

***Bacteroides* in the Infant Gut Consume Milk Oligosaccharides via Mucus-Utilization Pathways**

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SUMMARY

Newborns are colonized with an intestinal microbiota shortly after birth, but the factors governing the retention and abundance of specific microbial lineages are unknown. Nursing infants consume human milk oligosaccharides (HMOs) that pass undigested to the distal gut, where they may be digested by microbes. We determined that the prominent neonate gut residents, *Bacteroides thetaiotaomicron* and *Bacteroides fragilis*, induce the same genes during HMO consumption that are used to harvest host mucus glycans, which are structurally similar to HMOs. Lacto-*N*-neotetraose, a specific HMO component, selects for HMO-adapted species such as *Bifidobacterium infantis*, which cannot use mucus, and provides a selective advantage to *B. infantis* in vivo when biassociated with *B. thetaiotaomicron* in the gnotobiotic mouse gut. This indicates that the complex oligosaccharide mixture within HMOs attracts both mutualistic mucus-adapted species and HMO-adapted bifidobacteria to the infant intestine that likely facilitate both milk and future solid food digestion.

INTRODUCTION

The infant gut undergoes a complex and unpredictable process of colonization during the first months of life, characterized by extreme fluctuations in overall density and membership (Koening et al., 2011; Palmer et al., 2007). Factors that drive microbiota assembly remain poorly understood. Several studies indicate that breast-milk versus formula feeding can play a large role in infant intestinal microbiota composition (Adlerberth and Wold, 2009). Human milk oligosaccharides (HMOs), a class of carbohydrates within human milk, may promote intestinal colonization of specific subsets of microbes (Engfer et al., 2000). HMOs are composed of more than 200 structurally distinct oligosaccha-

rides that occur at high concentrations (~20 g/L in colostrums; ~5–12 g/L in mature milk). The wide diversity of structures includes linear or branched lactosamine chains (Gal β 1-3/4, GlcNAc β 1-3/6) extended from a lactose (Gal β 1-4Glc) core. Structural variability is due to the addition of the terminating sugars *N*-acetylneuraminic acid (Neu5Ac) in α 2-3 or α 2-6 linkages and/or fucose in α 1-2, α 1-3, or α 1-4 linkages (reviewed in Chichlowski et al., 2011). The complete degradation of HMOs requires an extensive set of glycoside hydrolases and/or membrane transporters that humans lack in the small intestine. In the lower part of the intestine, HMOs theoretically can be used by members of the infant intestinal microbiota and thereby promote HMO-utilizing microbes. The ability of certain *Bifidobacterium* species to consume HMOs has been demonstrated and genome sequencing of HMO-utilizing *Bifidobacterium* isolates has revealed genes involved in HMO uptake, degradation, and metabolism (Sela et al., 2008; LoCasio et al., 2007). Recently, it has been described that HMO utilization is not exclusive to certain *Bifidobacterium*, since members from the genus *Bacteroides* can use milk glycans as a sole carbon source (Marcobal et al., 2010).

Bacteroides species are variably abundant in the infant gastrointestinal microbiota, but by the first year of life they are consistently present in the gut (Palmer et al., 2007). These species are well adapted to use a multitude of both dietary polysaccharides and host-derived glycans (e.g., mucus), due to specialized machinery encoded by polysaccharide utilization loci (PULs) (Martens et al., 2008). PULs have been widely expanded within the genomes of *Bacteroides*, and each appears to be specialized in the use of a particular class of carbohydrates (Martens et al., 2008; Sonnenburg et al., 2010). The ability of *Bacteroides* to utilize HMOs suggests that specific PULs within these *Bacteroides* genomes are involved in HMO use.

Using *Bacteroides thetaiotaomicron* (*Bt*) and *Bacteroides fragilis* (*Bf*), two of the predominant *Bacteroides* species found in the gut of neonates (Rotimi and Duerden, 1981), we have demonstrated that each species induces a common set of genes when growing in host mucin glycans or HMOs. However, these genes are not conserved between *Bt* and *Bf*. These data suggest that HMO use is either a convergent functionality between members of the genus or relies upon pathways that have been

