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The evolution of cooperation within the gut microbiota

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Abstract

Cooperative phenotypes are considered central to the functioning of microbial communities in many contexts, including communication via quorum sensing, biofilm formation, antibiotic resistance, and pathogenesis¹⁻⁵. The human intestine houses a dense and diverse microbial community critical to health^{1,2,4-9}, yet we know little about cooperation within this important ecosystem. Here we experimentally test for evolved cooperation within the Bacteroidales, the dominant Gram-negative bacteria of the human intestine. We show that during growth on certain dietary polysaccharides, the model member *Bacteroides thetaiotaomicron* exhibits only limited cooperation. Although this organism digests these polysaccharides extracellularly, mutants lacking this ability are outcompeted. In contrast, we discovered a dedicated cross-feeding enzyme system in the prominent gut symbiont *Bacteroides ovatus*, which digests polysaccharide at a cost to itself but at a benefit to another species. Using in vitro systems and gnotobiotic mouse colonization models, we find that extracellular digestion of inulin increases the fitness of *B.ovatus* due to reciprocal benefits when it feeds other gut species such as Bacteroides vulgatus. This is a rare example of naturally-evolved cooperation between microbial species. Our study reveals both the complexity and importance of cooperative phenotypes within the mammalian intestinal microbiota.

A major challenge facing the study of host-associated microbiotas is to understand the ecological and evolutionary dynamics that shape these communities^{2,5,10-13}. A key determinant of microbial dynamics is the balance of cooperation and competition both within and between populations^{2,5,14,15}. Here we test for the evolution of cooperation within

Author Contributions

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S.R-N performed mutant construction bacterial cultures, gnotobiotic experiments, protein purification and TLC, L.E.C. assisted with mutant construction. S.R-N, and L.E.C analyzed the data. S.R-N, K.R.F and L.E.C designed the study and wrote the paper.

the mammalian microbiota by focusing on the Bacteroidales, the most abundant order of Gram-negative bacteria of the human intestine with species that co-colonize the host at high densities of 10¹⁰-10¹¹ CFU/gram feces^{16,17}. Members of this order breakdown polysaccharides outside of their cell using outer surface glycoside hydrolases^{6,18}, some which are secreted on outer membrane vesicles^{3,19}. This suggests a significant potential for one cell to cooperatively feed other cells. As extracellular digestion is considered essential for growth of Bacteroidales on polysaccharides^{1,6,8}, we focused on this trait as a candidate for cooperative interactions within the gut microbiota and asked whether a bacterium that breaks down a polysaccharide extracellularly receives most, or all²⁰, of the benefits of its efforts.

We first made isogenic mutants in genes responsible for the extracellular digestion of polysaccharides in the well-studied human gut strain, *Bacteroides thetaiotaomicron (Bt)* VPI-5482^{1,8}, Specifically, we deleted the genes encoding the outer surface glycoside hydrolases BT3698 of the amylopectin/starch utilization locus and BT1760 of the levan utilization locus (Fig.1a), required for growth on amylopectin and levan, respectively (Fig. 1b,c, Extended Data Fig.1a,b)^{1,8}. Consistent with previous observations^{1,8}, neither mutant grew with the specific polysaccharide in monoculture (Fig.1b,c, Extended Data Fig.1a,b). However, co-culture of BT3698 or BT1760 with WT in amylopectin or levan increased the fitness of the mutants (Fig.1d,e, Extended Data Fig.1c,d). This is consistent with cooperation via public good availability of amylopectin and levan breakdown products.

One of the key questions in evolutionary biology is how cooperative systems can be evolutionarily stable^{4,5,14,21,22}. If certain cells invest in the production of an enzyme that helps others, what prevents these cells from being outcompeted by cells that consume the breakdown products without making the enzyme? In the *Bt* amylopectin and levan polysaccharide utilization systems, while receiving public goods benefits provided by WT cells (Extended Data Fig.1e), mutant cells do not outcompete the WT. Cells that make the enzyme receive more benefits than non-producing neighboring cells suggesting that private²³ benefits are central to the evolutionary stability of polysaccharide breakdown in these systems.

We extended our analysis to another prominent member of the human Bacteroidales known to extensively utilize polysaccharides, *Bacteroides ovatus* $(Bo)^{24}$. During growth on inulin, a dietary fructan known for health promoting effects²⁵, *Bo* extracellularly digests and liberates considerable amounts of inulin breakdown products³. The predicted inulin utilization locus of the *Bo* type strain ATCC 8483 encodes two similar outer surface glycoside hydrolases, BACOVA_04502 and BACOVA_04503 (Fig.2a), both of which are predicted to target the β 1,2 inulin fructose polymer^{1,3}. Both of these enzymes are required for inulin breakdown (Fig.2b). We therefore predicted that a single mutant of either BACOVA_04502 or BACOVA_04503 would be unable to grow on inulin. Surprisingly, neither of the single deletion mutants (04502 or 04503) nor the double mutant (04502/3) demonstrated impaired fitness with inulin as the sole carbohydrate source (Fig.2c, Extended Data Fig.2a), even at limiting concentrations (Extended Data Fig.2a).

