Enhancing nutritional niche and host defenses by modifying the gut microbiome

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Abstract

The gut microbiome is essential for processing complex food compounds and synthesizing nutrients that the host cannot digest or produce, respectively. New model systems are needed to study how the metabolic capacity provided by the gut microbiome impacts the nutritional status of the host, and to explore possibilities for altering host metabolic capacity via the microbiome. Here, we colonized the nematode Caenorhabditis elegans gut with cellulolytic bacteria that enabled C. elegans to utilize cellulose, an otherwise indigestible substrate, as a carbon source. Cellulolytic bacteria as a community component in the worm gut can also support additional bacterial species with specialized roles, which we demonstrate by using Lactobacillus plantarum to protect C. elegans against Salmonella enterica infection. This work shows that engineered microbiome communities can be used to endow host organisms with novel functions, such as the ability to utilize alternate nutrient sources or to better fight pathogenic bacteria.

Keywords: bacteria community; cellulose; gut microbiome; nutrition; pathogen

Subject Categories: Biotechnology & Synthetic Biology; Microbiology; Virology & Host Pathogen Interaction

DOI: 10.15252/msb.20209933 | Received 21 August 2020 | Revised 15 August 2022 | Accepted 14 October 2022

Mol Syst Biol. (2022) 18: e9933

Introduction

The gut microbiome is a complex community of microbes, the ecology of which is intimately tied to the physiology of its host organism (Sampson & Mazmanian, 2015; Barratt et al., 2017). These symbiotic microbes have been shown to contribute to the nutritional status and metabolism of the host (Shreiner et al., 2015). Understanding how various species of microorganisms constituting the microbiome affect their host is fundamental to deriving potential benefits from manipulating the microbiome.

Gut microbe metabolism affects the health of animal hosts in numerous ways, from the liberation of indigestible nutrients (Tremaroli & Bäckhed, 2012) to the modification of host-secreted bile acids (Rowland et al., 2018), to the synthesis of certain vitamins and neurotransmitters at significant levels (Sampson & Mazmanian, 2015). Moreover, metabolic interactions among members of the microbiota help maintain community structure, thereby, preventing pathogen invasion or other dysbioses (Shreiner et al., 2015).

Carbohydrates are important sources of energy for both microbes and animals (Tremaroli & Bäckhed, 2012). Simple sugars such as glucose and lactose can be easily absorbed by hosts, but more complex carbohydrates and plant polysaccharides, including cellulose, xylans, resistant starch, and inulin, are not as easily digested. The gut microbiome can confer caloric benefits upon its host by breaking down these ingested plant carbohydrates, which the host enzymes cannot digest (Rowland et al., 2018), as has been demonstrated for termites (Bourguignon et al., 2018) and ruminants (Clemmons et al., 2019). The gut microbiome can also evolve to adapt to new carbohydrate sources when hosts change their eating habits. For example, the Japanese, as a result of their diet, have intestinal...
microbiomes that have acquired enzymes for processing the algal carbohydrates in seaweed through horizontal gene transfer from marine bacteria (Hehemann et al., 2012). However, research into the use of non-native bacteria to perform nutritional functions for other animal hosts has been limited.

In this study, we engineered a functional microbiome in a simple animal gut to allow host utilization of a complex carbohydrate source. We used the nematode Caenorhabditis elegans as the animal host as it has been extensively used as a model system to elucidate mechanisms of interaction between prokaryotes and their hosts (Jones & Ashrafi, 2009; Zhang et al., 2017). Specifically, we colonized C. elegans with cellulolytic bacteria that break down cellulose in the gut, such that the released glucose is available as a nutrient for both C. elegans itself and the colonizing bacteria. To move this system toward higher community complexity, as occurs in a natural gut microbiome, we added an additional bacterial species with the specialized function of preventing infection by pathogens. Our work indicates that by assembling a functional community in the C. elegans gut, we can extend the nutrition processing and pathogen inhibition capacities of the animal host.