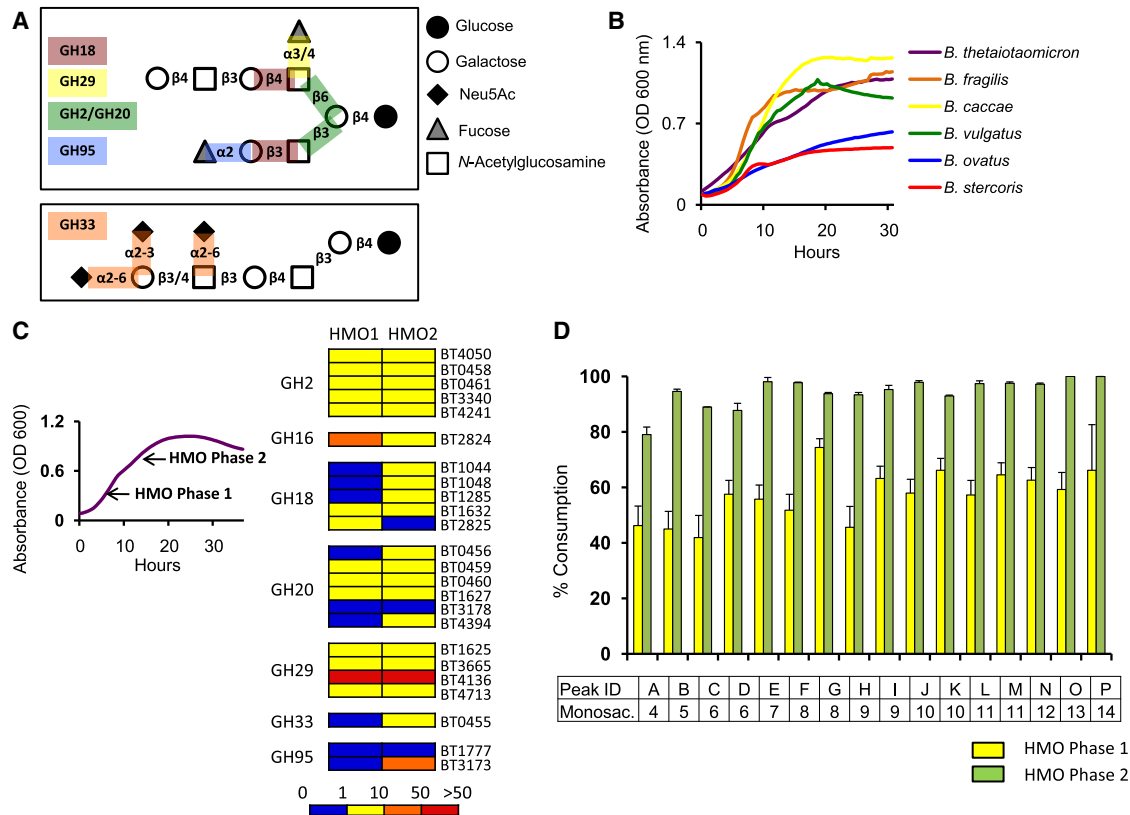


Figure 1. *B. thetaiotaomicron* Upregulates Numerous Glycoside Hydrolases during Consumption of HMO

(A) Schematic of HMO linkages (branched, top box; linear, bottom box) and putative HMO active glycoside hydrolase families (GH) (Wu et al., 2010). Linkages are to the 1-carbon of the underlying sugar unless otherwise noted.

(B) In vitro growth of *Bacteroides* species in MM-HMO.

(C) *Bt* gene expression at two time points (HMO1, HMO2) in MM-HMO relative to MM-glucose for 24 putative HMO active GHs (predicted to hydrolyze linkages found in milk glycans in A). See Table S1 for full list of genes upregulated in all tested carbon sources.

(D) *Bt* HMO consumption at two time points (HMO1, HMO2), determined by MALDI-FTICR-MS. Peak IDs correspond to a characteristic oligosaccharide, with the following discrete mass to charge ratios (m/z): A, 732.25; B, 878.31; C, 1024.36; D, 1097.38; E, 1243.44; F, 1389.50; G, 1462.51; H, 1535.55; I, 1608.57; J, 1754.63; K, 1827.64; L, 1900.69; M, 1973.70; N, 2119.76; O, 2265.82; P, 2484.89. Number of monosaccharides for each mass is indicated. See Figure S1 for detailed structural information for each oligosaccharide. Error bars represent standard deviation for three biological replicates.

differentially shaped by other selective forces, such as mucus utilization. The ability of the short HMO, lacto-*N*-neotetraose (LNnT), to selectively expand the HMO-using *Bifidobacterium infantis* (*Bi*) relative to *Bt* in gnotobiotic biassociated mice suggests that milk oligosaccharides play a dual role in selecting for species that are HMO or mucus adapted.

RESULTS

B. thetaiotaomicron Exhibits an Expansive Glycoside Hydrolase Response during Consumption of HMOs In Vitro

Bt possesses a repertoire of predicted glycoside hydrolases (GH) capable of accommodating the structural diversity found in milk oligosaccharides. Among *Bt*'s > 260 glycoside hydrolases, 67 make up six GH families with predicted activities required to process linkages found in HMOs (Figure 1A). *Bt* grows efficiently when cultured in minimal medium containing HMO (1.5% w/v; MM-HMO) as the sole carbon source. Five

additional sequenced *Bacteroides* species, *Bf*, *B. caccae*, *B. vulgatus*, *B. ovatus*, and *B. stercoris*, all grow in the presence of HMOs. Growth of *Bf*, *B. vulgatus*, and *B. caccae* are comparable to that of *Bt* (doubling times of 2.9 hr, 3.3 hr, and 2.8 hr, respectively; saturating OD₆₀₀ > 0.9 for each strain). *B. ovatus* and *B. stercoris* do not exhibit exponential growth in MM-HMO, indicating that efficient use of milk oligosaccharides is not universal in the gut resident *Bacteroides* (Figure 1B).

