-regulation of *B. thetaiotaomicron* PULs involved in harvesting and degrading host polysaccharides, and a downregulation of several PULs involved in the degradation of dietary plant polysaccharides (Fig. S6A). *E. rectale*’s response to the HF/HS and LF/HS diets was to down-regulate several of its GHS and a number of its sugar transporters (Fig. S6B). Moreover, levels of butyrate were 5-fold lower in colonized mice fed these compared with the standard chow (LF/PS) diet ([0.496 ± 0.0051] μmol per gram of wet weight cecal contents; LF/PP vs. 0.095 ± 0.002 (HF/HS) vs. 0.080 ± 0.008 (LF/HS) (*P < 0.05* ANOVA)].

These dietary manipulations lend further support to the view that *B. thetaiotaomicron* with its large repertoire of PUL-associated GHS functions in this model 2-member human microbiota to process complex dietary plant polysaccharides and to distribute to the products of digestion to *E. rectale*, which, in turn, synthesizes butyrate. The reduced colonization response of *E. rectale* to the HF/HS and LF/HS diets can be explained by a number of factors: (i) this Firmicute does not have predicted GHS and PULs that can process host glycans (Fig. S3); (ii) it cannot use most of the sugars we tested that are derived from mucosal polysaccharides (Table S3); and (iii) the host possesses enzymes in its glycometabolism that can directly process the simple sugars present in these 2 diets. Indeed, human subjects that are fed diets deficient in complex polysaccharides harbor lower levels of butyrate-producing gut bacteria, including members of the *E. rectale*-containing clade (30). Our simplified gnotobiotic model of the microbiota underscores the functional implications of diet-associated changes in the representation of this clade, not only as they relate to the operations of the microbiota itself but also potentially as they relate to butyrate-mediated changes in gut epithelial homeostasis.

### Proteomic Studies of This Simplified 2-Component Model of the Human Gut Microbiome

Model communities such as the one described above, constructed in gnotobiotic mice, where microbiome gene content is precisely known and transcriptional data are obtained under conditions where potentially confounding host variables such as diet and host genotype can be constrained, provide a way to test the efficacy of high-resolution mass spectrometric methods for characterizing gut microbial community proteomes. Therefore, we assayed the proteins present in luminal contents, collected from the ceca of 8 gnotobiotic mice fed the standard polysaccharide-rich LF/PP diet (germ-free, monoassociated, and cocolonized; *n* = 2 mice per treatment group representing 2 independent biological experiments; see SI Methods for additional details).

The measured proteomes had high reproducibility in terms of total number of proteins observed and spectra matching to each species. Table S5 and SI Results provide a summary of our analyses, including the percentage of mRNAs called “Present” in the GeneChip datasets for which there was an identified protein product. The most abundant identified products from both microbes included ribosomal proteins, elongation factors, chaperones, and proteins involved in energy metabolism (for a full list of identified proteins, see Table S6; note that Table S4 A and B, which list differentially expressed genes in monoassociation versus bias-association experiments, also indicate whether protein products from their transcripts were identified in these mass spectrometry datasets). Many conserved hypothetical and pure hypothetical proteins were identified, as were proteins encoded by 10 genes in *B. thetaiotaomicron* whose presence had not been predicted in our initial annotation of the finished genome (Table S6A). Together, the results provide validation of experimental and computational procedures used for proteomic assays of a model gut microbiota, and illustrate some of the benefits in obtaining this type of information.

### Prospectus

These studies of a model 2-member human gut microbiota created in gnotobiotic mice support a view of the Bacteroidetes, whose genomes contain a disproportionately large number of glycan-degrading enzymes compared with sequenced Firmicutes, as responding to increasing microbial diversity in the distal intestine by modulating expression of their vast array of polysaccharide utilization loci. *B. thetaiotaomicron* adapts to the presence of *E. rectale* by up-regulating a variety of loci specific for host-derived mucin glycans that *E. rectale* is unable to use. *E. rectale*, which like other Firmicutes has a more specialized capacity for glycan degradation, broadly downregulates its available GHS in the presence of *B. thetaiotaomicron*, even though it does not grow efficiently in the absence of carbohydrates. It also becomes more selective in its harvest of sugars and its transcriptional profile suggests improved access to other nutrients (e.g., there is a generalized up-regulation of amino acid biosynthetic genes and a set of nutrient transporters that can harvest peptides). Thus, this simplified, model microbial community illustrates some of the basic ecologic principles that likely shape the operations of the human gut microbiota: nutrient interchange and the observed reciprocal effects on metabolism of these 2 organisms provide examples of classic syntrophy while “character displacement”, where cooccurrence drives (niche) divergence, also operates.

