Developing a *Bacteroides* System for Function-Based Screening of DNA from the Human Gut Microbiome

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**ABSTRACT** Functional metagenomics is a powerful method that allows the isolation of genes whose role may not have been predicted from DNA sequence. In this approach, first, environmental DNA is cloned to generate metagenomic libraries that are maintained in *Escherichia coli*, and second, the cloned DNA is screened for activities of interest. Typically, functional screens are carried out using *E. coli* as a surrogate host, although there likely exist barriers to gene expression, such as lack of recognition of native promoters. Here, we describe efforts to develop *Bacteroides thetaiotaomicron* as a surrogate host for screening metagenomic DNA from the human gut. We construct a *B. thetaiotaomicron*-compatible fosmid cloning vector, generate a fosmid clone library using DNA from the human gut, and show successful functional complementation of a *B. thetaiotaomicron* glycan utilization mutant. Though we were unable to retrieve the physical fosmid after complementation, we used genome sequencing to identify the complementing genes derived from the human gut microbiome. Our results demonstrate that the use of *B. thetaiotaomicron* to express metagenomic DNA is promising, but they also exemplify the challenges that can be encountered in the development of new surrogate hosts for functional screening.

**IMPORTANCE** Human gut microbiome research has been supported by advances in DNA sequencing that make it possible to obtain gigabases of sequence data from metagenomes but is limited by a lack of knowledge of gene function that leads to incomplete annotation of these data sets. There is a need for the development of methods that can provide experimental data regarding microbial gene function. Functional metagenomics is one such method, but functional screens are often carried out using hosts that may not be able to express the bulk of the environmental DNA being screened. We expand the range of current screening hosts and demonstrate that human gut-derived metagenomic libraries can be introduced into the gut microbe *Bacteroides thetaiotaomicron* to identify genes based on activity screening. Our results support the continuing development of genetically tractable systems to obtain information about gene function.

**KEYWORDS** *Bacteroides thetaiotaomicron*, anaerobic sulfatase maturing enzyme, chondroitin sulfate utilization, fosmid library, functional metagenomics, functional screening, gut microbiota, human gut microbiome, metagenomic library, surrogate host

As the microbes that live within the human body are implicated in a growing number of human disease states, there has been corresponding growing interest in the role of the different organisms that comprise the gut microbiota. This interest has been supported by advances in DNA sequencing technology that allow the generation of large metagenome sequence data sets, and yet, study of the human gut microbiota is hampered by a lack of knowledge of gene function that makes annotation of those data sets incomplete; for example, approximately 50% of genes identified in Human

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Microbiome Project (HMP) stool samples lacked a functional assignment using standard (GO, EC, or homology-based) annotation methods (1). As research on the microbiota continues, there will be an increased need for effective methods that can provide knowledge of microbial gene function (2). Functional metagenomics is an approach in which environmental DNA is cloned to generate metagenomic libraries that are maintained in *Escherichia coli* (Fig. 1A) and the cloned DNA is screened en masse for specific functions of interest. This powerful approach allows the isolation of genes whose roles may not have been predicted based on their DNA sequence alone (3), but crucially, the method is dependent on the ability to express cloned metagenomic DNA in a surrogate host.

The human gut microbiota is dominated by members of the *Firmicutes* and *Bacte-
roidetes phyla, with the Bacteroides genus often the most abundant (4). Though *E. coli*, a member of the Gammaproteobacteria, is often used as a surrogate host to screen gut-derived DNA, there is evidence of a barrier to the expression of Bacteroides genes at the level of transcription (5), due to differences in promoter consensus recognition between the Bacteroidetes primary sigma factor and that of *E. coli* (6). The development of a more suitable host will likely improve hit rates from activity screens (7), and in particular, a Bacteroides host would offer an array of potentially selectable glycan utilization phenotypes (8, 9) that are less feasible in *E. coli*. The human gut symbiont *Bacteroides thetaiotaomicron* is a natural choice as a surrogate host to screen gut metagenomic DNA, given that molecular genetic methods for this organism are reasonably well developed. Here, we describe our attempt to develop *B. thetaiotaomicron* for functional metagenomics, through the construction of a *B. thetaiotaomicron*-compatible fosmid cloning vector, generation of a human gut metagenomic library, and screening of the library to achieve functional complementation of a *B. thetaiotaomicron* glycan utilization mutant.