Given the importance of extracellular polysaccharide digestion for growth of *Bacteroides* on numerous polysaccharides^{1,6,8} (Fig.1, Extended Data Fig.1), we predicted that *Bo* synthesizes other enzymes that breakdown inulin extracellularly, allowing the *Bo* mutants to grow on this polysaccharide. However, analysis of the growth media of 04502, 04503 and

04502/3 revealed no released inulin breakdown products demonstrating that BACOVA_04502 and BACOVA_04503 are solely responsible for extracellular digestion of inulin (Fig.2d, Extended Data Fig.2b; see Extended Data Fig.2c for complementation). Deletion of BACOVA_04504 or BACOVA_04505 encoding SusD and SusC orthologs, respectively, encoding predicted inulin binding and import machinery^{6,26} resulted in significant impairment of growth on inulin (Fig.2e, Extended Data Fig.3a; see Extended Data Fig.3b for complementation). Growth of 04502, 04503 and 04502/3 in limiting concentrations of inulin revealed depletion of inulin (Fig.2d, right panel; Extended Data Fig. 2b). Together, these data demonstrate that surface enzymes 04502 and 04503 are not needed for *Bo* to utilize inulin, and that inulin is directly imported via 04504-04505 without prior extracellular digestion.

Why would *Bo* synthesize surface/secreted enzymes that potently digest inulin outside of the cell if not necessary for its growth on the polysaccharide? A key evolutionary explanation for the release of secreted products by microbes is that they feed clonemates in a manner that is beneficial at the level of the clonal group^{2,5,11,14,15}. We hypothesized that the importance of extracellular digestion may be realized during spatially-structured growth on plates where not all cells are in direct contact with the polysaccharide. However, mutant bacteria showed no significant differences in growth yield compared to WT on defined inulin plates (Fig.3a; Extended Data Fig.4a). In addition, these enzymes were not required for optimal growth *in vivo as* WT and 04502/3 showed equal colonization levels in gnotobiotic mice fed a polysaccharide-free diet with inulin added as the sole dietary polysaccharide (Fig.3b). Therefore, we could find no evidence that inulin breakdown by 04502 and 04503 benefits *Bo* in three-dimensional growth or during monocolonization of the mammalian gut.

Bacteroidales polysaccharide utilization loci can be induced by monomers or oligomers of the utilized polysaccharide^{1,26}. This raised the possibility that 04502 and 04503 may be important for optimal growth on inulin during induction. However, while addition of trace amounts of fructose monomers or oligosaccharides led to accelerated growth on inulin (Fig. 3c, Extended Data Fig.4b) this did not require 04502/3 (Fig.3c, Extended Data Fig.4b). Rather than a benefit from the presence of the enzymes, we observed a cost based on reduced yield of the WT compared to 04502/3 mutant during induced growth (Fig.3c, Extended Data Fig.4b) that was independent of a direct energetic cost of synthesis of 04502/3 (Fig.2c, e, Fig.3c, d, Extended Data Fig.2a, Extended Data Fig.4b, Extended Data Fig.5a, b, c). We find that Bo grows better on undigested inulin than an equal concentration of inulin breakdown products (Fig.3d, Extended Data Fig.5a,b). In addition, the yield advantage to the mutant does not occur under limiting inulin concentrations (Extended Data Fig.5d,e). Furthermore, Bo preferentially consumes longer inulin digestion products over shorter oligomers and fructose (Fig.2d, right panel, Extended Data Fig.2b, right panel). Together, these data suggest that undigested inulin is the preferred substrate of Bo, and that extracellular digestion by 04502/3 can be costly for fitness.

We found no evidence that extracellular digestion of inulin by *Bo* evolved for cooperation with clonemates. Therefore, we speculated that this trait might have evolved for cooperation with other species in the gut. We first sought evidence of cooperation in the setting of a natural gut ecosystem. Germ-free mice were fed inulin and colonized with either *Bo* WT or 04502/3 followed by the introduction of the cecal microbiota of conventionally raised

O(4502/3) followed by the introduction of the cecar interobiota of conventionally faised mice. *Bo* WT and O(4502/3) equally colonized mice before the introduction of the microbiota (Extended Data Fig.6a), but *Bo* WT received a significant fitness benefit compared to O(4502/3) in the context of a complex microbiota (Fig.4a). These data suggested that while not required for *Bo* to utilize inulin, O(4502) and O(4503) provides a benefit to *Bo* only realized in a community setting. The conditions for the evolution of cooperation between species are much more restrictive than those within a clone. In particular, theory predicts that costly interspecies cooperation will only be stabilized if there are reciprocal feedback benefits, such as a plant providing nectar for an insect that pollinates it^{21,22}. From this experiment, we identified two dominant mouse microbiota Bacteroidales strains that thrived on *Bo* derived inulin breakdown products (Extended Data Fig.6b), with delayed growth on inulin (Extended Data Fig.6b). These data suggest that cross-feeding Bacteroidales members may provide reciprocal benefits to wild type *Bo* in the mammalian gut.