We identified the *Bt* genes induced by HMOs using transcriptional profiling of *Bt* during growth in MM-HMO at the midpoints of the two logarithmic growth phases, using a *Bt* GeneChip (n = 2 biological replicates/growth phase, four datasets total) (Figure 1C). As a baseline for comparison, we used previously reported data of *Bt* grown in MM-glucose (Martens et al., 2008). A total of 156 genes are significantly upregulated during the first phase, and 230 genes are upregulated during the second phase, relative to MM-glucose (see Experimental Procedures for criteria used to define significance; see Table S1 for complete list). Forty-six genes of the 253-gene response to HMOs are

predicted to encode glycoside hydrolases, and over half of those (24 genes) belong to the seven GH families that target the most common linkages found within HMOs (Figures 1A and 1C).

The biphasic growth of *Bt* in MM-HMO suggests a sequential, ordered degradation of glycans. We used laser desorption/ionization coupled with mass spectrometry to characterize the consumption of 16 structurally defined neutral milk oligosaccharides, which represents >85% of the total HMO pool (LoCascio et al., 2007). After the completion of each exponential phase, HMOs were purified from culture supernatants, reduced, and profiled (Marcobal et al., 2010). During the first growth phase, *Bt* consumes the full spectrum of HMOs; a slight preference for larger oligosaccharides is apparent (Figure 1D). After the second phase all the glycans are depleted >80%, with the exception of the smallest oligosaccharide mass (Figure 1D, Peak A), which corresponds to two isomers present at high concentration in the HMO pool: lacto-*N*-tetraose (LNT, Gal β 1-4GlcNAc β 1-3Gal β 1-4Glc) and LNnT (Gal β 1-3GlcNAc β 1-3Gal β 1-4Glc) (see Figure S1). *Bt* exhibits no preference for fucosylated or nonfucosylated glycans. These data demonstrate *Bt*'s capacity to utilize a broad range of HMOs, adding to its previously described saccharolytic capacity for mucus glycans and plant polysaccharides (Sonnenburg et al., 2005).

***B. thetaiotaomicron* HMO Use Is Coupled to Upregulation of Mucin Glycan Degradation Pathways**

To identify which portion of the *Bt* transcriptional response in MM-HMO was due to the complex oligosaccharides versus the simple core sugars, lactose and galactose, we performed transcriptional profiling of *Bt* grown in MM-galactose and MM-lactose ($n = 2$ mid-log phase for each monophasic growth) (Figure S2A). Comparison of these data with the baseline MM-glucose revealed 40 and 34 *Bt* genes are significantly upregulated (>5-fold) in MM-galactose and MM-lactose, respectively. Thirty-two of the 137 genes that were upregulated at least 5-fold in the MM-HMO response were also upregulated in the presence of galactose and/or lactose, consistent with the presence of these simple sugars in the core structures of all HMOs (Figure S2B). The 105 genes that were upregulated in either phase of HMO growth and not in MM-galactose or MM-lactose were highly enriched in the COG functional group "carbohydrate metabolism and transport" (32.1% compared with 11% representation across the genome) (Figure S2C). These data suggest that this 105-gene signature captures the *Bt* response to the structural complexity within HMOs.

Eighty of the 105 HMO-specific genes are found within ten PULs (loci involved in oligosaccharide acquisition and degradation). The larger 253-gene group that is upregulated in both phases of HMO growth included regions of three additional PULs, which resulted in 13 PULs or partial PULs upregulated in one or both phases of HMO growth (Figure 2A). Three of these PULs encode putative fucosidases, key enzymes that hydrolyze the terminal fucose residues from HMOs (BT1624–BT1632, BT3172–BT3173, BT4132–BT4136) and three endo- β -*N*-acetylglucosaminidases. Average fold changes for genes within each PUL or partial PUL ranged from 8- to 173-fold (Table S2).

Bt profiles in previously defined conditions were examined to search for overlap with the HMO response. Substrate specificities of several *Bt* PULs have been inferred using growth

conditions in which *Bt* is reliant upon host-derived gut mucus glycans. We compared our in vitro *Bt* HMO growth expression data with that from three different experimental paradigms in which *Bt* is reliant upon host mucus glycans: (i) in vitro growth in purified porcine mucus glycans (PMG) ($n = 3$ replicates/growth phase, 2 time points, during exponential phases from biphasic growth) (Martens et al., 2008); (ii) in vivo *Bt*-colonized 17-day-old gnotobiotic suckling mice ($n = 6$ samples) (Bjursell et al., 2006); and (iii) in vivo *Bt*-colonized adult gnotobiotic mice that were fed a diet lacking complex glycans ($n = 3$ samples) (Sonnenburg et al., 2005). Transcriptional profiles were analyzed using *Bt* grown in MM-glucose in vitro as a baseline (Table S2 lists genes upregulated in at least one condition).

Nine of the 13 upregulated PULs or partial PULs in MM-HMO were also highly upregulated (≥ 10 -fold induction) in one or more of the conditions of *Bt* grown in mucin glycans (Figure 2A). This overlap between HMO- and mucin-induced genes presents the possibility that *Bt* responds to common structural motifs found in oligosaccharides from mother's milk and intestinal mucin glycans. For instance, in all datasets we observed increased expression of the PUL BT2818–BT2826, which encodes glycoside hydrolases predicted to cleave linkages from Gal β -GlcNAc β -Gal, a structure common to HMOs and mucins. Alternatively, four *Bt* PULs exhibited increased expression specific to HMOs: three complete PULs (BT2618–BT2633, BT3172–BT3173, and BT3958–BT3965) and one partial PUL (BT3172–BT3173) (Figure 2A). These data indicate that *Bt* responds to aspects of the milk-derived glycans that are not found appreciably in the mucin preparations.