We have previously used gnotobiotic mice to show that the
efficiency of fermentation of dietary polysaccharides to short chain fatty acids by *B. thetaotaomicron* increases in the presence of *M. smithii* (21). Cocolonization increases the density of colonization of the distal gut by both organisms, increases production of formate and acetate by *B. thetaotaomicron* and allows *M. smithii* to fix CO₂ and produce methane, thereby preventing the build-up of these fermentation end-products (and NADH) in the gut bioreactor, and improving the efficiency of carbohydrate metabolism (21). Removal of H₂ by this methanogenic archaeon allows *B. thetaotaomicron* to regenerate NAD⁺, which can then be used for glycolysis. This situation constitutes a mutualism, in which both members show a clear benefit. The present study, characterizing the cocolonization with *B. thetaotaomicron* and *E. rectale*, describes a more nuanced interaction where both species colonize to similar levels if carbohydrate substrates are readily available. Moreover, certain aspects of bacterial-host mutualism become more apparent with cocolonization, including increased microbial production and host transport of butyrate, and increased host production and microbial consumption of mucosal glycans.

It seems likely that as the complexity of the gut community increases, interactions between *B. thetaotaomicron* and *E. rectale* will either by subsumed or magnified by other “similar” phylogenetic types (as defined by their 16S rRNA sequence and/or by their glycozymes). Synthesizing model human gut microbios of increasing complexity in gnotobiotic mice using sequenced genomes will either by subsumed or magnified by other “similar” phylogenetic types. *E. rectale* is very useful for further testing this idea, as well as a variety of ecologic concepts and principles that may operate to influence the assembly and dynamic operations of our gut microbial communities.

**Materials and Methods**

**Genome Comparisons.** All nucleotide sequences from all contigs of completed genome assemblies containing both capillary sequencing and pyrosequencer data, produced as part of the HGMI, were downloaded from the Washington University Genome Sequencing Center’s website (http://genome.wustl.edu/pub/organism/Microbes/Human.Gut.Microbiome) on September 27, 2007.

The finished genome sequences of *B. thetataomicron* VPI-5482, *Bacteroides vulgatus* ATCC 8482, and *B. fragilis* NCTC3343 were obtained from GenBank.

For comparison purposes, protein-coding genes were identified in all genomes using YACOP (32). Each proteome was assigned InterPro numbers and GO terms using InterProScan release 16.1. Statistical comparisons between genomes were carried out as described in ref. 4, using perl scripts that are available upon request from the authors.

**GeneChip Analysis.** Previously described methods were used to isolate RNA from a 100- to 300-mg aliquot of frozen fecal contents, synthesize cDNA, and to biotinylate and hybridize the cDNAs to a custom bacterial GeneChip (21). The only modification was that in RNA isolation protocol, 0.1 mm zirconia/silica beads (Biospec Products) were used for lysis of bacterial cells in a bead mill (60-s min run at highest speed). Genes in given bacterial species that were differentially expressed in mono-versus biassociation experiments were identified using CyberT (default parameters) after probe masking and scaling with the MAS5 algorithm (Affymetrix; for details about the methods used to create the mask, see the Methods section of SI Text).

RNA was purified from proximal colon using Mini RNeasy kit (Qiagen) with on-column DNase digestion. Biotinylated cRNA targets were prepared from each sample (n = 4 per treatment group). cRNA was hybridized to Affymetrix Mouse Genome Mo430 2 GeneChips, and the resulting datasets analyzed using Probe Logarithmic Error Intensity Estimate method (PLIER + 16). Fold-changes and p-values were calculated using Cyber-T. Significance was defined by maintaining a FDR <1% using Benjamini–Hochberg correction (33).

**Other Methods.** Details about bacterial culture, genome sequencing and finishing, animal husbandry, quantitative PCR assays of the level of colonization of the ceca of gnotobiotic mice, GeneChip design and masking, plus proteomic and metabolite assays of cecal contents are provided in SI Methods.