To experimentally test for reciprocity and benefits of inulin digestion between species, we used an inulin co-culture system with *Bo* WT or 04502 and *Bacteroides vulgatus* ATCC 8482 (*Bv*), which is commonly found together with *Bo* at high densities in humans ^{16,17} and thrives on inulin breakdown products³ but cannot use inulin^{1,3}. Co-culture and proximate plating with WT *Bo* increased the fitness of *Bv* compared to that with 04502 (Fig.4b,c Extended Data Fig.7a,9a); however, *Bv* is able to persist better with 04502 than when alone (Fig.4b, Extended Data Fig.7a) due to a small (<2000 Da MW), secreted molecule (s) that contributes to the survival of *Bv* (Extended Data Fig.8a,b). This 04502/3-independent survival is not mediated by a universal factor made by *Bacteroides* during growth on inulin nor *Bo* derived short chain fatty acids (Extended Data Fig.7a), the greatest being crossfeeding mediated by 04502/3.

We next addressed the critical question of whether *Bo* receives reciprocal benefits from *Bv*. Co-culture of *Bo* with *Bv* on plates increased the fitness of *Bo* WT and *Bo* 04502 (Fig.4d, Extended Data Fig.7b), but did not increase the fitness of *B. fragilis* (Extended Data Fig.7c), If inulin breakdown can be costly, and *Bo* receives benefits from *Bv* irrespective of whether inulin is broken down and fed to *Bv*, natural selection is expected to favor the loss of the genes encoding the secreted inulin glycoside hydrolases²². However such pairwise partnerships would not reveal the possibility that *Bo* WT receives more reciprocal benefits from *Bv* when in direct competition with the non-cross-feeding mutant^{21,27}. Therefore, we co- and tri-cultured these strains on plates and compared the yields of the two *Bo* strains (WT and 04502/3) with or without *Bv*. Addition of *Bv* leads to an increased proportion of WT *Bo* compared to the 04502/3 mutant (Fig.4e, Extended Data Fig.9b). We extended these studies to a gnotobiotic mouse colonization model. WT *Bo* had no advantage in direct competition with 04502/3 on a polysaccharide-free diet or when inulin is added (Fig.4f). However, introduction of *Bv* increased the fitness of the *Bo* WT relative to the mutant (Fig.

4f, Extended Data Fig.9c,d). Together, these data suggest that extracellular breakdown of inulin increases the fitness of *Bo* via reciprocal benefits from another species. These findings are consistent with the evolution of cooperation between species within the gut microbiota.

We find evidence of distinct forms of cooperativity within the Bacteroidales of the human intestinal microbiota (Extended Data Fig.10). For *Bt*, amylopectin and levan digestion provide mostly private benefits and modest social benefits to other cells. By contrast, *Bo* releases large amounts of inulin digestion products via a pair of dedicated cross-feeding secreted enzymes unnecessary for its use of inulin. These enzymes allow for cooperation with cross-fed species, which provide benefits in return. Potential mechanisms by which *Bv* may provide return benefits to *Bo* include detoxification of inhibitory substances, or production of a depleted or growth promoting factor, the latter supported by early growth benefits to *Bo* via *Bv* secreted factors (Extended Data Fig.9e).

Understanding whether microbial communities are formally cooperative is central to predicting their evolutionary and ecological stability. While cooperative systems can be productive, they are prone to instabilities on both ecological and evolutionary timescales that can undermine them^{5,14,15,21,22}. The ability of one species to utilize the waste product of another is prevalent, but waste production alone does not signify cooperative evolution. As opposed to waste product utilization or exploitive interactions²⁸⁻³⁰, there are few well-documented cases of evolved cooperation between microbial species¹⁴. We have found evidence of strong eco-evolutionary interactions within the microbiota that are likely to be central to both the functioning and stability of these complex communities.

Methods

Bacterial strains and media

Bacteroidales type strains used in this study are, *Bo* ATCC 8483, *B. thetaiotaomicron* VPI 5482 *Bv* ATCC 8482, and *B. fragilis* NCTC 9343. Bacteria were grown in media formulation as previously described³.

Bacterial culture

For growth in defined media, bacteria were inoculated from brain heart infusion plates containing hemin and vitamin K (BHIS) plates into basal medium (BS), cultured overnight to stationary phase, then diluted 1:10 in fresh BS and grown to mid log. At mid log, bacteria were pelleted by centrifugation and washed with sterile phosphate buffered saline (PBS) and then inoculated in defined media. Carbohydrates used to supplement defined media include fructose (F2543, Sigma), fructose oligosaccharides (FOS; OraftiP95, Beneo-Orafti group), levan (L8647, Sigma), amylopectin (10120, Sigma), and inulin (OraftiHP, Beneo-Orafti group). Levan and amylopectin were autoclaved as 1% w/v in H₂O and dialyzed using 3.5kD MW membranes (Slide-A-Lyzer Dialysis Cassettes, ThermoScientific). Short chain fatty acids acetate, propionate and succinate were purchased from Sigma. Stock solutions of 2 mM were pH neutralized to pH 7.2-7.3 with 10N NaOH. All cultures were grown at 37° C under anaerobic conditions. Bacterial growth was quantified by optical density (OD₆₀₀) using 200 µl of bacterial culture in 96 well flat-bottom microtiter plates using a Powerwave spectrophotometer (Biotek). Murine gut Bacteroidales from the cecal preparations used in for the colonization experiments were grown on BHIS plates. Resulting colonies were tested for growth in inulin minimal medium or inulin breakdown products from the conditioned media of *Bo* grown in inulin, containing inulin breakdown products as previously described³.