Comparing the structures of HMOs and human intestinal mucin glycans, it is apparent that HMOs exhibit less structural complexity (Karlsson et al., 1997; Robbe et al., 2004; Robbe et al., 2003a; Robbe et al., 2003b; Thomsson et al., 2002) (Figure 2B). Mucin glycans are typically built upon an *N*-acetylglucosamine that is *O*-linked to serine and threonine residues of the mucin protein, and the most abundant are based on five different core structures. Milk oligosaccharides are elaborated from a galactose of the "core" lactose disaccharide that is analogous to the reducing GalNAc of mucin *O*-linked glycans; structures similar to core 3 [GlcNAc β 1-3Gal] and core 4 [GlcNAc β 1-3(GlcNAc β 1-6)Gal] are present in HMOs. Structures very similar to human mucin glycans are found in the porcine and mouse mucin glycans, which have been used experimentally to define *Bt*'s mucus use capability (Martens et al., 2008) (Figure 2B). In both the intestinal mucins and in HMOs, repeated motifs containing galactose and *N*-acetylglucosamine are present and terminate with fucose and sialic acid residues. The extensive structural similarity between HMOs and mucins provides a parsimonious explanation for *Bacteroides* species upregulating the same PULs for the utilization of glycans from these two different sources.

We tested if the HMO-induced PULs are required for HMO consumption by creating *Bt* mutants in four of the HMO-specific loci. In-frame deletions for the respective *susC*-like genes, involved in carbohydrate binding, and the fucosidase BT4136 containing PUL (the most upregulated HMO-responsive glycoside hydrolases in *Bt*) were constructed, but showed no defect in growth in HMOs in vitro (see Figure S2D). These data indicate that extensive degeneracy exists within *Bt*'s HMO response,

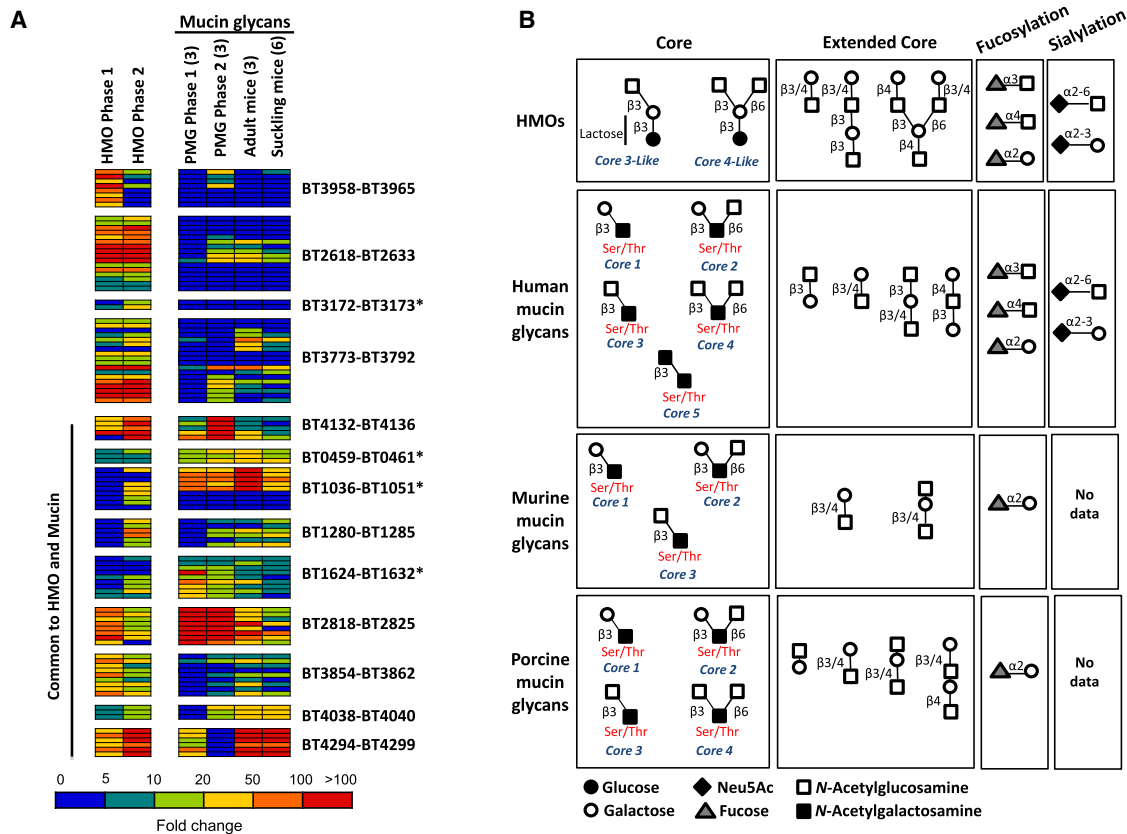


Figure 2. *B. thetaiotaomicron* Upregulates Mucus-Utilization Loci during HMO Consumption

(A) Gene expression profile of *Bt*'s induced PULs or partial PULs (*) in MM-HMO, MM-porcine mucin glycans (PMG), or host intestinal glycans from adult or suckling mice relative to MM-glucose. Parentheses denote sample number per condition. See also Figure S2 and Table S2.