**ACKNOWLEDGMENTS.** We thank Maria Karlsson and David O’Donnell for help with gnotobiotic husbandry; Jan Crowley, Janaki Guruge, Jill Manches- ter, and Sabrina Wagoner for technical assistance; Ruth Ley for valuable comments during the course of this work; and Marne Shaw for proteomics data-mining and computational aspects related to mass spectrometry. This work was supported by National Science Foundation Grant 0332384 and National Institutes of Health Grants DK30922, DK70977, DK52574, GM07200, and T32-AI07172 and by the Laboratory Directed Research Program Oak Ridge National Laboratory.

Supporting Information

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SI Methods

Bacterial Culture. Bacterial strains were stored frozen at −80 °C in a prerduced mixture of 2 parts TYG medium (1) to 1 part glycerol. Bacteria were routinely cultured in TYG medium in an anaerobic chamber (Coy Lab Products, Grass Lake, MI) under an atmosphere of 40% CO2, 5% nitrogen, and 2% H2. To assay growth of _E. rectale_ on specific carbon sources, the organism was cultured on medium containing 1% tryptone, 100 mM potassium phosphate buffer (pH 7.2), 15 mM NaCl, 180 μM CaCl2, 100 μM MgCl2, 50 μM MnCl2, 42 μM CoCl2, 15 μM FeSO4, 1% trace element mix (ATCC), 2 μg/mL folinic acid (calcium salt), 1.2 μg/mL hematin, and 1 mg/mL menadione. Growth curves for different carbon sources were acquired at 37 °C in the Coy anaerobic chamber using a 96-well plate spectrophotometer (Tecan). Growth was scored as positive if the OD600 measurement rose by ≥0.2 over a 72-h incubation at 37 °C.

Genome Sequencing. _E. rectale_ and _E. eligens_ were grown to late log phase under anaerobic conditions in TYG medium. Cells harvested from a 50 mL culture were lysed by incubation for 30 min at 37 °C in 11 mL of Buffer B1 (Qiagen Genomic DNA buffer set; Qiagen) supplemented with 2.2 mg of RNase A, 50 units lysozyme, 50 units mutanolysin, and 600 units achromopeptidase (all from Sigma) followed by addition of 4 mL of Buffer B2 (Qiagen) together with 10 mg (300 units) proteinase K (Sigma) and incubation at 50 °C for 2 h. DNA was precipitated by adding 1.5 mL of 3M sodium acetate and 30 mL of isopropanol, removed with a sterile glass hook, and washed several times with ethanol.

Unlike _E. eligens_, genomic DNA from _E. rectale_ was very resistant to standard cloning techniques. This cloning bias made efforts to produce fosmids ineffective, and left vast regions of the genome uncloned in our primary sequencing vector, pOT. Only half (1.7 Mb) of its genome was represented in our initial assembly containing 228 contigs from >9x plasmid shotgun reads with an ABI 3730xl capillary instrument. Therefore, we generated >40x coverage of the _E. rectale_ genome through pyrosequencing with a 454 GS20 instrument, and used an additional vector (pJAZZ) for capillary sequencing to obtain a finished genome sequence.

Protein-coding genes were identified with Glimmer 2.13 (2) and GeneMarkS (3), using the start site predicted by GeneMarkS where the two overlapped. “Missed” genes were then added by using a translated BLAST of intergenic regions against the nonredundant (NR) protein database and finding conserved ORFs. Additional missed genes were added to the _E. rectale_ genome using YACOP (trained by Glimmer 2.13) (4), tRNA, rRNA and other non-coding RNAs were identified and annotated using tRNAscan-SE (5), RNAMMER (6), and RFAM (7), respectively. Protein-coding genes were annotated with the KEGG Orthology group definition using a National Center for Biotechnology Information BLASTP search (8) of the KEGG genomes database (9), with a minimum bit score of 60.

Animal Husbandry. All experiments using mice were performed using protocols approved by the animal studies committee of Washington University. NMRI-KI mice (10) were maintained in flexible plastic film isolators under a strict 12 h light cycle, and fed an irradiated standard low-fat, high plant polysaccharide chow (LF/PP, diet 2018 from Harlan Teklad, www.tekladcus-tomdiets.com) or a high fat, high-sugar (HF/HS) Western-style diet (Harlan Teklad 96132) or a corresponding control low fat, high-sugar (LF/HS; Harlan Teklad 03317).