Creation of Bacteroides mutants

Deletion mutants were created whereby the genes encoding BT3698 or BT1760 in *Bacteroides thetaiotaomicron* VPI 5482, BACOVA_04502 in *Bo* ATCC 8483 BACOVA_04503, BACOVA_04502/3, BACOVA_04504, or BACOVA_04505 in *Bo* ATCC 8483 were removed. DNA segments upstream and downstream of the region to be deleted were PCR amplified using the primers outlined in Extended Data Table 1. PCR products were digested with BamHI, EcoRI and/or MluI engineered into the primers (Supplementary Information Table 1) and cloned by three way ligation into the appropriate site of pNJR6³¹. The resulting plasmids were conjugally transferred into the *Bacteroides* strain as indicated using helper plasmid R751 and cointegrates were selected by erythromycin resistance. Cross outs were screened by PCR for the mutant genotype.

Cloning of PUL genes for expression in deletion mutants

BACOVA_04502, BACOVA_04503, BACOVA_04502/3, BACOVA_04504, BACOVA_04505 or BACOVA_04504/5 genes were PCR amplified using the primers listed in Extended Data Table 1. The PCR products were digested and ligated into the BamHI or KpnI site of the *Bacteroides* expression vector pFD340³². Plasmids containing the correct orientation of the insert in relation to the vector-borne promoter were introduced into mutant *Bacteroides* strains by mobilization from *E. coli* using helper plasmid RK231.

Mono-, co- and tri- culture experiments

For bacterial mono and co-culture experiments in defined liquid media, bacteria were grown as indicated for monoculture prior to addition to the defined media. Sterile magnetic stir bars were added to culture tubes within a rack placed on a stir plate within the anaerobic chamber. For conditioned media experiments, Bo WT, Bo 04502, Bo 04502/3, Bv, or Bf were grown to early log in inulin defined media or 0.125% fructose defined media (for Extended Data Fig.9e), conditioned media were collected, filter sterilized, and incubated at 37°C for 72 hrs. Conditioned media was replenished with defined media without additional carbohydrate and used for cultivation of Bv. Bo 04502/3 conditioned media was dialyzed in defined media without carbohydrate using 2 kD MW membranes (Slide-A-Lyzer Dialysis Cassettes, ThermoScientific). For monoculture of Bo WT and mutants on solid agarose, 4 µl of the indicated concentration of bacteria were dotted onto minimal inulin agarose plates. At the indicated timepoints, the bacteria were cut out, diluted and plated onto BHIS for CFU enumeration. For co-plating experiments, $10^6 Bo$ (WT or mutant) or Bf were co-plated with $10^5 Bv$ or a control volume of PBS. Four μ l were then dotted on to defined inulin agarose plates. At the indicated timepoints, the dotted patches were cut from the agarose plates and resuspended in PBS, diluted and plated to BHIS for enumeration. Quantification and differentiation of WT and isogenic glycoside hydrolase or polysaccharide lyase mutant was performed by plating dilutions of mixed liquid culture or the cut-out patch on agarose plates

onto BHIS, followed by picking ~100 colonies and determining WT or mutant genotype by PCR using primers listed in Extended Figure Table 1. For genotypic screening of *Bt* WT and *Bt* 3698, two sets of primers were used (Extended Table 1). For *Bo/Bv* co-culture in Fig. 4B,C and Extended Data Fig.7, *Bo* 03533 (WT) and *Bo* 03533 04502 (04502) arginine auxotrophic mutants were used in co-culture with *Bv* on minimal inulin agarose plates supplemented with 50 µg/ml of arginine (Sigma) which does not impair or limit growth as compared to WT³. Colonies on BHIS plates were replica plated onto defined glucose defined plates, which support the growth of *Bv* but not *Bo* 03533 or *Bo* 03533 04502.

Thin-layer chromatography

Thin-layer chromatography (TLC) was employed to specifically detect carbohydrates as previously described³. Standards for TLC included glucose (G7528, Sigma), fructose (F2543, Sigma), fructose oligosaccharides (FOS; OraftiP95, Beneo-Orafti group) and inulin (OraftiHP, Beneo-Orafti group). See Supplemental Information Figure 1 for uncropped TLCs.

Gas Chromatographic Analysis of Culture Media

Chromatographic analysis was carried out using a Shimadzu GC14-A system with a flame ionization detector (FID) (Shimadzu Corp, Kyoto, Japan). A volatile acid mix containing 10 mM of acetic, propionic, isobutyric, butyric, isovaleric, valeric, isocaproic, caproic, and heptanoic acids was used (Matreya, Pleasant Gap PA). A non-volatile acid mix containing 10 mM of pyruvic and lactic and 5 mM of oxalacetic, oxalic, methy malonic, malonic, fumaric, and succinic was used (Matreya, Pleasant Gap PA).