(B) Schematic of mucin glycans based on previous reports (Hurd et al., 2005; Karlsson et al., 1997; Robbe et al., 2004; Robbe et al., 2003b; Thomsson et al., 2002). Structural information includes the core, extended core, and terminal (fucosylation, sialylation) motifs.

which contrasts to the strict requirement of genes within the *Bt*'s fructan-utilization locus (Sonnenburg et al., 2010).

B. fragilis Upregulates a Distinct Set of Mucin-Use Genes when Consuming HMOs

The HMO-induced *Bt susC/susD* homologs and genes located adjacent to the *susC/susD* genes were used as markers of HMO-utilization genes to identify the presence or absence of orthologs across five sequenced *Bacteroides* (*Bf*, *B. caccae*, *B. ovatus*, *B. ovatus*, and *B. stercoris*). In cases where a species shares an orthologous *susC/susD* pair with *Bt*, adjacent genes within the PUL generally display a lack of conservation (Figure S3). *Bf* grows efficiently in MM-HMOs (Figure 1B), but does not show conservation of any of the HMO-utilization PULs identified in *Bt*. These results suggest that *Bacteroides* species have developed diverse strategies for using HMOs with varying levels of efficiency.

Whole genome transcriptional profiling of *Bf* at mid-log phase of its monophasic growth in either MM-HMO, MM-lactose, MM-galactose, or MM-glucose was used to identify the genes upregulated by *Bf* during HMO consumption. MM-glucose served as a baseline to define the genes that were upregulated in HMO, but were not upregulated in MM-lactose or MM-galac-

tose ($n = 2$ per condition). These data revealed a *Bf* response to HMO composed of a much smaller set of genes compared to *Bt*. We identified 21 genes specifically upregulated (>5-fold) by *Bf* in HMO (See Table S3) compared to 105 genes by *Bt*. Just four *Bf susC/susD*-homolog-containing PULs were upregulated in HMO compared to 13 in *Bt* (Figure 3A).

Twelve of the 21 HMO-specific *Bf* genes were distributed within two loci that are dedicated to sialic acid acquisition and catabolism (Brigham et al., 2009) (Figure 3B). These genes include a neuraminidase (*nanH*, BF1806), an *N*-acetyl neuraminase permease (*nanT*, BF1724), an *N*-acetylneuraminase lyase (*nanL*, BF1712), and an *N*-acetylmannosamine 2-epimerase (*nanE*, BF1713) (upregulated 7.8-fold, 5.8-fold, 6.9-fold, and 6.0-fold, respectively) and provide the machinery necessary to cleave and catabolize Neu5Ac from sialylated glycans. To confirm that *Bf* does in fact consume sialic acids from HMOs, sialic acid content in the MM-HMO after growth was measured before and after acid hydrolysis using derivatization with 1,2-diamino-4,5-methylenedioxybenzene followed by reverse phase HPLC. The sialic acid Neu5Ac was completely depleted by *Bf* after growth in HMO. In contrast, *Bt* can cleave Neu5Ac from sialylated HMOs, presumably to access underlying sugars, but is unable to catabolize it (Figure 3C).