Results

Cellulolytic bacterial colonization and function in C. elegans gut

To introduce a novel carbon processing capability to C. elegans, we started by selecting cellulose-degrading microbes as potential intestinal colonists. To identify these functional bacteria members, we assembled a collection of bacteria native to the C. elegans gut (Dirksen et al., 2016; see Appendix Table S1 for a full list) and tested these species for their ability to utilize carboxymethyl cellulose (CMC) and other carbon sources by measuring bacterial optical density following in vitro incubation (Appendix Fig S1). Additionally, we checked the cellulose degradation capacity for each bacterial strain using a Congo Red assay (Teather & Wood, 1982; Houfani et al., 2017). As Congo Red stains glucose polymers, a halo surrounding the colony indicates cellulase activity (Fig 1A). Although the investigation of the functional capacity of the C. elegans native microbiome indicates that some rare members of this microbiome can degrade cellulose (Zimmermann et al., 2020), none of the native C. elegans microbiome isolates in this particular set of strains (Fig 1A) exhibited detectable cellulase activity, which reflects their lack of cellulose processing capability. To further identify cellulolytic bacteria, we screened a set of soil bacteria (Appendix Table S1) and identified Pseudomonas cellulosa and Bacillus subtilis as cellulase-active bacteria (Fig 1A). Of these strains, P. cellulosa produced the highest levels of reducing sugars, including glucose (Fig 1B), suggesting that this strain could potentially be useful as a nutritional endosymbiont for the C. elegans host, supplying nutrients that would otherwise be unavailable. After screening, we also kept one of the non-cellulolytic bacterial strains, Pseudomonas putida, as a negative control and for use in future engineering.

We next sought to determine whether a cellulolytic organism could colonize the worm intestine. C. elegans feed on bacteria, a certain percentage of which escape digestion and colonize the gut (Schulenburg & Félix, 2017; Vega & Gore, 2017). In these experiments, synchronized adult N2 worms were used for colonization to provide a genetically homogeneous population of mature hosts. We allowed the worms to be colonized by culturing them in liquid bacterial culture to ensure that all the worms experienced a uniform environment (Vega & Gore, 2017). Worms were then collected, washed, and mechanically disrupted to release the intestinal bacteria. These suspensions were then plated on agar to estimate the extent of bacterial colonization (Fig 1C). Naturally cellulolytic bacteria, P. cellulosa and B. subtilis, as well as the natively non-cellulolytic bacterium, Pseudomonas putida, which had been engineered to produce endoglucanase (see Materials and Methods), each achieved a colonization density of ~10^2–10^4 colony-forming units (CFU) per worm (Fig 1D). Colonization was stable for at least 2 days in the C. elegans gut. All of these bacterial species were therefore able to colonize the host intestine, albeit at different population sizes, at colonization levels comparable to those of native C. elegans gut strains (Appendix Figs S2 and S3).

To determine whether bacteria identified as being cellulolytic in vitro could hydrolyze complex sugars in the gut environment, we pre-colonized C. elegans with the cellulolytic strain P. cellulosa. This strain was chosen for its strong ability to degrade CMC and release reducing sugars, including glucose (Fig 1B and Appendix Fig S4). Since P. cellulosa grows to low densities in culture, we used Escherichia coli OP50 as a supplemental food source to ensure that worms had adequate nutrition during growth. We used OP50 because C. elegans is usually grown monoxenically in the laboratory using E. coli OP50 as a food source (Brenner, 1974). In these experiments, synchronized adult N2 worms were grown on solid plates instead of liquid media to allow efficient feeding on these bacteria and therefore increase the rate of colonization. As we intended to use fecundity as one measure of carbon uptake in these experiments and as C. elegans fecundity shows a strong response to starvation (adult reproductive diapause) when food is withdrawn in the mid- to late L4 stage, feeding and colonization on live bacteria were stopped at 46 h post-L1 on plates (Seidel & Kimble, 2011). Worms raised on mixed lawns of P. cellulosa and E. coli OP50 showed intestinal colonization by P. cellulosa (Fig 1D), indicating that this bacterial species is able to colonize the host. Pre-colonized worms were then incubated for 24 h in liquid media (M9 worm buffer) with: (i) no carbon source, (ii) cellulose (in the form of CMC), or (iii) glucose. After incubation, worms were washed and digested to check gut bacterial density. We found that M9 buffer with CMC as a carbon source enhanced the colonization density of P. cellulosa compared with no carbon source (Fig 1E). This experiment demonstrated that a cellulolytic bacterium, P. cellulosa, could utilize CMC as a carbon source in the C. elegans gut. Glucose as a carbon source further enhanced colonization density; this result was not surprising because free sugar monomers are an easier-to-access carbon source than CMC, which can be utilized only after it has been hydrolyzed to glucose.