Cloning, purification, and enzymatic analysis of BACOVA_04502-3

To obtain purified BACOVA_04502 and BACOVA_04503 protein, these genes were cloned individually into the BamHI site of pET16b (Novagen) using the primers listed in Extended Figure Table 1. The constructs were designed so that the His-tag encoded by pET16b replaced the SpII signal sequence of these proteins allowing for their solubility. The recombinant plasmids were transformed into E. coli BL21 (DE3), grown to an OD₆₀₀ of 0.6 - 0.7, and expression of the recombinant gene was induced by the addition of 0.4mM IPTG for an additional 4 h. The His-tagged proteins were isolated essentially as described³³ using Dynabeads TALON paramagnetic beads. For enzymatic analysis (Fig.2A) the proteins were added to inulin media in their magnetic bead-bound form. This allows for easy removal of the enzymes following digestion. As a control, the beads resulting from the same procedure performed with E. coli BL21 (DE3) containing only the vector (pET16b) were used. For Bo growth assays (Fig.3d), 50 µl of beads (25 µl containing His-04502 and 25 µl containing His-04503) or equivalent volume of Dynabead buffer (for undigested inulin) were added to inulin defined medium. After 24 h at 37°C, the beads containing the enzyme were removed with a magnet, and the media (digested or undigested inulin) was used to culture Bo WT and 04502.

Gnotobiotic mouse experiments

All animal experiments were approved by the Harvard Medical School IACUC. Swiss Webster germ free male mice (6-10 weeks old) were purchased from the Harvard Digestive Diseases Gnotobiotic Core facility. Littermates were randomly allocated for different gnotobiotic experimental arms. Experiments were conducted in either sterile Optima cages (Fig.3b, 4a, Extended Data Fig.9d) or gnotobiotic isolators (Fig.4f). When appropriate, animals were numbered by sterile tail markers for longitudinal analysis and were not blinded. Number of animals used for experimentation was determined by precedence for gnotobiotic studies. Longitudinal analysis in the experiment of Fig.4f of the ratios of wild type to mutant Bo under changing environmental conditions allowed for internallycontrolled, within-individual comparisons. Mice were placed on polysaccharide free special chow (65% w/v glucose, protein-free, supplemented with all essential amino acids except arginine; BioServe). Arginine at 50 µg/ml was supplemented in all drinking water. As indicated, mice were given 1% inulin (w/v) in sterile drinking water. Mice were inoculated with the indicated bacteria by applying $\sim 10^8$ live bacteria (grown to mid log) onto mouse fur. Dilutions of feces at various time points following inoculation were plated to BHIS plates and genotyped as above (Fig.4f). Predetermined exclusion criteria for gnotobiotic experiments contamination as determined by either the presence of colonies with distinct morphology on anaerobic plates than Bo or Bv or colonies present at $>10^2$ CFU/ml (limit of detection) under aerobic conditions.

For gavage of Bo WT or 04502/3 monocolonized mice with the cecal content of conventionalized raised mice, two ~ 8 week old male Swiss Webster mice purchased from Taconic and housed in a specific pathogen free facility were sacrificed under sterile conditions. Intestine was excised and care was made to leave cecum intact. Within 2 minutes of excision, the ceca were transferred into an anaerobic chamber where the cecal contents were pooled and diluted with ~10 ml of pre-reduced phosphate buffered solution supplemented with 0.1% cysteine. Contents were vigorously vortexed and immediately divided equally into two conicals (one for each group of mouse) to ensure equality of suspension between both conicals. Tightly sealed conicals were transferred to the mouse facility and which point 200 µl were gavaged to each mouse housed in sterile Optima cages. Cecal contents were plated to LKV (Remel) plates for enrichment of Bacteroides. Different colony morphologies were speciated by 16S PCR as previously described¹⁷. For each group of mice, mice were housed in cages of 2 and 3 mice per cage. At time of sacrifice, cecal contents were collected and plated to defined inulin plates by which Bo was enumerated based on distinct colony morphology of Bo on inulin agarose plates that was not present in conventionally raised cecal population.

Statistical analysis

Replicate experiments are shown in Extended Data. All p values are derived from Student's t test except as indicated in Fig.4f and Extended Data Fig.9c where Fisher exact test was performed. Statistical significance of variance reported as indicated per experiment in Figure Legends All center values are mean. Error bars are standard error of the mean.

Extended Data



Extended Data Figure 1. Limited cooperation during polysaccharide utilization by *B. thetaiotaomicron*

a-d, Independent experiments for Fig. 1b-e. (**a,b**) Upper panels n=2 biological replicates, lower panels n=1; (**c,d**) n=2 biological replicates. **e**, TLC analysis of conditioned media from *Bt* grown in amylopectin (left panel) or levan (right panel) minimal media. EL = early log, ML = late log, LL = late log, Stat = stationary phase. Glu = Glucose, Fru = fructose, Suc = sucrose. See SI Figure 1 for uncropped scanned images. In all panels, error bars represent standard error; *p* values derived from two-tailed Student's t test.