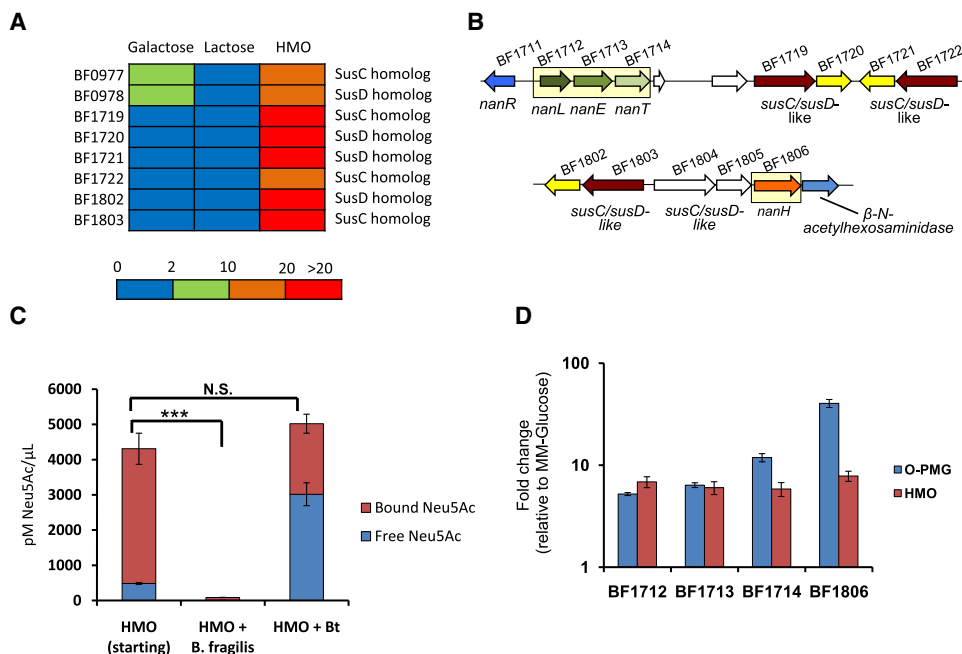


Figure 3. *B. fragilis* Response to HMOs Includes Sialic Acid Catabolism

(A) *Bf susC/susD* homologs upregulated (fold change) in vitro in MM-galactose, MM-lactose, and MM-HMO relative to MM-glucose. See also Figure S3.

(B) Genomic organization of *Bf* genes with >5-fold induction in MM_HMO relative to MM-glucose. Yellow boxes frame genes related to sialic acid consumption. White genes are upregulated <5 fold. Table S3 lists all significant HMO upregulated genes.

(C) HMO-bound versus liberated Neu5Ac content in MM-HMO and in MM-HMO after *Bf* and *Bt* growth. Error bars represent standard error for three biological replicates.

(D) Fold-induction of sialic acid-related genes from *Bf* grown in MM-O-PMG and MM-HMO relative to growth in MM-glucose, as measured by qRT-PCR. Error bars represent standard error for three biological replicates.

We wondered if *Bf* exhibited an overlap in the general strategies used for accessing HMOs and mucin glycans similar to that observed in *Bt*. The expression of genes that represent the *Bf* response to HMOs (BF1712, BF1713, BF1714, and BF1806) in MM supplemented with the O-glycan fraction from porcine mucin (MM-O-PMG) and MM-glucose was measured by qRT-PCR. We found that all four genes are upregulated in the presence of mucin, confirming that *Bf* upregulates sialic acid-use pathways in the consumption of both HMOs and intestinal mucin (Figure 3D). Therefore, while the specific PULs employed for HMO use by *Bf* and *Bt* differ, the mobilization of mucin-use PULs for HMO consumption is conserved between these *Bacteroides* species.

***B. thetaiotaomicron* and *Bifidobacterium infantis* Differentially Consume the Structurally Similar Mucin Glycans and HMO**

Previous work has demonstrated that *Bifidobacterium* species are well adapted to HMO use (Sela and Mills, 2010). We grew an HMO-consuming *Bifidobacterium* strain that is abundant in the microbiota of breast-fed infants, *Bi*, in purified porcine mucin glycans (MM-O-PMG) to elucidate its competence in mucin consumption (Haarman and Knol, 2006). Despite its proficiency at HMO use, *Bi* fails to grow in the MM-O-PMG (Figure 4A). These data suggest that structures unique to HMOs (i.e., not found in mucin glycans) are responsible for supporting growth of the HMO-adapted *Bifidobacterium* strain. *Bi* has previously

been shown to exhibit a preference for the smaller oligosaccharides found in HMOs, some of which are structurally distinct from mucin glycans (LoCascio et al., 2007). One such simple oligosaccharide is the four-sugar LNNt (Gal β 1-3GlcNAc β 1-3Gal β 1-4Glc). Therefore, we grew both *Bt* and *Bi* in a pure preparation of this single human milk oligosaccharide (MM-LNNt). *Bi* grew efficiently reaching a high OD, while *Bt* did not (Figure 4A).

These results suggest a model in which the structural complexity of HMOs includes structures that represent mucin glycans that may promote colonization of mucin-adapted symbionts, like *Bacteroides*, early in life. HMOs also contain structures distinct from mucins, such as LNNt, which may provide a niche for species that are important in microbiota assembly but that are not well adapted to mucus, like *Bi*, and would otherwise be outcompeted by mucus-adapted species like *Bacteroides*. The incomplete overlap of these glycan structural features allows HMOs to attract both HMO-adapted *Bifidobacterium* species and mucus-adapted *Bacteroides* species simultaneously (Figure 4B).

We performed an in vivo experiment to test whether LNNt could exhibit selectivity in vivo, enabling the expansion of a *Bifidobacterium* species in the presence of a *Bacteroides* species. Two groups of 6-week-old germ-free mice were fed a polysaccharide-deficient diet, forcing a reliance on host mucus glycans for carbon and energy. One group received LNNt supplementation in the water (1% w/v; average consumption of 75 mg of LNNt daily), the control group received plain water.