Animals were colonized via gavage with 10⁸ CFU from an overnight culture of _B. thetaiotaomicron_ or a log-phase culture of _E. rectale_. Gavage with _E. rectale_ was repeated on 3 successive days using cells from separate log-phase cultures begun from separate colonies. Cecal contents were flash frozen in liquid nitrogen immediately after animals were killed.

Quantitative (q) PCR Measurements of Colonization. A total of 100–300 mg of frozen cecal contents from each gnotobiotic mouse was added to 2 mL tubes containing 250 μL of 0.1 mm-diameter zirconia/silica beads (Biospec Products). 0.5 mL of Buffer A (200 mM NaCl, 20 mM EDTA), 210 μL of 20% SDS, and 0.5 mL of a mixture of phenol:chloroform:isoamyl alcohol (25:24:1; pH 7.9; Ambion). Samples were lysed with a bead beater (BioSpec; “high” setting for 4 min at room temperature). The aqueous phase was extracted after centrifugation (8,000 × g at 4 °C for 5 min), and the extraction repeated with another 0.5 mL of phenol:chloroform:isoamyl alcohol and 1 min of vortexing. DNA was precipitated with 0.1 volume of 3M sodium acetate (pH 5) and 1 volume of isopropanol (on ice for 20 min), pelleted (14,000 × g, 20 min at 4 °C) and washed with ethanol. The resulting pellet was resuspended in water and 1/2 (for _E. rectale_ mono-associations) or 1/10 of the DNA (for _B. thetaiotaomicron_-colonized samples) cleaned up further using a DANEasy column (Qiagen).

qPCR was performed using (i) primers specific to the 16S rRNA gene of _B. thetaiotaomicron_ (11) or to the _Clostridium coccosides_/E. rectale group (forward: 5'-CGGTACCTGAAGAAAAGACC-3'; reverse: 5'-AGTTT(C/T)ATTCTTGCAACG-3') (12), and (ii) conditions described previously for _B. thetaiotaomicron_ (11). The amount of DNA from each genome in each PCR was computed by comparison to a standard curve of genomic DNA prepared in the same manner from pure cultures of each bacterial species. Data were converted to genome equivalents by calculating the mass of each finished genome (2.8 × 10⁶ genome equivalents (GEq) per ng _E. rectale_ DNA, and 1.5 × 10⁶ GEq per ng _B. thetaiotaomicron_ DNA).

GeneChip Design, Hybridization, and Data Analysis. A custom, 6-species human gut microbiome Affymetrix GeneChip was designed using the finished genome sequences of _B. thetaiotaomicron_, _B. vulgatus_, _P. distasonis_ and _M. smithii_ (13–15), plus draft versions of the _E. rectale_ and _E. eligens_ genomes. GeneChip predictions for the Firmicute assemblies were made using Glimmer3 (2). The design included 14 probe pairs (perfect match plus mismatch) per CDS (protein coding sequence) in each draft assembly, and 11 probe pairs for each CDS in a finished genome.

Non-specific cross-hybridization was controlled in 2 ways. _First_, probe masks for each genome were developed as follows. For analysis of _E. rectale_- _B. thetaiotaomicron_ cocolonizations, probes resulting from misassembly and missing sequences later identified (from the finished genome) in the _E. rectale_ draft assembly were removed to avoid cross-hybridization. _Second_, a custom, 6-spe- cies human gut microbiome Affymetrix GeneChip was designed using the finished genome sequences of _B. thetaiotaomicron_, _B. vulgatus_, _P. distasonis_, and _M. smithii_ (13–15), plus draft versions of the _E. rectale_ and _E. eligens_ genomes. GeneChip predictions for the Firmicute assemblies were made using Glimmer3 (2). The design included 14 probe pairs (perfect match plus mismatch) per CDS (protein coding sequence) in each draft assembly, and 11 probe pairs for each CDS in a finished genome.

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Primers used for these reactions are available from the authors computed using Bioconductor. Expression values were calculated using Bioconductor.