Extended Data Figure 2. *Bo* 04502 and 04503 mutants grow equivalently to wild type on limiting concentrations of inulin and do not require surface digestion for utilization of inulin a, Growth of *Bo* WT, 04502, 04503 and 04502/3 mutants in varying concentrations of inulin as indicated. Biological replicates of each condition are plotted as individual lines (n=2 cell culture biological replicates). Upper and lower panels are independent experiments. **b**, Independent experiments for Fig. 2d. EL = early log, ML = mid log, LL = late log, St = stationary phase. See SI Figure 1 for uncropped scanned images. **c**, Complementation of *Bo* 04502 and 04502/3 mutants with the respective genes *in trans*. TLC analysis of conditioned media from *Bo* 04502/3 (left panel) complemented *in trans* with BACOVA_04502, BACOVA_04503, BACOVA_04502/3 or vector alone (pFD340) and *Bo* 04502 (right panel) with BACOVA_04502 or vector alone grown in defined inulin media. See SI Figure 1 for uncropped scanned images. **S** = stationary phase.



Extended Data Figure 3. SusC and SusD homologs BACOVA_04505 and BACOVA_04505 are required for inulin utilization

a, Independent experiments for Fig. 2e. Left panel n=2 biological replicates, right panel each line represents n=1 sample per condition. **b**, Complementation of *Bo* 04504, *Bo* 04505 and 04504/5 mutants with the genes *in trans*. Growth of *Bo* 04504, *Bo* 04505, *Bo*

04504/5 with BACOVA_04504, BACOVA_04505, BACOVA_04504/5 or vector alone (pFD340) *in trans* in defined inulin media. Each line represents n=1 sample per condition. In all panels, error bars represent standard error; *p* values derived from two-tailed Student's t test.



Extended Data Figure 4. Costs of extracellular inulin digestion by *B. ovatus*

a,b, Independent experiments for Fig. 3a,c. n=3 biological replicates at day 1,2; n=2, biological replicates at day 4 (**a**). In upper and upped middle panels biological replicates of each condition are plotted as individual lines (n=2 cell culture biological replicates); in lower middle and lower panels each line represents n=1 sample per condition (**b**). **c**, TLC analysis of conditioned media from *Bo* WT and *Bo* 04502/3 cultured in 0.5% inulin with trace (0.06%) amounts of fructose. St = stationary phase. See SI Figure 1 for uncropped scanned images. In all panels, error bars represent standard error; *p* values derived from two-tailed Student's t test



Extended Data Figure 5. Preferential utilization of undigested in ulin by B. ovatus and costs of inulin digestion by 04502/3

a, Time to mid-log (estimated at 50% maximal OD (OD50) by linear regression analysis) of Fig. 3d. **b**, Additional independent experiments for Fig. 3d. Each line represents n=1 sample per condition. **c**, Competition of *Bo* WT and *Bo* 04502/3 co-cultured in 0.5% inulin with trace (0.06%) amounts of fructose. n=2 cell culture biological replicates; **d**,**e**, Growth (**d**) and TLC analysis of conditioned media (**e**) of *Bo* WT and *Bo* 04502/3 cultured in 0.1% inulin with trace (0.06%) amounts of fructose. See SI Figure 1 for uncropped scanned images. For **d**, each panel is an independent experiment. Each condition is plotted as individual lines in each panel (n=2 cell culture biological replicates). In all panels, error bars represent standard error; *p* values derived from two-tailed Student's t test.



Extended Data Figure 6. A complex mouse microbiota differentially affects *B. ovatus* WT and 04502/3 pre-colonized gnotobiotic mice and analysis of cross feeding of the predominant murine Bacteroidales of the murine gut microbiota

a, Germ-free mice were monocolonized with *Bo* WT or 04502/3 and maintained on a diet supplemented with inulin as the sole polysaccharide and housed under gnotobiotic conditions for 2 weeks. Bacteria were enumerated from feces prior to gavage with intestinal microbiota of conventionally raised mice (n=5 mice, cell culture biological replicates). b, Growth of two dominant mouse microbiota Bacteroidales strains (*Parabacteroides goldsteinii* and a strain with 96% 16S rRNA gene identity to *B. thetaiotaomicron*) with inulin breakdown products derived from the conditioned media of *B. ovatus* grown in inulin (all inulin had been digested) or undigested inulin minimal media. Each data point is a different isolate of the indicated species from the ceca of the conventionally raised mice used for gavage (n=11 isolates for upper panel, n=6 for lower panel. In all panels, error bars represent standard error.



Extended Data Figure 7. *B. ovatus*, but not *B. fragilis* benefits from *B. vulgatus* in co-culture in inulin

a, **b**, Independent experiments for Fig. 4b,d. The left panel corresponds to starting culture with ~ 10^7 CFU *Bo* corresponding to starting culture of ~ 10^6 CFU *Bo* in Fig. 4d. The two right panels are a duplicate pair of experiments of starting CFU of 10^6 and 10^7 . In each panel n=2 biological replicates. **c**, Enumeration of *B. fragilis* (**c**) in monoculture or co-culture with *Bv* on defined inulin plates, n=3 cell culture biological replicates. Letters in parentheses refer to values correlating to color of line used for statistical analysis. In (**a**), for example, p (g,r) refers to comparison of values of green (g) and red (r) values at the time-point indicated. In (**b**) color of p value indicated. For all panels, error bars represent standard error; *p* values derived from two-tailed Student's t test.