Intestinal cellulolytic bacteria allow host utilization of cellulose

Next, we sought to determine whether colonization by cellulolytic bacteria would allow the host to use the otherwise indigestible cellulose carbon source. C. elegans is able to use simple carbohydrates, including glucose (Lu & Goetsch, 1993), but does not produce enzymes that degrade cellulose. We hypothesized that if cellulolytic
organisms in the gut could liberate carbon in a form that is available and metabolizable by the host, colonization of the gut with a cellulolytic microbiome would provide this host with the ability to use cellulose as a growth substrate. Again, we chose *P. cellulosa* as the colonizing bacteria. Three approaches, described below, were used to investigate whether *C. elegans* utilized glucose released by gut bacteria.

First, we used radionucleotide (14C) incorporation as a direct measurement of cellulose-derived carbon incorporation into worm biomass. Briefly, adult wild-type worms were colonized with *E. coli* OP50 (non-cellulose degrader) or *P. cellulosa*, then incubated in S medium with heat-killed *E. coli* OP50 and trace 14C labeled CMC (radioabeled cellulose purified from *Arabidopsis thaliana*; Faik *et al.*, 2002) for 24 h to allow bacterial degradation of the substrate to proceed. After incubation on the radionucleotide substrate, worms were washed and treated with an antibiotic cocktail for 24 h to remove bacteria, in order to eliminate the effects of bacterial biomass on scintillation counts and observe the incorporation of cellulose-derived carbon into the germ-free worm biomass. A high-throughput antibiotic susceptibility screen of the bacterial strains (including bacteria native to the *C. elegans* gut and the soil bacteria listed in Appendix Table S1) was conducted to determine an appropriate antibiotic cocktail to eliminate these species from the *C. elegans* gut (Appendix Fig S5). Clearance by the antibiotic cocktail was tested by imaging worms fed with fluorescently labeled *Pseudomonas putida* (Appendix Fig S6). After clearing the bacterial biomass, worms were mechanically digested in batches (200 worms per sample), and 14C incorporation was measured using scintillation counts. 14C incorporation was significantly higher for *P. cellulosa* colonized worms than for *E. coli* colonized worms, before and after clearance with antibiotics (Fig 2A), demonstrating that cellulolytic bacteria in the *C. elegans* gut had hydrolyzed 14C labeled CMC and that the hydrolyzed glucose had been taken up by the host, as well as by the microbes.
Second, we confirmed that incorporation of carbon from microbiome-digested CMC affected host metabolism by using a fluorescent reporter of host nutritional status (Lee et al., 2015). We first validated that fat-7p::GFP, which is expressed in the intestinal cells of C. elegans (Nomura et al., 2010; Ma et al., 2015), was a reliable reporter for carbon uptake by the host, and determined an optimal glucose concentration of 0.1–0.2% (w/v) for these assays (Appendix Fig S7). Young adult wild-type worms (N2 strain, 46 h after L1 instar) were transferred from plates of live OP50 + E. coli OP50 alone, produced more larvae per adult worm when CMC was provided in the medium (Fig 2C and Appendix Fig S6), again

fat-7p::GFP (Fig 2B), indicating an increase in nutrients available to the host.