Expression of selected genes was confirmed by qRT-PCR (11). Primers used for these reactions are available from the authors on request.

Proteomic Analyses of Cecal Contents. Cecal contents were processed via a single tube cell lysis and protein digestion method as follows. Briefly, the cell pellet was resuspended in 6 M Guanidine/10 mM DTT, heated at 60 °C for 1 h, followed by an overnight incubation at 37 °C to lyse cells and denature proteins. The guanidine concentration was diluted to 1 M with 50 mM Tris/10 mM CaCl2 (pH 7.8), and sequencing grade trypsin (Promega) was added (1:100; wt/wt). Digestions were run overnight at 37 °C. Fresh trypsin was then added followed by additional 4 h incubation at 37 °C. The complex peptide solution was subsequently de-salted (Sep-Pak C18 solid phase extraction; Waters), concentrated, filtered, aliquoted and frozen at −80 °C.

All 8 samples were coded and mass spectrometry measurements conducted in a blinded fashion. Cecal samples were analyzed in technical triplicates using a 2-dimensional (2D) nano-LC MS/MS system with a split-phase column (SCX-RP) (18) on a linear ion trap (Thermo Fisher Scientific) with each sample consuming a 22-h run as detailed elsewhere (19, 20). The linear ion trap (LTQ) settings were as follows: dynamic exclusion set at 1; and 5 data-dependent MS/MS. Two microscans were averaged for both full and MS/MS scans and centroid data were collected for all scans. All MS/MS spectra were searched with the SEQUEST algorithm (21) against a database containing the entire mouse genome, plus the B. thetaiotaomicron, and E. rectale, genomes (common contaminants such as keratin and trypsin were also included). To find potential food proteins, yeast and rice databases were included. The break down of each database can be found and downloaded from http://compbio.ornl.gov/gnotobiotic/mouse_cecal.metaproteome/databases/. The SEQUEST settings were as follows: enzyme type, trypsin; Parent Mass Tolerance, 3.0; Fragment Ion Tolerance, 0.5; up to 4 missed cleavages allowed (internal lysine and arginine residues), and fully tryptic peptides only (i.e., both ends of the peptide must have arisen from a trypsin-specific cut, except the N and C termini of proteins). All datasets were filtered at the individual run level with DTASelect (22) [Xcorrs of at least 1.8 (+1 ions), 2.5 (+2 ions) 3.5 (+3 ions)]. Only proteins identified with 2 fully tryptic peptides were considered. All resulting DTASelect Contrast files used in this study are available from http://compbio.ornl.gov/gnotobiotic/mouse_cecal.metaproteome. Also accessible from this site are MS/MS spectra for all identified peptides.

For this study, false-positive rates (FPR) were used to estimate the error associated with peptide identifications. The overall FPR was estimated using the formula: FPR = \( \frac{2(n_{\text{rev}}/\text{N}_{\text{rev}} + n_{\text{real}})}{\text{N}_{\text{real}}} \times 100 \) where \( n_{\text{rev}} \) is the number of peptides identified from the reverse database and \( n_{\text{real}} \) is the number of peptides identified from the real database (23). Reverse and shuffled databases were created to calculate FPRs (23, 24). A reverse database was created by precisely reversing each protein entry (i.e., N terminus became C terminus in each case) and then appending these reversed sequences onto the original database. Two runs—samples 705, Run 1 and 710, Run 2—were randomly selected for estimating a FPR. The observed FPR rates were 0.55% and 0.31% respectively for these 2 runs. An additional database was created by randomly shuffling the amino acids of each protein rather than simply reversing the N terminus and C terminus. A FPR was estimated using a similar formula as that described above except that the number of identified reverse peptides was replaced with the number of shuffled peptides. A FPR was estimated for both samples, 705, Run 1 (0.45%) and 710, Run2 (0.31%) and was similar to the rate determined by the reverse database method. Datasets for calculating FPR rates are available on the web site mentioned above.

In addition to differentiating between true and false peptide identifications with FPRs, label-free quantitation methods were used to estimate relative protein abundance. Several protein quantitation methods are currently available and routinely performed for shotgun proteomics. To estimate relative protein abundance in complex protein mixtures and communities, spectral counts and normalized spectral abundance factors (NSAF) (25) are commonly used. Spectral counting is based on the theory that the more abundant peptides are typically sampled more frequently, resulting in higher spectral counts. Lui et al. (26) has shown that spectral copy number provides a more accurate correlation to protein abundance than peptide count and % coverage. NSAF, however, is based on spectral counts, but takes into account protein size and the total number of spectra from a run, thus normalizing the relative protein abundance between samples (25).