RESULTS

Construction of a Bacteroides-compatible human gut metagenomic library. To be able to screen a library in a *B. thetaiotaomicron* host, the library must be constructed using a vector that is able to replicate in *B. thetaiotaomicron*. To construct a suitable library cloning vector, we chose to build on pCC1FOS, a commercial fosmid vector that has been widely used for constructing metagenomic libraries from diverse environments (10). Although cosmid vectors for *B. thetaiotaomicron* have been constructed in the past using pBR322 and RSF1010 origins (11, 12), we desired the potentially increased insert stability offered by a single-copy F-based vector, as cloned Bacteroides DNA may be unstable in *E. coli* (12) and instability may be exacerbated by maintenance at a higher copy number (13). We chose to use a self-replicating rather than an integrating vector because the former allows fosmid DNA to be isolated from *B. thetaiotaomicron* cells by plasmid minipreparation, facilitating DNA sequencing of the complementing insert. A fosmid vector was especially desirable for two reasons: (i) lambda packaging to generate fosmid clone libraries is very efficient and (ii) large-fragment libraries would be suitable for capturing the polysaccharide utilization loci of Bacteroides, which may contain over 20 genes (9).

pCC1FOS was modified by the addition of an origin of transfer (oriT) to allow plasmid conjugation from *E. coli* to *B. thetaiotaomicron*, as well as plasmid replication elements (repA) and an erythromycin-selectable marker (ermF) for *B. thetaiotaomicron*. The constructed *B. thetaiotaomicron*-compatible vector was designated pKL13 (Fig. 1B) and used to generate a human gut metagenomic library, called CLGM3, that contained over 100,000 unique clones with an estimated average insert size of 26 ± 10 kb. To assess the level of transfer from *E. coli* to *B. thetaiotaomicron*, a triparental conjugation was carried out (Fig. 1C), resulting in conjugation efficiencies (relative to recipient) of 2.6 × 10⁻² for the vector alone and 1.1 × 10⁻² for the CLGM3 metagenomic library. Though transfer into *B. thetaiotaomicron* was not as efficient as that for other surrogate hosts, such as the legume symbiont *Sinorhizobium meliloti* (14), it was sufficient for initial attempts at functional complementation.

Proof-of-principle functional complementation of a *B. thetaiotaomicron anSME* mutant. As a host for a proof-of-principle functional complementation, we chose a *B. thetaiotaomicron ΔanSME* mutant, also called the ΔchuR mutant (15). The 1,245-bp *chuR/anSME* gene (BT_0238) was first identified through transposon mutagenesis as a regulator of chondroitin sulfate and heparin utilization (16). Knocking out this gene renders *B. thetaiotaomicron* unable to grow on the glycan chondroitin sulfate or heparin as a sole carbon source. It was later characterized as an anaerobic sulfatase maturating enzyme: the breakdown of these glycans by *B. thetaiotaomicron* requires the action of sulfatase enzymes that must be modified posttranslationally by the product of the *anSME* gene (17); without this modification, the sulfatases are inactive. The mutant phenotype being dependent on the single *anSME* gene, as well as the clean
phenotype of the ΔanSME mutant on chondroitin sulfate (Fig. 2A), made it a good candidate for functional complementation.

To screen the CLGM3 metagenomic library for genes able to complement the anSME mutant, the library was conjugated from E. coli EPI300 into the B. thetaiotaomicron ΔanSME strain, selecting on minimal medium with chondroitin sulfate as the sole carbon source. As a positive control, a B. thetaiotaomicron genomic library (constructed using B. thetaiotaomicron VPI 5482 DNA; called BT3) was screened simultaneously. Colonies arising on the selective medium were streak purified to confirm the restored phenotype, providing evidence that the mutant’s ability to grow on chondroitin sulfate was restored by complementation with clones from the B. thetaiotaomicron genomic library or the gut metagenomic library (Fig. 2B or C, respectively).

Possible fosmid clone recombination into the B. thetaiotaomicron host genome. Plasmid DNA can be prepared from B. thetaiotaomicron cultures using standard alkaline lysis, and we confirmed that plasmid preparations of empty vector DNA from B. thetaiotaomicron can be used to successfully transform E. coli EPI300. We applied this same strategy to obtain the fosmid DNA from cultures of the complemented anSME mutant; however, plasmid minipreparations followed by transformation of EPI300 yielded no transformants for the samples, indicating that there was no fosmid DNA isolated from these cultures despite the restored ability to use chondroitin sulfate as the sole carbon source. We hypothesized that the anSME-complementing fosmid DNA may have recombined into the host genome, an unfortunate scenario as the screening of pooled-clone metagenomic libraries hinges on being able to retrieve the complementing DNA for sequence analysis.

To test this hypothesis, we isolated genomic DNA from 5 clones from the BT3 complementation and 7 clones from the CLGM3 complementation and used the DNA as the template in a PCR to test for the presence of the fosmid’s oriT sequence. As suspected, the genomic DNA preparations from all clones were positive for the oriT (Fig. 3A). We also confirmed the anSME mutant background; this mutant strain carries a deletion of the ~1,200-bp anSME open reading frame (ORF), and primers designed to 300 bp upstream and 300 bp downstream of the ORF amplify only 600 bp from the mutant versus ~1,800 bp from the wild type. As expected, genomic DNA preparations from all of the BT3- and CLGM3-complemented clones carried the mutant anSME genomic context (Fig. 3B).