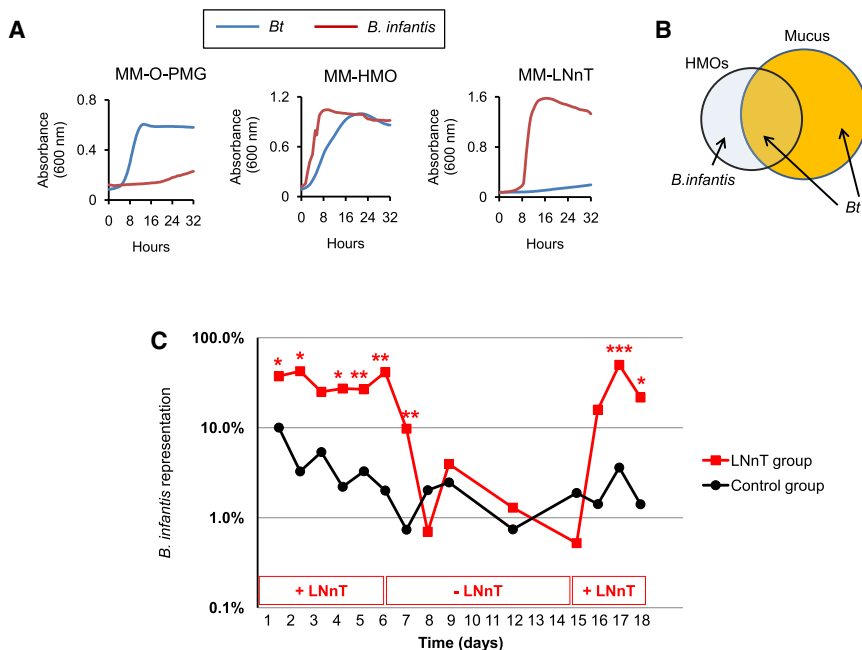


Figure 4. Selective Use of the HMO Lacto-N-Neotetraose by *B. infantis* Provides In Vivo Advantage

(A) In vitro growth of *Bt* and *B. infantis* in the presence of MM-O-PMG, MM-HMO, or MM-LNnT.

(B) Venn diagram representing the structural relationship of mucin and milk glycans. HMOs include a subset of structures found in mucus that can be consumed by mucus-adapted mutualists (e.g., *Bacteroides*). *B. infantis* is adapted to use simple structures within HMOs (e.g., LNnT) and is unable to use the structures found in mucin glycans.

(C) *Bt* and *B. infantis* biassociation of adult germ-free mice fed a polysaccharide-deficient diet without (black circles) or with LNnT (red squares). Values represent average of fecal communities within each group (n = 4 mice/group).

After one day of LNnT feeding, both groups were colonized with equivalent quantities of *Bt* and *Bi* (10^8 CFU each). At day 6 post-inoculation, the group receiving LNnT was switched to normal water; at day 15 postinoculation, LNnT was readministered for 3 additional days. Total bacterial colonization density was determined by assessing the CFUs in feces over the course of the 18 day experiment.

The administration of LNnT resulted in an expanded population of *Bi* relative to *Bt* throughout the first period (day 1–6) compared to control ($41.5 \pm 6.4\%$ versus $2 \pm 0.5\%$ on day 6; $p = 0.009$, day 6, n = 4 mice) (Figure 4C). LNnT supplementation was withdrawn on day 6 and colonization density was determined on days 7, 8, 9, 12, and 15 post colonization (1, 2, 3, 6, and 9 days after removing LNnT from the water). Removal of LNnT resulted in a drop in *Bi* to levels similar to controls within 2 days. At day 3 post withdrawal of LNnT, *Bi* representation was $3.9 \pm 0.2\%$ compared to $41.5 \pm 6.4\%$ just prior to withdrawal ($p = 0.006$, day 6 versus day 9, n = 4 mice). LNnT was reintroduced into the water on day 15, and bacterial colonization was monitored at days 16, 17, and 18 post colonization. Two days after LNnT reintroduction, *Bi* expanded from $0.5 \pm 0.5\%$ at day 15 to $49.9 \pm 2\%$ at day 17 ($p = 0.012$, day 15 versus day 17, n = 4 mice). By day 18, *Bi* represented $25.9 \pm 9.6\%$ of the community in LNnT-fed mice, compared to $1.9 \pm 1.5\%$ in the control group ($p = 0.014$, n = 4 mice) (Figure 4C). These results confirm that this short milk oligosaccharide provides a selective advantage to a *Bifidobacterium* species over a *Bacteroides* species in vivo.

DISCUSSION

HMO use by *Bt* and *Bf* results in the induction of PULs involved in mucin consumption. One important question is whether HMOs or mucin was the primary selective force shaping locus functionality. For strains that are strictly adapted to one of the glycan

sources, the selective pressure can be more confidently assigned, such as *Bt*'s adaptation to HMOs. However for *Bt*, it is less clear whether (i) adaptation to mucin glycans provides extant pathways that may be mobilized for HMO use or (ii) HMOs play a significant role in *Bt* propagation and provide a selective force, in addition to mucin, that shapes multifunctional loci adapted to accommodate the unique structural aspects of HMOs and mucin glycans. The inability of *Bt* to readily consume LNnT argues against the latter. If *Bt*'s entire HMO response is a result of HMO glycans co-opting extant pathways, it is unclear why a subset of *Bt*'s transcriptional response to HMOs includes loci that are not induced by mucin use. Considering the more diverse carbohydrate structures that human mucus contains, compared to the pig or mouse mucin glycans, it is possible that the set of *Bt* genes identified as mucin-glycan responsive is not exhaustive. Specifically, the complete set of *Bt* genes responsive to human mucus, once defined, may include the loci that now appear to be "HMO-specific."