Extended Data Figure 8. Secreted factors from Bo and isogenic mutants, but not B. fragilis (Bf), support Bv survival

a, Growth of Bv in conditioned media from Bo WT, 04502, 04502/3 or Bf grown in defined media with inulin as the sole carbohydrate and in inulin media. End time-point corresponded to peak growth of Bv in conditioned media derived from Bo WT. B. fragilis, which utilizes inulin^{1,3} but similar to Bo = 04502/3 does not liberate inulin breakdown products³, does not support the survival of Bv during co-culture. Left and right panels are independent experiments. Left panel; start, n=4 biological replicates; end n=2 biological replicates. Right panel; t0, n=4 cell culture biological replicates; t13 and 24, n=2 biological replicates. b, Growth of Bv in dialyzed (2 kD MW membrane) or undialyzed conditioned media from *Bo* 04502/3 grown in inulin, n=2 cell culture biological replicates. c, Gas chromatographic analysis of acetate, propionate and succinate in conditioned media during growth of Bo WT and 04502/3 in defined media with inulin as the sole carbohydrate. Other volatile and non-volatile substances (as listed in Methods) were undetectable, n=2 cell culture biological replicates, except 04502/3 stationary phase, n=1. **d**, Growth of Bv in defined medium with inulin as the sole carbohydrate with or without addition with 15 mM of acetate, propionate or succinate, t0, n=4 biological replicates; t13 and 24, n=2 biological replicates. CM = conditioned media, UD = undetected. For all panels, error bars represent standard error; p values derived from two-tailed Student's t test



Extended Data Figure 9. Spatial aspects of mutualism between *Bo* and *Bv* via cross feeding and interspecies cooperation *in vivo* via 04502/3

a, Independent experiments for Fig. 4c, n=4 biological replicates. b, Independent experiments for Fig. 4e, n=3 cell culture biological replicates. c, Scatter plot of experiment in Fig. 4f, n=3 cell culture biological replicates. d, Ratios of WT and 04502/3 in feces 21 days after co-colonization of three germ-free mice (n=3 mice biological replicates) on a diet of inulin as the sole dietary polysaccharide (pre-Bv) and then four days after introduction of Bv (post-Bv). Each panel shows the ratio pre- and post-Bv of an individual mouse. p values are Fisher exact test comparing the frequency of Bo WT and 04502/3 pre- and postcolonization with Bv for each individual mouse. At day 21, all mice were colonized with a higher ratio of the mutant (ranging to 86% in the mouse depicted in the lowest panel), with each mouse showing a statistically significant increase in the proportion of the WT after introduction of Bv. e, Growth of B. ovatus in 0.5% inulin (upper panel) or fructose (lower panel) in minimal media to which filter sterilized conditioned media from early log, OD_{600} matched growth of Bv or Bo in 0.125% fructose minimal media or fresh 0.125% fructose minimal media control was added at 1:1 ratio. In (e) numbers refer to p values (* <0.05, **<0.01, ***<0.001) of comparison of values of green (g), red (r) or black (b) values by unpaired, two-tailed student t test at the time-point indicated, n=2 cell culture biological replicates. For all panels, error bars represent standard error, for all panels except (\mathbf{d}) p values derived from two-tailed Student's t test.



Extended Data Figure 10. Schematic of forms of cooperativity via polysaccharide digestion among Bacteroidales

a, Limited cooperation. Privatization of extracellularly digested public goods by the individual performing the digestion leads to greater individual (dark arrow) than shared benefits (light arrow) as seen in *Bt* during growth on levan and amylopectin. **b**, Cooperation between species is seen between *Bo* and *Bv* during growth on inulin. Surface digestion of inulin by *Bo* creates breakdown products that it does not need to grow on inulin. Rather, inulin breakdown represents a dedicated cross-feeding system that provides benefits to *Bv*, with reciprocal fitness benefits to *Bo*.

Supplementary Material

Refer to Web version on PubMed Central for supplementary material.

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Figure 1. Direct and cooperative benefits of polysaccharide digestion by surface glycoside hydrolases (GH)

a, Polysaccharide utilization loci of *B. thetaiotaomicron* (*Bt*) for amylopectin and levan with the products or properties each gene encodes listed above and color coded. SPI or SPII = signal peptidase I or II cleavage site. **b**, **c**, Growth of *Bt* WT and surface GH mutants in media with amylopectin (n=2, cell culture biological replicates) (**b**) or levan (n=2, cell culture biological replicates) (**c**). See Extended Data Fig.1 for additional independent experiments. **d**,**e**, Growth of *Bt* WT and surface GH mutants in media with amylopectin (n=2, cell culture biological replicates) (**d**) or levan (n=2, cell culture biological replicates) (**d**) or levan (n=2, cell culture biological replicates) (**e**). See Extended Data Fig.1 for additional independent experiments. In all panels error bars represent standard error; *p* values derived from two-tailed Student's t test.