We next asked whether the metagenomic fosmid clones were carrying DNA from B. thetaiotaomicron or closely related species, which may explain the propensity for homologous recombination. To answer this, we performed PCR for the anSME ORF using primers based on the B. thetaiotaomicron VPI 5482 anSME sequence, which would likely amplify only exact or close matches to B. thetaiotaomicron. All clones from the
B. thetaiotaomicron BT3 library produced PCR products (Fig. 3C), which was expected as this library was constructed using B. thetaiotaomicron DNA. From the CLGM3 metagenomic library, all but clone 2 showed amplification, confirming our suspicion that most complementing clones were probably closely related to the host. We purified and Sanger sequenced the PCR products, finding that 5 of the 6 metagenomic anSME sequences were an exact match to VPI 5482. The last of the 6 products, from CLGM3 clone 5, was not identical but highly similar (see Fig. 5).

**FIG 3**  PCR analysis of genomic DNA isolated from anSME-complemented B. thetaiotaomicron clones. As controls, genomic DNAs from the wild-type (WT) B. thetaiotaomicron VPI 5482 and the ΔanSME mutant were included, as well as plasmid DNA for the pKL13 fosmid. PCR was carried out to amplify the oriT sequence of the fosmid vector backbone (~800 bp) (A), a product corresponding to 300 bp upstream and 300 bp downstream of the anSME ORF (~1,800 bp for VPI 5482 and 600 bp for ΔanSME) (B), the anSME ORF (~1,200 bp for VPI 5482) (asterisks indicate products confirmed as identical to B. thetaiotaomicron VPI 5482 by Sanger sequencing) (C), and the anSME ORF plus 300 bp downstream (~1,500 bp for VPI 5482) (D).
The result of the PCR for the anSME ORF and flanking region (Fig. 3B) was surprising in that the BT3 library clones did not exhibit both the 600-bp and 1,800-bp bands—the former from the B. thetaiotaomicron ΔanSME background and the latter from the complementing fosmid DNA carrying the B. thetaiotaomicron anSME gene. To determine if the smaller product may be preferentially amplified in PCR, we designed primers to the anSME locus such that the smaller PCR product was not possible, amplifying the anSME ORF plus 300 bp downstream (Fig. 3D). The ~1,500-bp product of this PCR confirmed that indeed the complementing anSME locus was present in the clones carrying VPI 5482 DNA (BT3 clones) or closely related metagenomic DNA (CLGM3 clones).

**Genome sequencing of B. thetaiotaomicron ΔanSME complemented with metagenomic DNA.** We were interested in further characterizing CLGM3 clone 2 and clone 5; these derived from the metagenomic library and appeared to be carrying DNA distinct from VPI 5482. We decided to carry out genome sequencing in the hope that we could (i) gain insight into the sequence similarity between fosmid clone and host that may contribute to recombination and (ii) identify the complementing anSME gene from clone 2, which we were unable to retrieve using PCR (Fig. 3C). After sequencing, we first aligned reads to the B. thetaiotaomicron VPI 5482 genome; second, we de novo assembled reads to identify pKL13 vector DNA and the complementing anSME genes.

By mapping reads from clone 2 back to VPI 5482, we were able to confirm the ΔanSME mutant background through the zero read depth observed at the cleanly deleted anSME ORF (Fig. 4A). Assembly of reads resulted in a fosmid-sized 45-kb contig that included the pKL13 vector backbone and the complementing anSME gene (Fig. 4B). The insert carried by the fosmid had 99% nucleotide identity to Bacteroides vulgatus ATCC 8482 and was sufficiently different from VPI 5482 to enable essentially complete assembly; this dissimilarity also validates the lack of an amplicon from PCR using primers that were designed against the VPI 5482 anSME sequence (Fig. 3C and D). Interestingly, the ends of this contig share sequence similarity with regions that flank a gene annotated as a transposase in the host genome, suggesting a possible integration mechanism and/or locus, although there exist other regions along the contig of lower similarity to the host genome (see Fig. S1 in the supplemental material).

When we mapped reads from clone 5 back to VPI 5482, we found that there were reads with high-enough similarity to map to the VPI 5482 anSME locus. However, the higher read depth at that locus as well as the low identity observed indicated a foreign source of DNA; perhaps most tellingly, the stretch of low identity was consistent with the size of a typical fosmid insert (Fig. 4C). The high sequence similarity between the fosmid insert and host genome likely also contributed to a fragmented de novo assembly: a fosmid-sized contig was not assembled, although the pKL13 vector and anSME sequence were found on smaller contigs (Fig. 4D). The sequences adjacent to the vector (on both ends) as well as the contig carrying the anSME gene exhibited high nucleotide identity to B. thetaiotaomicron VPI 5482 (84 to 99%). Due to the fragmented nature of assembly and the high similarity between metagenomic DNA and host DNA, it was difficult to speculate on possible integration loci for this clone.