Third, we measured the benefits of carbon incorporation into the new host biomass via larval output, to determine whether colonization by cellulose-digesting bacteria could improve the reproductive fitness of the host. Larval output in the worm is a function of nutritional status and can be initiated by adding nutrients after starvation at a specific point in the young adult-to-mature adult transition (Seidel & Kimble, 2011). We, therefore, transferred worms from solid media colonization plates to liquid culture supplemented with glucose or CMC 46 h after plating L1 larvae, to capture worms at the vulnerable point in this transition. Here, worms were placed into individual wells of a 384-well plate, to allow enumeration of larval yield for single adults. After 48 h in liquid media with a carbon source, the number of larvae produced by individual bacteria was counted. Worms colonized with P. cellulosa, but not with E. coli OP50 alone, produced more larvae per adult worm when CMC was provided in the medium (Fig 2C and Appendix Fig S6), again
indicating that colonization by the cellulolytic bacterial species allows the worms to benefit from CMC as a carbon source made available by these resident bacteria.

**Bacteria community protects C. elegans against pathogen invasion of the gut**

Alongside these efforts, we sought to determine whether adding a co-colonizing microbial species would produce a community with additional functionality for the host. As an alternative to adding an additional cellulose-degrading bacterium, we sought to take advantage of the probiotic properties of *Lactobacillus plantarum*, as lactic acid bacteria have previously been shown to have probiotic effects in their hosts, including protection against pathogens (Ikeda et al., 2007; Bucková et al., 2018; Mao et al., 2018). We hypothesized that adding this probiotic to the gut community could prevent disruption of the community by an intestinal pathogen (here, *Salmonella enterica* LT2). To determine the effect of *L. plantarum* on pathogen colonization, worms were pre-colonized with *P. cellulosa* on solid media as previously described, with and without the addition of *L. plantarum* (10⁵ CFU/plate). After colonization, worms were exposed to *S. enterica* LT2 (10⁸ CFU/ml) for 1 h in liquid culture (Fig 3A). Following this initial exposure, worms were washed to remove external pathogens and incubated for 36 h to allow infection to progress (Fig 3B). We found that *P. cellulosa* and *L. plantarum*, alone or in combination, modestly suppressed pathogen proliferation relative to the control condition (colonization by *E. coli* OP50), but there was no significant difference between the *P. cellulosa* only and the *P. cellulosa* and *L. plantarum* combination conditions without a carbon source (Fig 3B).

In the presence of CMC as carbon source, however, *P. cellulosa* and *L. plantarum* in combination suppressed the proliferation of *Salmonella* by 10-fold compared with what was seen in worms colonized by either one of these strains alone (Fig 3B and Appendix Fig S8). These data suggest that the progression of infection in *C. elegans* can be affected by the resident microbiome. Moreover, the presence of CMC as a carbon source enhanced the community’s capability to suppress pathogens, demonstrating that provision of a specific substrate for the cellulose degrader improved overall community performance. Our hypothesis is that simple sugars released by *P. cellulosa* promote *L. plantarum*’s capability to fight against the pathogen. Although there have been efforts using single-species probiotics to fight against pathogen colonization (Mao et al., 2018; Kissoyan et al., 2019), our work shows that a simple microbial community can suppress pathogens in the *C. elegans* intestine. The strains in a community, compared with the single-strain system, appear to benefit from the complex mix of available nutrients and acquire an enhanced ability to suppress organisms that are pathogenic to the host.

**Discussion**

In this study, we developed a model of metabolic microbe-host interactions and directly demonstrated that colonization with heterologous bacteria enables *C. elegans* to digest and incorporate carbon from previously indigestible long-chain carbohydrates. In addition to isotopic measurement of carbon incorporation, we demonstrate direct benefits both to the host, with increased larval yield, and to other gut species, which in turn may provide protective effects against pathogens (Fig 4).

We developed assays to quantify the effects of an exogenously introduced microbiome on the nutritional status of *C. elegans*. Data sets collected toward this end include high-throughput in *vivo* carbon source growth screens to identify bacteria capable of degrading complex carbon sources, and in *vivo* screens to identify antibiotic cocktails for the removal of colonized bacteria from the *C. elegans* gut. These assays can support the use of these organisms as a convenient model for future studies of microbiota-host interactions.