Biochemical Analyses. Measurements of acetate, propionate, butyrate, NAD+, and NADH in cecal contents were performed as described in ref. 11, with the exception that acetate acid-1-13C4 (Sigma) was used as a standard to control for acetate recovery.

SI Results

Comparative Genomic Studies of Human Gut-Associated Firmicutes and Bacteroidetes. Although the sequenced gut Bacteroidetes all harbor large sets of polysaccharide sensing, acquisition and degradation genes, the gut Firmicutes, including E. rectale and E. eligens, have smaller genomes and a significantly smaller proportion of genes involved in glycan degradation (Fig. S2). The gut-associated Bacteroidetes possess large families of SusC and SusD paralogs involved in binding and import of glycans, while the genomes of E. rectale and other gut Firmicutes are enriched for phosphotransferase systems and ABC transporters (Fig. S2).

Lacking adhesive organelles, the ability of gut Bacteroidetes to attach to nutrient platforms consisting of small food particles and host mucus via glycansusC/SusD outer membrane binding proteins likely increases the efficiency of oligo- and monosaccharide harvest by adaptively expressed bacterial GHs, and preventing washout from the perfused gut bioreactor. Unlike the surveyed Bacteroidetes, several Firmicutes, notably E. rectale, E. eligens, E. siraeum, and Anaerotruncus colihominis (the later belongs to the Clostridium leptum cluster) possess genes specifying components of flagellae (Fig. S2). These organelles may contribute to persistence within the continuously perfused gut ecosystem and/or enable these species to move to different microhabitats to access their preferred nutrient substrates.

Table S2 lists predicted GHs and PLs present in the Firmicutes and Bacteroidetes surveyed, sorted into families according to the scheme incorporated into the Carbohydrate Enzymes (CAZy)
results of in vitro assays of the growth of family 28, PL families 1 and 9) (Table S2). Among the Bacteroidetes “glycobiomes”, there is also evidence of niche specialization: Although B. vulgatus has fewer GHS and PLs overall than B. thetaiotaomicron, it has a larger assortment of enzymes for degrading pectins (e.g., GH family 28 and PL families 1, 10 and 11) and possesses a number of enzymes for degrading pectins (e.g., GH and possesses enzymes that degrade pectins (GH family 28 and PL families 1 and 9) (Table S2). Among the Bacteroidetes “glycobiomes”, there is also evidence of niche specialization: Although B. vulgatus has fewer GHS and PLs overall than B. thetaiotaomicron, it has a larger assortment of enzymes for degrading pectins (e.g., GH family 28 and PL families 1, 10 and 11) and possesses enzymes that B. thetaiotaomicron lacks that should enable it to degrade certain xylans (GH family 10 and Carbohydrate esterase (CE) family 15] (Fig. S3 and Table S2). The results of in vitro assays of the growth of B. thetaiotaomicron and E. rectale in defined medium containing mono- di- and polysaccharides are summarized in Table S3.

Proteomic Analysis. For a complete list of the total number of identified spectra, peptides and proteins per sample and run, see Table S6B. Interestingly, the total number of identified spectra were, for the most part, distinct and unique to each bacterial species. Unlike B. thetaiotaomicron and E. rectale, the number of identified spectra belonging to mouse were redundant; Thus, a higher number of spectra were non-unique spectra. The difference is evident when the total spectra counts are compared with unique spectra counts only. The total average spectra count identified in the control (germ-free) mouse was 10,767 for sample 700 and 11,221 for sample 799. The total average unique spectra count, however, decreased to 4,394 and 4,168. Therefore, the majority of mouse peptides are not unique within the database.