**Gut-derived anSME genes identified by functional complementation.** Using functional complementation, we were able to identify two anSME gene sequences from a gut metagenomic library (Fig. 4B and D). Comparison of the translated sequences to the VPI 5482 anSME 415-residue protein sequence revealed a number of changes at the amino acid level (Fig. 5), none of which were in the three conserved cysteine clusters thought to be involved in the ability of the anSME gene product to mature sulfatase enzymes (17).

The anSME gene of clone 2 was identical to B. vulgatus ATCC 8482, whereas the anSME gene of clone 5 appeared to be novel though nearly identical to B. thetaiotaomicron VPI 5482. The identification of anSME genes from a human gut metagenomic library that are different in sequence from the B. thetaiotaomicron VPI 5482 host used for screening indicates that functional screening of metagenomic libraries using B. thetaiotaomi-
**B. theta ΔanSME carrying CLGM3 clone 2**

(A and C) Mean read depth per 5,000 bp after mapping reads to VPI 5482 genome. (Pullout) Read depth and percent identity per base pair at the anSME locus; red lines delineate anSME open reading frame.

(B and D) Relevant contigs from de novo assembly; pKL13 vector sequence and complementing anSME gene are indicated in red. B. theta, B. thetaiotaomicron.

**FIG 4** Genome sequencing and de novo assembly results for B. thetaiotaomicron ΔanSME carrying CLGM3 clone 2 and clone 5. (A and C) Mean read depth per 5,000 bp after mapping reads to VPI 5482 genome. (Pullout) Read depth and percent identity per base pair at the anSME locus; red lines delineate anSME open reading frame. (B and D) Relevant contigs from de novo assembly; pKL13 vector sequence and complementing anSME gene are indicated in red. B. theta, B. thetaiotaomicron.
**DISCUSSION**

In this study, *B. thetaiotaomicron* was chosen as a host for screening gut-derived metagenomic DNA because it is anticipated to be able to express a greater fraction of the cloned DNA than would *E. coli*. Previous work on the *B. thetaiotaomicron* 16S rRNA gene operon showed that while the *B. thetaiotaomicron* ribosome-binding site was recognized by *E. coli*, the barrier to gene expression was due to lack of promoter recognition (5). Though there have been reports in the literature of functional screens in *E. coli* yielding positive clones carrying *Bacteroides*-derived DNA (18–22), these hits may be due to spurious transcription of foreign DNA in *E. coli* rather than transcription from native *Bacteroides* promoters.

To develop a system that may be better suited for screening gut metagenomic libraries, we generated *B. thetaiotaomicron*-compatible fosmid libraries with which we demonstrated successful functional complementation of a *B. thetaiotaomicron* anSME mutant. Analysis of the complemented mutants, however, indicated that recombination may have occurred between the host genome and the DNA carried on the fosmid clones, suggesting high sequence similarity between the host genome and complementing DNA. Consistent with this, we found that nearly all complementing anSME genes from the metagenomic library were exact or close matches to *B. thetaiotaomicron*, which may not be surprising given that *B. thetaiotaomicron* is often a dominant species in the human distal gut (23). We carried out full genome sequencing on 2 of the 7 metagenomic clones to obtain the complementing anSME genes and found both deriving from *Bacteroides* species. It is possible that recombination may be due to instability of the pKL13 vector backbone; however, we did confirm that empty vector could be introduced into and reisolated from *B. thetaiotaomicron*.

To demonstrate that *B. thetaiotaomicron* can be used as a screening host, we chose a ΔanSME mutant for the simplicity of its phenotype being dependent on a single gene. In retrospect, this choice may have limited the diversity of hits obtained. Although functional metagenomics is an approach that can uncover novel genes, here we found that the anSME-complementing genes obtained from the gut were either identical or closely related to *B. thetaiotaomicron* VPI 5482. To enrich for more diverse hits, it may be useful to choose a target with a larger known sequence space, including multigene operons; indeed, the strength of the functional metagenomic approach is that the large inserts of cos-based vectors can carry entire operons involved in polysaccharide utili-
zation. Here, we showed that an ~45-kb fosmid carrying human gut metagenomic DNA can be successfully conjugated into *B. thetaiotaomicron*, demonstrating that large, multigene operons characteristic of polysaccharide utilization loci can be transferred for functional complementation.