The simple systems described here have yielded insights about the intricacies of inter-species interactions in the potential pitfalls of designing a novel microbiome. In our experiments, the use of a facultatively interacting taxon (*L. plantarum*) allowed us to add a new functional capacity (resilience against pathogen invasion) by capitalizing on the particular properties of this facultative species. This new function of the community was boosted when we exposed it to CMC, which can be utilized by cellulolytic *P. cellulosa* in the community, demonstrating that distinct bacteria can interact to combat pathogens via cellulose utilization. These results, consistent with other research (Miller et al., 2018) in this area, indicate that understanding interactions between bacterial strains within an *in vivo* host environment will be critically important in selecting novel colonists for a host microbial ecosystem. Community assembly starting from modular orthogonal interactions offers a promising approach for designing microbiome communities (Chu & Adler, 2015; Friedman et al., 2017; Kong et al., 2018; Medlock et al., 2018; Venturelli et al., 2018; Lopez et al., 2020). Our results are also consistent with current thought on the use of “prebiotic” nutritional adjuvants to support the engraftment and function of specific beneficial microbes in the gut. Recent studies that utilize porphyran as an orthogonal metabolite to enable strain engraftment (Kearney et al., 2018; Shepherd et al., 2018) are likewise consistent with this view.

Engineering the gut microbiome could permit hosts to digest new food sources or enhance the efficiency of current nutrition processing capabilities. With rapid progress in enzyme evolution (Nielsen & Keasling, 2016) and microbiome engineering (Johns et al., 2016) techniques, facilitated by developments with model host organisms, engineering the gut microbiome could accelerate...
A  
Colonize → Expose to *Salmonella* → Disrupt & plate

B  
**Salmonella Inhibition**

<table>
<thead>
<tr>
<th>Media</th>
<th>None</th>
<th>CMC</th>
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<tbody>
<tr>
<td>OP50</td>
<td>100,000</td>
<td>3E-2</td>
</tr>
<tr>
<td><em>P. cellulosa</em></td>
<td>10,000</td>
<td>10,000</td>
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<tr>
<td><em>L. plantarum</em></td>
<td>1,000</td>
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<tr>
<td><em>P. cellulosa + L. plantarum</em></td>
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Figure 3.
nutrient processing in livestock or expand diets to include entirely new food sources.

**Materials and Methods**

**Bacterial species, worm strains, and culturing**

Bacterial species used in this study are listed in Appendix Table S1, along with species-specific growth media. In absence of specific culture requirements, bacterial cultures were routinely inoculated from glycerol stock and grown in LB media in a 30 or 37 \(^\circ\)C shaking incubator or a benchtop shaker (23 \(^\circ\)C) for 24–48 h before use.

Unless otherwise stated, N2 wild-type *C. elegans* (Caenorhabditis Genetic Center) were used in these experiments. Cultivation of all worms used here was performed at 25 \(^\circ\)C on NGM plates seeded with *E. coli* OP50. Synchronization of worms for these experiments was performed using standard protocols (Stiernagle, 2006); briefly, worms were cultivated on NGM to obtain large numbers of gravid adults hermaphrodites, and eggs were isolated from these plates via bleach-NaOH lysis of adults. Eggs were rinsed 5–6× in M9 worm buffer to remove the hypochlorite solution and allowed to hatch out in 5 ml of M9 worm buffer overnight, after which the synchronized and starved L1s could be used for experiments. When reproducitively sterile adult worms were desired, L1 larvae were plated onto NGM + 1 mM IPTG + 50 \(\mu\)g/ml plates seeded with *E. coli* expressing *pos-1* RNAi from a plasmid (Rual et al., 2004); worms grown to adulthood on these plates produce inviable eggs and are therefore reproductively sterile.