a, Predicted inulin utilization locus of *B. ovatus* (*Bo*). Gene designations shown are preceded by BACOVA_. The color coding of gene products is as in Fig.1. **b**, Thin layer chromatography (TLC) analysis of inulin defined medium incubated for the indicated times with purified His-tagged 04502, 04503 or 04502 and 04503, or vector control. See SI Figure 1 for uncropped scanned images. **c**, Growth of *Bo* WT and mutants in inulin defined medium. **d**, TLC analysis of conditioned media during the growth of *Bo* WT and mutants in 0.5% and 0.1% inulin. EL = early log, ML = mid log, LL = late log, St = stationary. **e**, Growth of *Bo* WT, 04502/3, 04504 (*susD* ortholog) and 04505 (*susC* ortholog) mutants in inulin defined medium. For **c**, **e**, each line represents n=1 sample per condition. See Extended Data Fig.2 for additional independent experiments. In all panels error bars represent standard error; *p* values derived from two-tailed Student's t test.



Figure 3. Cost of inulin digestion by surface glycoside hydrolases

a, Growth yield of *Bo* WT and mutants on inulin agarose plates. Each line represents n=1 sample per condition. See Extended Data Fig. 4 for additional independent experiments. **b**, Bacteria per gram feces of gnotobiotic mice seven days after monocolonization with *Bo* WT or 04502/3 on a polysaccharide free diet or supplemented with inulin (n=5 biological replicate mice per condition). **c**, Growth curves (upper and middle panels) and maximal OD₆₀₀ (bottom panel) of *Bo* WT or 04502/3 in minimal media with carbon sources as indicated. Each line represents n=1 sample per condition. See Extended Data Fig. 4 for

additional independent experiments. In the bottom panel, *p* values are displayed as paired (matched WT and 04502/3 within same experiment) followed by unpaired (all experiments) two-tailed Student's t test (n=7, biological replicates). **d**, Growth of *Bo* WT or mutants in inulin or a stoichiometric equivalent amount of inulin breakdown products after digestion with purified 04502 and 04503 enzymes (n=2, cell culture biological replicates); *, p < 0.05; ** p < 0.01; ***, p < 0.001; See Extended Data Fig. 5 for additional independent experiments. In all panels error bars represent standard error; *p* values derived from two-tailed Student's t test.





a, Germ-free mice were monocolonized with *Bo* WT or 04502/3 and maintained on a polysaccharide free diet supplemented with inulin and housed for 2 weeks. Mice were then gavaged with the cecal contents of conventionally raised mice. At day 5 post gavage, cecal contents were plated for enumeration of *Bo* (n=5 biological replicate mice per group). **b**, Enumeration of *Bv* in monoculture or co-culture with *Bo* WT or 04502 on defined inulin plates. α , p = 0.001 for number of *Bv* in *Bv/Bo* WT co-culture vs. *Bv* alone; ψ , p = 0.001

for number of Bv in Bv/Bo WT co-culture vs. Bv/04502 co-culture; a, p = 0.003 for number of Bv in Bv/ 04502 co-culture vs. Bv alone (n=2, cell culture biological replicates). See Extended Data Figs. 7 and 9 for additional independent experiments. c, (left) Photos of patches of Bv plated at varying distances around Bo WT or Bo 04502/3 or alone on inulin agarose plates. (right) Enumeration of Bv after five days of culture plated at the same distance to Bo WT or Bo 04502/3 or alone on inulin plates. See Extended Data Fig. 9 for additional independent experiments. d, Enumeration of Bo WT or 04502 in monoculture or co-culture with Bv on defined inulin plates. p values correlate to color of line/genotype used for statistical analysis. p value indicates comparison of monoculture and co-culture for the given condition at the time-point indicated. The benefit Bo receives from Bv is most robust when starting with fewer *Bo*. Depicted are starting CFU of $\sim 10^6$ *Bo*. (n=3, cell culture biological replicates at day 1,2; n=2, biological replicates at day 4). See Extended Data Fig. 7 for additional independent experiments and Extended Data Fig.7b for starting CFU of $\sim 10^7$ Bo. e, Ratios of WT and 04502/3 at the start and day 2 of culture when co-plated with or without Bv on inulin plates (n=3, cell culture biological replicates). See Extended Data Fig. 9 for additional independent experiments. f, Ratios of WT and 04502/3 in the inoculum (day 0) and in feces at various time points (days 4, 7, 11, 14, 18) post cocolonization of gnotobiotic mice (n=3 cell culture biological replicates) with Bo WT and

04502/3. Polysaccharide free diet was supplemented with inulin at day 7. Bv was introduced at day 14. A Fisher exact test comparing the frequency of Bo WT and 04502/3 pre- (day 14) and post- (day 18) colonization with Bv was significant with a p value of 0.0001 for each individual mouse. Bo CFU in feces were maximal after the switch to inulin diet and addition of Bv changed the abundance of Bo WT compared to 04502/3 but not total CFU of Bo. See Extended Data Fig. 9 for additional independent experiments. In all panels error bars represent standard error; p values displayed are derived from two-tailed Student's t test.