Although the propensity for homologous recombination presents difficulties for the screening of pooled-clone metagenomic libraries, it is not a barrier to the functional metagenomics approach. One possible solution is to use arrayed libraries in which clones are stored and conjugated into the recipient individually, enabling clone tracking and eliminating the need for clone DNA retrieval. Individual conjugations may also be advantageous over en masse conjugations if the relatively low conjugation efficiency is a bottleneck for transferring large clone libraries into a *B. thetaiotaomicron* recipient. Screening of arrayed libraries containing hundreds of thousands of clones, however, often requires specialized equipment to achieve the necessary throughput. Another possible solution may be to use a recombination-deficient *B. thetaiotaomicron* strain, although a *B. thetaiotaomicron* recA mutant has been reported to have increased sensitivity to oxygen (24).

Our results show that the development of a *B. thetaiotaomicron* system for functional screening was not as straightforward as hoped; however, the genetic tractability of *B. thetaiotaomicron* and the generation of genetic tools with which to manipulate it (25) provide support for continued efforts. In addition, a more quantitative comparison between *E. coli* and *B. thetaiotaomicron* as expression hosts for metagenomic library screening would be valuable. Although the development of a *B. thetaiotaomicron* expression host may offer an advantage over *E. coli* for screening DNA derived from the *Bacteroidetes*, *B. thetaiotaomicron* would not be ideal for screening DNA from other phyla present in the gut, particularly *Firmicutes*. A comprehensive screening strategy will likely require the use of multiple expression hosts. Indeed, expanding the range of screening hosts will be important for the functional metagenomics field and for the characterization of microbial genes with currently unknown function. The fosmid vector that we have described in this report, and the strategy for functional complementation via conjugal transfer, will provide a strong foundation for further refinements.

**MATERIALS AND METHODS**

**Bacterial strains, plasmids, and oligonucleotides.** The bacterial strains and plasmids used in this study are provided in Table 1. Oligonucleotide names and sequences are provided in Table 2.

**Culture of *E. coli***. *E. coli* was routinely cultured in LB broth or agar at 37°C with appropriate antibiotics. Antibiotics used in solid medium were chloramphenicol (10 μg/ml), ampicillin (100 μg/ml), kanamycin (25 μg/ml), and tetracycline (10 μg/ml); antibiotic concentrations were halved for liquid culture.

**Culture of *B. thetaiotaomicron***. *B. thetaiotaomicron* was routinely cultured in broth using brain heart infusion medium (BD Biosciences), supplemented with 1.2 μM histidine, 1.9 μM hematin, 1 μg/ml menadione, and 500 μg/ml cysteine (BHI†). *B. thetaiotaomicron* was cultured in liquid using the pyrogallol method (26): after inoculation, a cotton ball was inserted into the mouth of the culture tube and set aflame; after the flame was extinguished, 200 μl of 20% (wt/vol) NaCO₃ and 200 μl of 35% (wt/vol) pyrogallol were added to the cotton, and the tube was immediately plugged with a rubber stopper. Cultures of *B. thetaiotaomicron* were incubated at 37°C, without shaking. Typically, resazurin was added to the liquid medium as an indicator of oxidizing/reducing conditions (1 μg/ml final concentration). *B. thetaiotaomicron* was cultured on complex medium agar plates using brain heart infusion medium, supplemented with 10% defibrinated horse blood (Bio-Media Unlimited) (BHIH). Minimal medium agar with chondroitin sulfate as the sole carbon source was prepared by dissolving chondroitin sulfate (Sigma-Aldrich or Toronto Research Chemicals) completely in distilled water (for a 5-g/liter final concentration) and autoclaving with agar, followed by adding salts and supplements as previously described (27) as well as trace elements (1,000× stock solution; concentrations per liter: 0.247 g H₃BO₃, 0.1 g CuSO₄·5H₂O, 0.338 g MnSO₄·H₂O, 0.282 g ZnSO₄·7H₂O, 0.056 g CoSO₄·7H₂O, and 0.048 g Na₂MoO₄·2H₂O) and the appropriate antibiotics. Antibiotics used in solid medium were gentamicin (200 μg/ml), kanamycin (200 μg/ml), nalidixic acid (25 μg/ml), and erythromycin (10 μg/ml); antibiotic concentrations were halved for liquid culture. Agar plates were incubated in airtight containers with GasPak EZ anaerobe sachets (BD Biosciences).

**Construction of pKL13 fosmid vector.** The fosmid cloning vector pCC1FOS was modified for screening in a *B. thetaiotaomicron* host. Briefly, the RK2 oriT fragment was amplified from pJC8 using primers KL12/KL13 and RK2 oriT fragment was cloned into the intermediate vector pJET1.2 (Thermo Fisher).