**Cellulolytic bacteria screening**

A Congo Red assay was used to determine cellulose hydrolyzing capability of bacteria. 1% CMC plates seeded with bacteria were cultured overnight at 30\(^\circ\)C. The plates were then flooded with 0.1% Congo Red for 15–20 min and subsequently rinsed with 1 M NaCl solution. A halo formed around the colony indicates cellulase activity. For resting cell assays, cells in liquid culture were first washed once with buffer containing 50 mM Tris–HCl (pH 8.0) and suspended in 1% CMC (carboxymethyl cellulose) from Sigma. Samples were collected after 1 h and immediately mixed with 0.5 ml of DNS reagents (10 g/l dinitrosalicylic acid, 10 g/l sodium hydroxide, 2 g/l phenol, 0.5 g/l sodium sulfite). After incubation at 95\(^\circ\)C for 10 min, 1 ml of 40% Rochelle salts was added to the color before measuring the absorbance at 575 nm (Sun & Chen, 2016).

**Bacteria colonization and analysis**

Washed adult synchronized worms were resuspended in S medium and moved in 500 \(\mu\)l aliquots to 15 ml culture tubes with a final concentration of ~1,000 worms/ml. In total, 500 \(\mu\)l of bacterial suspension at 2× desired concentration was added to each culture tubes. Culture tubes were incubated with shaking at 300 rpm at...
25°C. After 24 h, colonized worms were washed 3× with M9 worm buffer +0.1% Triton X-100 and 2× with M9 worm buffer to remove external bacteria. Worms were subsequently chilled to 4°C for 15 min to stop peristalsis and treated for 10 min with a 1:2,000 solution of commercial bleach to remove any remaining external bacteria, then washed 2× in M9 worm buffer +0.1% Triton X-100 to remove bleach prior to disruption.

For manual disruption, worms were transferred to 3 ml M9 worm buffer +1% Triton X-100 in a small (35 cm) petri dish (Fisher Scientific). Individual worms were pipetted out and transferred to 0.5 ml clear microtubes (Kimble Kontes) for manual disruption with a motorized pestle (Kimble Kontes Pellet Pestle with blue disposable pestle tips, Fisher Scientific, all digestions in 20 μl of buffer). Magnification was provided by a magnifying visor (Magni-Focuser Hands Free Binocular Magnifier 3.5×). After disruption, tubes were centrifuged 2 min at 16,099 × g to collect all material, and the resulting pellet was resuspended in 180 μl M9 worm buffer (final volume 200 μl) before transfer to 96-well plates for serial dilution in 1× phospho-buffered saline (PBS). Ten-, 100- and 1,000-fold dilution samples were plated on LB or ATCC Medium 2720 plates to check the colonization density.

Colonized bacteria benefit from CMC

Synchronized L1 worms from the N2 wild-type lineage were plated on solid agar on mixed lawns of E. coli OP50 and P. cellulosa, and allowed to grow to adulthood. Following this colonization, day 0 adult worms were separated into two aliquots. One went through disruption for colonization density as described above. The second aliquot was divided between wells of a 24-well plate with S medium, S medium with 0.1% glucose and S medium with 0.1% CMC (final volume 1 ml/well), covered with a BreathEasy gas-permeable membrane, and incubated with shaking at 200 rpm at 25°C for 24 h. Individual samples went through disruption and plating to check on nutritional effects on colonization density.

Measurement of 14C-carbon incorporation

Each gut colonization type was done in triplicate using 200 N2 synchronized worms. Samples were incubated at room temperature, with gentle mixing, in 1 ml M9 Worm buffer containing 0.5% Carboxy-methyl cellulose and 1 μCi/ml 14C-Uniformly labeled cellulose from A. thaliana for 72 h. Antibiotic cocktails (100 μg/ml ciprofloxacin and 500 μg/ml carbenicillin) were provided at 48 h. Scintillation counts were performed after washing the worms with M9 buffer containing 0.1% TritonX. Washing of the worms was repeated until the scintillation counts in the wash supernatant reached background levels. Scintillation counts were performed using liquid scintillation analyzer (PerkinElmer Tri-Carb 2910 TR) in a high-efficiency LSC-cocktail (PerkinElmer Ultima Gold).