The total number of unique spectra counts per species and run can be found in Table S6C. The 2 cocolonized mice (710 and 810) had a total of ~77% unique spectra belonging to B. thetaiotaomicron, 20% unique spectra belong to E. rectale, and only 3% of the 2 species’ combined spectra counts were non-unique. This suggests that the majority of identified proteins belonging to B. thetaiotaomicron and E. rectale are true unique identifications and these species can be easily differentiated by proteomics.
Fig. S1. Phylogenic relationships of human gut-associated Firmicutes and Bacteroidetes surveyed in the present study. A phylogeny based on 16S rRNA gene sequences showing the relationships between representatives from 2 dominant bacterial phyla in the gut microbiota. Green, Firmicutes; red, Bacteroidetes; arrows, organisms used for cocolonization studies described in the present study. 16S rRNA gene sequences were aligned with the NAST aligner (27). Likelihood parameters were determined using Modeltest (28) and a maximum-likelihood tree was generated using PAUP (www.paup.csi.fsu.edu). Bootstrap values represent nodes found in >70 of 100 repetitions.
Fig. S2. Genes involved in carbohydrate metabolism and energy production whose representation is significantly enriched or depleted in sequenced human gut-associated Firmicutes and Bacteroidetes. The number of genes assigned to each GO term in each genome is shown. Significance is judged by a binomial test, with multiple hypothesis testing correction (see Methods) comparing the proportion of genes assigned to a GO term in one genome versus the average number assigned to the same GO term across all of the Firmicutes. The Firmicutes also use distinct mechanisms for environmental sensing and membrane transport. The Bacteroidetes employ a large number of paralogs of the SusC/D system to bind and import sugars (classified as receptors, GO:0004872), while the Firmicutes use ABC transporters and phosphotransferase systems (classified as active membrane transporters, GO:0022804). Red, enriched; blue, depleted; darkercolor $P < 0.001$ and light color $P < 0.05$ relative to the average of all Firmicute genomes excluding the one tested.
### CAZy Family