**Table 1**

| Bacterial Strains, Plasmids, and Oligonucleotides |
|-----------------------------------------------|
| *E. coli*                                      |
| *B. thetaiotaomicron*                         |
| *B. thetaiotaomicron* recA mutant             |

**Table 2**

| Oligonucleotide | Name          | Description                              |
|-----------------|---------------|------------------------------------------|
| Primer          | KL12/KL13     | Suitable for screening in *E. coli*      |
| Primer          | KL14/KL15     | Suitable for screening in *B. thetaiotaomicron* |

**Figure 1**

- Pyrogallol method
- Minimal medium with chondroitin sulfate
- BHIH agar plates
**TABLE 1** Bacterial strains and plasmids used in this study

| Strain or plasmid | Description | Source or reference |
|-------------------|-------------|---------------------|
| **Strains** E. coli | F– mcrA Δ(mrr-hsdRMS-mcrBC) ϕ80lacZAM15 ΔlacX74 recA1 endA1 araD139 Δ(ara leu)7697 galU galK rpsL (S'^r) nupG traA dfrF | Epicentre |
| | HB101 F – mcrB mrr hsdS20(r– mR–) recA13 leuB6 ara-14 proA2 lacY1 galK2 xyl-5 mtl-1 rpsL20 (S'^r) glnV44 | 40 |
| B. thetaiotaomicron | VPI 5482 B. thetaiotaomicron type strain; the VPI 5482 type strain is the same as ATCC 29148 | 23 |
| | BtUW24 Derivative of VPI 5482 with deletion of BT-2275 (tdk); used in conjunction with pExchange-tdk to construct deletion mutants | 30 |
| | BtUW25 Derivative of Δtdk with deletion of BT-0238 (ansME); unable to grow on chondroitin sulfate as sole carbon source | 15 |
| | BtUW4 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from BT3 genomic library designated BT3_chuR2 | This study |
| | BtUW7 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from BT3 genomic library designated BT3_chuR5 | This study |
| | BtUW8 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from BT3 genomic library designated BT3_chuR6 | This study |
| | BtUW11 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from BT3 genomic library designated BT3_chuR9 | This study |
| | BtUW12 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from BT3 genomic library designated BT3_chuR10 | This study |
| | BtUW14 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from CLGM3 metagenomic library designated CLGM3_chuR1 | This study |
| | BtUW15 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from CLGM3 metagenomic library designated CLGM3_chuR2 (clone 2) | This study |
| | BtUW16 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from CLGM3 metagenomic library designated CLGM3_chuR3 | This study |
| | BtUW17 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from CLGM3 metagenomic library designated CLGM3_chuR4 | This study |
| | BtUW18 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from CLGM3 metagenomic library designated CLGM3_chuR5 (clone 5) | This study |
| | BtUW20 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from CLGM3 metagenomic library designated CLGM3_chuR8 | This study |
| | BtUW21 B. thetaiotaomicron BtUW25 carrying anSME-complementing clone from CLGM3 metagenomic library designated CLGM3_chuR9 | This study |
| **Plasmids** | pRK2013 Mobilizer plasmid; ColE1 replication origin and Km' | 41 |
| | pAFD1 E. coli-Bacteroides shuttle vector with pUC replication origin | 42 |
| | pJC8 Cosmid vector with RK2 replication origin; NCBI accession no. KC149513 | 29 |
| | pJET1.2 Commercial vector for PCR product cloning; NCBI accession no. EF694056 | 43 |
| | pCC1FOS Commercial fosmid vector; NCBI accession no. EU140751 | Epicentre |
| | pKL13 Derivative of pCC1FOS; ermT and repA for selection and replication in Bacteroides, respectively; oriT for conjugation from E. coli into B. thetaiotaomicron; NCBI accession no. KU746975 | This study |

Fisher, and then digested and subcloned into the EcoRI site. The ~4-kb ermF–repA fragment was sequenced at the Centre for Applied Genomics (Toronto, Canada) to compile the complete sequence for the constructed vector pKL13, using primers KL14, KL16, KL33, KL42, KL43, KL45, and KL46 (Table 2). The vector was further modified to include transcriptional terminators (TTs) that flank the Eco72I cloning site to reduce insert-borne transcription, with both terminators from E. coli MG1655; the TT proximal to ermF incorporates the ilvGEDA terminator and the TT proximal to the RK2 oriT incorporates the repB T1 terminator (28). Finally, pKL13 contains a stuffer in the Eco72I site that aids in complete digestion of the restriction enzyme site.