Antibiotics cocktails for gut bacteria elimination

A high-throughput antibiotic susceptibility screen over our collection of soil bacteria was conducted with incubating bacteria antibiotics pooled and checked on bacteria density afterward. Once an effective antibiotic cocktail was identified, the best conditions to kill bacteria in the worm gut were determined. Worms were colonized with YFP-labeled Pseudomonas citronellolis (GmR) and treated with antibiotic cocktails (ciprofloxacin 100 μg/ml + carbenicillin 500 μg/ml). After 24 h, fluorescent gut-associated colonies were checked; disruption and plating of intestinal contents indicated if all live bacteria were cleared out from the worm intestine.

Larval yield assay

In these experiments, we chose to measure fecundity in individual worms rather than in groups of hermaphrodites on shared plates, to avoid statistical issues associated with taking average measurements on long-tailed distributions and thereby to increase our power to detect differences between groups. N2 wild-type worms were synchronized according to standard protocols (Stiernagle, 2006), and L1 worms were transferred to 10 cm NGM plates containing OP50 only, OP50 + P. cellulosa (50 μl 10⁻⁸ bacterial suspension from 48 h culture in 10 ml ATCC 2720, grown at room temperature with shaking), OP50 + B. subtilis (prepared identically to P. cellulosa using LB as growth medium). Worms were incubated on plates at 25°C for 48 h, then washed off plates using M9 worm buffer +0.1% Triton X-100, rinsed 3× to remove the bulk of extracellular bacteria, and picked as single worms into individual wells of a 384-well plate containing 25 μl S medium +0.25× heat-killed OP50 (from 50× stock, cells concentrated 50× from stationary phase) + 25 μg/ml gentamycin (to prevent external growth of bacteria) ± carbon (glucose or carboxymethylcellulose). For each combination of bacteria and carbon source, 24 worms were assayed in each experiment. After 48 h, larvae were counted in individual wells using 4× magnification and white light illumination on a Ti-E inverted microscope. This time point was chosen because at 24 h, there were still a large number of zero larval counts across conditions, as many worms had not yet started laying eggs and early eggs were still in the process of hatching, and dehydration in the plate began to become an issue by 72 h.

Assay for nutritional status using an integrated fluorescent reporter

We used transgenic worms expressing the fat-7p::GFP reporter fluorescent reporter as an indicator of host nutritional status (Lee et al., 2015). fat-7p::GFP worms were synchronized according to standard protocols and grown to day 0 adults on pos-1 plates to prevent reproduction, then transferred to 6 cm NGM plates containing live E. coli OP50 P. cellulosa, P. cellulosa was grown at 25°C, in 5 ml ATCC 2720 then pelleted at 9,056 g for 2 min and resuspended in S medium to concentrate the bacteria prior to seeding NGM plates with pre-existing OP50 lawns (50 μl of each suspension per plate). Worms were allowed to feed on these plates for 24 h at 25°C to allow bacterial colonization.

Nutritional status was read on a BioSorter large object sorter using the 250 micron nozzle. Adult worms were transferred from plates to individual wells of a 24-well plate in S medium with heat-killed E. coli (to provide nitrogen and other nutrients) + gentamycin 10 μg/ml (to prevent growth of bacteria outside the worm) ± glucose or CMC at the indicated concentrations, covered with a BreathEasy gas-permeable membrane, and incubated for 24 h at 25°C with shaking at 200 rpm. After incubation, worms were washed to remove heat-killed bacteria, and total GFP fluorescence per worm was measured on the BioSorter. Worms were gated based on
extinction and time of flight (TOF) to isolate full-sized adult worms in these data, due to imperfect penetrance of the pos-1 reproductively sterile phenotype in some experiments.