<table>
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<tr>
<th>CAZy Family</th>
<th>B. thetaiotaomicron</th>
<th>B. vulgatus</th>
<th>E. rectale</th>
<th>E. eligens</th>
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**Fig. S3.** Comparison of glycoside hydrolases and polysaccharide lyases repertoires of E. rectale, E. eligens, B. vulgatus and B. thetaiotaomicron. The number of genes in each genome in each CAZy GH or PL family are shown. Families that are significantly depleted relative to B. thetaiotaomicron are colored blue (P < 0.001), as judged by a binomial test followed by Benjamini-Hochberg correction. Families for which B. thetaiotaomicron has significantly more members are colored yellow. Families that are absent in B. thetaiotaomicron are highlighted in orange. B. thetaiotaomicron has a larger genome and a disproportionately larger assortment of GHs. Both Firmicutes have a reduced capacity to use host-derived glycans (hexosaminidases, mannosidases, and fucosidases; GH20, GH29, GH76). E. rectale has a large number of starch-degrading enzymes (GH13), while E. eligens has a capacity to degrade pectins (PL9, GH28). See Table S2 for a complete list of all CAZy enzymes among the sequenced gut Bacteroidetes and Firmicutes examined.
Fig. S4. Creation of a minimal human gut microbiota composed of a sequenced Firmicute (E. rectale) and a sequenced Bacteroidetes (B. thetaiotaomicron).
(A) Levels of colonization of the ceca of 11-week-old male gnotobiotic mice colonized for 14 days with one or both organisms. Animals were given an irradiated low-fat plant polysaccharide-rich (LF/PP) chow diet ad libitum. B. thetaiotaomicron and E. rectale colonize the ceca of mice to similar levels in both mono- and biassociation. Error bars denote standard error of the mean of 2–3 measurements per mouse, 4 mice per group. Results are representative of 3 independent experiments. (B) Summary of genes showing up-regulation in B. thetaiotaomicron with cocolonization. 55 of the 106 genes are within PULs, and of these, 51 (93%) were up-regulated. (C) Summary of B. thetaiotaomicron PUL-associated genes up-regulated with cocolonization and their representation in datasets of genes up-regulated during the suckling-weaning transition (29), and when adult gnotobiotic mice are switched from a polysaccharide-rich diet to one devoid of complex glycans and containing simple sugars (glucose, sucrose (30)). The latter 2 datasets are composed of genes that are also up-regulated 10-fold relative to log-phase growth in minimal glucose medium (30). (D) Heat map of GeneChip data from 3 loci up-regulated by E. rectale upon colonization with E. rectale; 2 are involved in degradation of α-mannans that E. rectale cannot access; the third is the Starch utilization system (Sus) locus, which targets a substrate that both species can use. Maximal relative expression across a row is red; minimal is green. Differential expression was judged using the MAS5 algorithm and CyberT (see Table S4A and Methods). (E) Overview of the response of E. rectale to cocolonization with B. thetaiotaomicron. Genes assigned to GO terms for carbohydrate metabolism (GO:0005975), transporters (GO:0006810) and predicted GHs are all significantly over-represented among down-regulated genes while genes with GO terms for biosynthesis (GO:0044249), in particular amino acid metabolism (GO:0006520), are significantly over-represented among up-regulated genes. All categories shown are significantly different from the genome as a whole. *, P < 0.05; **, P < 0.01; ****, P < 0.001 (binomial test). (F) Heat map from GeneChip data of all significantly regulated E. rectale GH genes showing that all but 2 are downregulated (both cytoplasmic phosphosugar processing enzymes) n = 4 mice/treatment group. (G) Heat map of all significantly regulated E. rectale genes assigned to the GO term for transporters (GO:0006810) showing that a number of simple sugar transporters are downregulated upon cocolonization, while peptide and amino acid transporters and 3 predicted simple sugar transporters (arrows; EUBREC.08479, a galactose ABC transporter; EUBREC.1075–6, a lactose/arabinose transport system; and EUBREC.3689, a predicted cellobiose transporter) are up-regulated. (H) Heat map of selected global regulators from E. rectale shows that CodY, a repressor in other Firmicutes of stationary-phase genes such as those needed to access lower-energy carbon sources, is significantly up-regulated upon cocolonization, suggesting increased accessibility of nitrogen and/or carbon sources. Glutamine synthetase and glutamate synthase, are also up-regulated, consistent with the observed up-regulation of various amino acid and peptide transporters. Differentially regulated genes were identified using the MA55 algorithm and Cyber-T (see Table S4B and Methods). Genes whose differential expression with cocolonization was further validated by qRT-PCR are highlighted with pink lettering (2 independent experiments, n = 4–5 mice per group, 2–3 measurements per gene; see Fig. 1).
Fig. S5. Unsupervised hierarchical clustering of average GeneChip expression intensity values from probesets representing differentially expressed genes (>1.5-fold; <1% FDR) in the proximal colons of germ-free versus cocolonized animals. Clustering was performed using host expression data from all treatment groups: germ-free, B. thetaiotaomicron and E. rectale monoassociations, and B. thetaiotaomicron-E. rectale biassociation. Data from all GeneChips in a given treatment group were averaged (n = 4 animals/group; total of 694 probesets analyzed). Clustering was performed by cluster 3.0 software (www.bonsai.ims.u-tokyo.ac.jp/~mdehoon/software/cluster/software.htm#ctv) and data were visualized using Tree View software. Each probeset is represented by a single row of colored bars. Maximal relative expression across a row is red; minimal is green.
hypothetical proteins were not included in the heat map. Differential expression was judged using the MAS5 algorithm and Cyber-t. Maximal relative expression

Fig. S6. Effects of low carbohydrate content diets on a B. thetaiotaomicron–E. rectale community gene expression. (A) Heat map of GeneChip data from B. thetaiotaomicron in the ceca of mice (cocolonized with E. rectale; n = 4 mice per group) that were either fed (i) a standard low-fat plant polysaccharide-rich diet (LFPP), (ii) a high-fat, high sugar (HFHS) “Western style” diet containing cellulose as the only complex plant polysaccharide or (iii) a control for diet (iii) that contained 4-fold less fat, high levels of simple sugars plus cellulose (LFHS). Polysaccharide utilization loci (PULs) whose specificities are known are indicated (31). Genes predicted to be regulated by complex plant polysaccharides are also highlighted. (B) Heat map of GeneChip data from E. rectale in the ceca of mice cocolonized with B. thetaiotaomicron fed as described in A. All genes were defined as significantly differentially expressed (>2 fold, PPDE >0.95). Genes encoding hypothetical proteins were not included in the heat map. Differential expression was judged using the MASS algorithm and Cyber-t. Maximal relative expression across a row is red; minimal is green.
Other Supporting Information Files

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