**Construction of genomic and metagenomic fosmid libraries.** The metagenomic library was constructed using DNA extracted from a human fecal sample pooled from seven volunteers, obtained with clearance from the Office of Research Ethics of the University of Waterloo. The metagenomic library was designated CLGM3, and the genomic library constructed using genomic DNA from *B. thetaiotaomicron* VPI 5482 was designated BT3. Library construction was based on methods described previously (29). Briefly, DNA was extracted from either feces or a pure culture of *B. thetaiotaomicron* VPI 5482 and size selected (~40 to 70 kb) by pulsed-field gel electrophoresis. The insert DNA was electroeluted from the gel fragment, end repaired, and purified for ligation. The fosmid vector pKL13 was prepared by Eco72I digestion followed by dephosphorylation. The insert and vector were ligated, and the ligation products were packaged into lambda phage heads using Gigapack III XL packaging extract (Stratagene). The phage were used to transduce EPI300, and transductants were recovered on LB supplemented with...
TABLE 2 Oligonucleotides used in this study

| Oligonucleotide | Purpose                  | Sequence                       |
|-----------------|--------------------------|--------------------------------|
| KL12            | F primer to amplify R2K orT from pJC8, with HindIII adapter | CCTAAGCTTTGAGCTTTGCTGTTCGGG    |
| KL13            | R primer to amplify R2K orT from pJC8, with HindIII adapter | CCTAAGCTTTGAGCTTTGCTGTTCGGG    |
| KL14            | F primer to amplify ermF-repA fragment from pAFD1, with EcoRI adapter | CCTGAATTCAATACACAGCGGTACACAGCGGC    |
| KL15            | R primer to amplify ermF-repA fragment from pAFD1, with EcoRI adapter | CCTGAATTCAATACACAGCGGTACACAGCGGC    |
| KL16            | Primer walking ermF-repA fragment | GTTCAACAAAGACTGTCGTTTTCAATAGC  |
| KL33            | Primer walking ermF-repA fragment | CAGGCTAGGCAAGCCTGGTTCTTCAAAATAGC  |
| KL42            | Primer walking ermF-repA fragment | GGAAGCTCTCAAAATTTCTTAAATACAC  |
| KL43            | Primer walking ermF-repA fragment | CAAGCCCGTACGCCGTCGCGCGGTGTGGG  |
| KL45            | Primer walking ermF-repA fragment | AACAGAACAAGCGGTATTAAAGGACTTG  |
| KL46            | Primer walking ermF-repA fragment | GTCAAGCAAACCAAGGTAGTATTTATATCC  |
| KL61            | F primer for B. thetaiotaomicron anSME ORF | ATGAAAGCCAAACACTTTATGACACCTTATGGC  |
| KL62            | R primer for B. thetaiotaomicron anSME ORF | TTAATAATCTTCATTCTTTTTAAGCTTCGTTTGTGTTTC  |
| KL63            | F primer for 300 bp upstream of B. thetaiotaomicron anSME ORF | TCTCCATCTCCTCCCTAAGGTCTTCATATACAGAGACAGACACCGCCGCTAC  |
| KL65            | R primer for 300 bp downstream of B. thetaiotaomicron anSME ORF | TAAACCAGCTGATGTGTAGTGACGAATCAAGG  |
| KL67            | Sequence anSME PCR product from clone CLGM3_chu5R | AACGGGACGACGTCAGCGTCTTCTCCACC  |
| KL69            | Sequence anSME PCR product from clone CLGM3_chu5R | TCTATTTGCGTCAGCGGAGAAGTGC  |

chloramphenicol (10 μg/ml). Colonies were counted to estimate library size and then resuspended, pooled, aliquotted to generate the CLGM3 and BT3 library stocks, and stored at –80°C.

Triparental conjugation from E. coli to B. thetaiotaomicron. The triparental conjugation protocol was adapted from a biparental protocol (30). Matings were carried out using 5 ml of each of the donor, mobilizer, and recipient strains. The E. coli donor and mobilizer were mobilized in 5 ml LB supplemented with the appropriate antibiotics, and the B. thetaiotaomicron recipient was cultured in 5 ml BHI + ; all were grown to an optical density at 600 nm (OD600) of ~0.4 (Spectronic Spec 20 D). Cells were pelleted by centrifugation at 7,000 × g at room temperature for 5 min. The supernatant was removed, and the cells were resuspended in either BHI + or 1× Bt salts [per liter: 13.6 g KH₂PO₄, 0.875 g NaCl, 1.125 g (NH₄)₂SO₄; pH 7.2 (27)]. Donor, mobilizer, and recipient were mixed in a final volume of 1 ml, and the mixture was swirled evenly over the surface of a BHI plate. The plate was dried for several minutes in a laminar flow hood and then incubated aerobically overnight with the agar side down. Overnight mating lawns were scraped and resuspended in 2 ml BHI + or 1× Bt salts. Typically, serial 10-fold dilutions were made from 10⁻¹ to 10⁻¹, and 100 μl of each dilution was plated on BHIH supplemented with appropriate antibiotics to select for transconjugants; typically, kanamycin or nalidixic acid was used to select against E. coli and erythromycin was used to select for the vector. If the mating was plated on minimal medium, then the mating resuspension was washed to remove complex medium components by three repetitions of centrifugation and resuspension in 1× Bt salts.