**High-throughput carbon source utilization screen**

Eighteen species were inoculated from frozen glycerol stock into 2 ml of their preferred rich medium (see Appendix Table S1), and grown in a shaking incubator at 30°C and 300 rpm for 48 h. Fast-growing species exceeding OD ~1.0 in the first 24 h of this period were diluted 1:1,000 into fresh rich media to grow for the remaining 24 h. All cultures were then centrifuged at 1,370 g for 5 min, then resuspended in an M9 media formulation lacking any carbon source. OD600 was measured in a 96-well plate on a SpectraMax plate reader, and culture densities were normalized to OD 1.0. These cultures were diluted 1:20 into deep 96-well plates containing 400 µl of 12 different M9 media formulations, each with a different carbon source at 0.5% concentration (arabinose, cellobiose, carboxymethyl cellulose, ethanol, glucose, lactose, raffinose, soluble starch, sucrose, xylan hemicellulose, and xylose, as well as a no-carbon control). Cultures were grown at 30°C and 900 rpm for 48 h. OD600 was measured again, and the M9 no-carbon control cultures were used to subtract out apparent growth that may have occurred from residual rich media or absorbance changes independent of carbon utilization. Triplicate assays were performed on different days, with pairwise correlations of $r^2 \geq 0.87$ between days.

**Antibiotic susceptibility screen**

Four species were grown from frozen glycerol stocks for 48 h in preferred rich media using the same protocol as in the carbon utilization screens. Cultures were centrifuged at 3,500 rpm, and resuspended in fresh rich medium and normalized to an OD600 of 1.0 by diluting with additional rich medium. Normalized cultures were next diluted 1:100 into deep 96-well plates containing 400 µl rich media dosed with one of seven antibiotics (carbenicillin, chloramphenicol, ciprofloxacin, kanamycin, nalidixic acid, streptomycin, and tetracycline, and an ethanol control) at 1x, 5x, and 10x working concentrations. Cultures were grown at 30°C and 900 rpm for 24 h, then OD600 was measured on a SpectraMax M5 plate reader to determine bacteriostatic potential. To determine whether each drug had proved bactericidal, five µl spots of each culture were grown on permissive nutrient agar. Colony counts were binned by order of magnitude (i.e., 0, 1–10, 11–100, lawn of > 100 colonies).

**Engineered cellulolytic *P. putida***

To engineer non-cellulolytic *P. putida* to cellulolytic bacteria, *P. putida* was transformed with pET24a with ice-nucleation protein fusion with endoglucanase Cel5A assembled through Gibson assembly. Induction of Cel5A surface display was achieved with 0.1 mM Isopropyl β-D-1-thiogalactopyranoside (IPTG) at OD 1 of bacteria culture. An overnight (16 h) induction at 16°C was conducted before bacteria were collected, and reducing sugar assay from the bacteria was checked. We used *P. putida* because it is a *Pseudomonas* strain, so it serves as a good negative control strain for our cellulolytic bacteria, *P. cellulosa*. Endoglucanase was chosen for these experiments because it hydrolyzes cellulose by cutting randomly on the cellulose chain. Ice-nucleation protein is a bacterial surface display tag that has been used to demonstrate the translocation of enzymes across the *P. putida* surface (Yang *et al.*, 2007). *P. putida* was transformed with the pET24a-INP-Endoglucanase plasmid, which had been sequenced prior to transformation, and enzyme activity was checked after protein expression. Since endoglucanase only works by being exposed to the substrate cellulose, the activity of the enzyme should come from the surface-displayed enzyme.

**Clearance of bacteria from the gut using antibiotics**

In order to drive clearance of bacteria from the gut, we screened antibiotics to develop a cocktail with efficacy against all species in our consortium. *In vitro* tests identified a combination of 50 µg/ml carbenicillin and 10 µg/ml ciprofloxacin that should have proven bactericidal to a broad range of species (Appendix Fig S5). However, during *in-vivo* experiments, we found that 10x higher concentrations of antibiotics were needed to effectively kill bacteria residing in the gut (Appendix Fig S5).

**Data availability**

Data supporting the findings of this study are available in the Appendix.
References


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