The lack of requirement of the HMO-specific loci for efficient growth in HMOs by *Bt*, as demonstrated by our gene deletion studies, may appear to support that these loci are not HMO-adapted, but rather are co-opted by the structurally similar HMOs. However, mutation of five mucin-glycan-use PULs in a quintuple KO strain of *Bt* did not reveal a profound growth phenotype during use of mucin glycans in vitro (Martens et al., 2008). Rather, the requirement of these PULs became apparent upon competition with the WT parent strain in vivo. Therefore, our results highlight the caveats associated with assessing the importance of a gene or locus of a gut resident in a noncompetitive environment. Due to the paucity of pure and complex HMOs, in vivo competition experiments are not currently possible.

One hypothesis that arises from this work is that HMO structures mimic mucus glycans to attract mucin-adapted resident mutualists to an infant microbiota. Attraction of *Bacteroides* may provide the additional benefit of seeding the community with species that are also well adapted to dietary glycan use, thus preparing the infant microbiota for a smooth transition upon the introduction of solid food. This possibility is supported

by recent metagenomic studies that have revealed an abundance of plant polysaccharide-degrading glycoside hydrolases within the gut microbiomes of breast-fed infants (Koenig et al., 2011; Vaishampayan et al., 2010). At the same time, unique structural features of HMOs, such as LNnT, are potentially important in shaping the infant microbiota in ways that are independent of mucus use.

EXPERIMENTAL PROCEDURES

Bacterial Strains and Culture Conditions

Bacterial strains used are listed in Supplemental Experimental Procedures. Type strains were used unless otherwise indicated. *Bacteroides* species were grown in tryptone-yeast extract-glucose (TYG) medium and minimal medium (MM) (Martens et al., 2008). *Bi* was grown in Reinforced Clostridial Medium (RCM) (Becton Dickinson, MD) and minimal medium consisting of modified de Man-Rogosa-Sharpe medium (MRS) (Oxoid, Basingstoke, Hampshire, UK), which lacks glucose. Carbon sources were added at 0.5% (w/v) to the respective MM for *Bacteroides* or *Bifidobacterium* with the exception of HMO, LNnT, and PMG, which were added at 1.5% (w/v). OD₆₀₀ was monitored using a BioTek PowerWave 340 plate reader (BioTek, Winooski, VT) every 30 min, at 37°C anaerobically (6% H₂, 20% CO₂, 74% N₂).

Whole Genome Transcriptional Profiling

Transcriptional profiling of *Bt* and *Bf* was performed in MM supplemented with milk oligosaccharides, glucose, galactose, and lactose (Sigma-Aldrich). *Bt* samples profiled in HMO were collected at OD₆₀₀ = 0.5 and OD₆₀₀ = 0.8. Cultures grown on galactose, lactose, and glucose were collected at OD₆₀₀ = 0.5. RNA targets were prepared and hybridized to custom *Bt* Affymetrix GeneChips or *Bf* GeneChips (Sonnenburg et al., 2005). (See Supplemental Experimental Procedures for *Bf* GeneChip generation and validation and GeneChip data analysis).

Genetic Manipulation of *B. thetaiotaomicron*

Genetic manipulation of *Bt* was performed using counterselectable allele exchange, resulting in “in-frame” gene deletion (Martens et al., 2008).

Glycoprofile Analysis of HMO Consumption

Bacteria culture supernatants in MM-HMO were collected by centrifugation. Remaining oligosaccharides were recovered and profiled by HiRes matrix-assisted laser desorption/ionization-Fourier transform ion cyclotron resonance mass spectrometry (MALDI-FTICR-MS) (Marcobal et al., 2010).

Sialic Acid Content Determination

Culture supernatant was isolated after growth in MM-HMO and clarified by centrifugation and filtration. Sialic acid concentrations were determined by the Glyco-technology Core Facility UCSD (University of California, San Diego, San Diego, CA), using the Sigma DMB labeling kit protocol (Prozyme, San Leandro, CA).

Competitive Colonization of Gnotobiotic Mouse

Germ-free Swiss-Webster mice were reared in gnotobiotic isolators and fed an autoclaved polysaccharide-deficient diet (BioServ, <http://bio-serv.com>) in accordance with A-PLAC, the Stanford IACUC. Mice were biassociated using oral gavage (10⁸ CFU of each species). Subsequent community enumerations from mice were determined from freshly collected feces, by selective plating of serial dilutions in RCM agar and BHI-blood agar supplemented with gentamicin (200 mg/ml). Significant differences between sample groups were determined using Student's *t* test. Synthetic LNnT (Glycom A/S, Denmark) was purified by crystallization to a final purity of >99%. Characterization was performed using multiple methods, including NMR (1D and 2D) mass spectrometry, and HPLC.

ACCESSION NUMBERS

GeneChip data files have been deposited to the Gene Expression Omnibus under accession numbers GSE26772 and GSE32207.

SUPPLEMENTAL INFORMATION

Supplemental Information includes three figures, Supplemental Experimental Procedures, Supplemental References, and three tables and can be found with this article online at [doi:10.1016/j.chom.2011.10.007](https://doi.org/10.1016/j.chom.2011.10.007).

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