Genomic DNA minipreparation of B. thetaiotaomicron. The minipreparation protocol is based on the method described by Charles and Nester (31). Briefly, B. thetaiotaomicron was cultured in 10 ml of liquid medium with the appropriate antibiotics, and the cell pellets were recovered after centrifugation at 7,000 × g for 5 min at room temperature. Cells were resuspended in 400 μl buffer (10 mM Tris [pH 8.0], 25 mM EDTA), 50 μl 5 M NaCl and 10 μl 10-mg/ml RNase A were added, and the tube was inverted several times. Twenty microliters of 1% SDS was added, and the sample was incubated at 65°C for 1 to 2 h. Two hundred sixty microliters of 7.5 M ammonium acetate was added, and the sample was incubated on ice for 20 min to precipitate proteins. The mixture was centrifuged at 21,000 × g for 20 min, the supernatant was decanted into a new microcentrifuge tube, and the mixture was extracted with chloroform in a 1:1 volume. The DNA was precipitated with 800 μl isopropanol and pelleted by centrifugation at 21,000 × g for 5 min. The pellet was washed with 100 μl 70% ethanol and centrifuged at 21,000 × g for 1 min, the supernatant was removed, and the pellet was allowed to dry. Finally, the pellet was added to lyophilization to dissolve the DNA. PCR analysis. Genomic DNA was isolated from the B. thetaiotaomicron clones carrying anSME-complementing fosmid DNA and used as the template. Taq-based 2× PCR master mix (Thermo Fisher) was used according to the manufacturer’s recommendations, with the exception that RNase A was added to remove RNA contamination (25-μg/ml final concentration). The touchdown PCR protocol used was 95°C for 3 min; 11 cycles of 95°C for 30 s, 60°C for 30 s (decrease of 1°C per cycle), and 72°C for 1 min/kb; 20 cycles of 95°C for 30 s, 50°C for 30 s, and 72°C for 1 min/kb; and 72°C for 5 min. PCR products chosen for Sanger sequencing were gel extracted, dissolved in binding buffer (140 mM 2-[N-morpholino]ethanesulfonic acid (MES)-NaOH [pH 7.0], 20 mM EDTA, 5.5 M guanidine isothiocyanate [33]) at 65°C for 5 min, applied to a silica spin column (BioBasic), washed several times (10 mM Tris-HCl [pH 7.5], 80% ethanol), and eluted in TE buffer. Sanger sequencing was completed at the Centre for Applied Genomics (Toronto, Canada).

Genome sequencing and analysis. Genome sequence data were generated on an Illumina MiSeq platform using 250-base paired-end sequencing. Sequence data were aligned to the B. thetaiotaomicron VPI 5482 reference genome (NCBI accession no. NC_004663.1 for the chromosome and NC_004703.1 for
the plasmids) using Bowtie 2 version 2.2.6 (34). Sequence data were de novo assembled using SPAdes version 3.8.0 (35), and functional annotations were obtained for contigs of interest using RAST (36). Regions of similarity between host and clone DNA were identified using Mauve (37). Data analyses were performed in R, including packages Rsamtools, Gviz, ape, and genoPlotR.

**Data availability.** Raw sequence data have been deposited at the NCBI Sequence Read Archive under accession numbers SRX3141910 (CLGM3 clone 2) and SRX3141914 (CLGM3 clone 5). Sequence data and other data may be accessed online at http://www.cm2bl.org/~data, including the raw data, alignment files, genome assemblies, and sequences of the anSME ORFs and ermF-repA fragment, as well as ab1 files from Sanger sequencing. The sequence of the constructed *B. thetaiotaomicron*-compatible vector, designated pKL13, may be found under NCBI accession no. KU746975. The metagenomic library designated CLGM3 may be found at NCBI BioSample accession no. SAMN04505233, and the genomic library constructed using genomic DNA from *B. thetaiotaomicron* VPI 5482, designated BT3, may be found at NCBI BioSample accession no. SAMN04505228.

**SUPPLEMENTAL MATERIAL**
Supplemental material for this article may be found at https://doi.org/10.1128/mSystems.00195-17.

**FIG S1**, EPS file, 0.6 MB.

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K.N.L. and T.C.C. conceived the ideas. K.N.L., E.C.M., and T.C.C. designed the experiments. K.N.L. performed the experiments, analyzed the data, made the figures, and wrote the manuscript. E.C.M. and T.C.C. provided constructive criticism, revised the manuscript, and provided reagents and materials.

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