

UCSF

UC San Francisco Previously Published Works

Title

Muribaculaceae Genomes Assembled from Metagenomes Suggest Genetic Drivers of Differential Response to Acarbose Treatment in Mice

Permalink

<https://escholarship.org/uc/item/22k027hs>

Journal

mSphere, 6(6)

ISSN

1556-6811

Authors

Smith, Byron J

Miller, Richard A

Schmidt, Thomas M

Publication Date

2021-12-22

DOI

10.1128/msphere.00851-21

Copyright Information

This work is made available under the terms of a Creative Commons Attribution License, available at <https://creativecommons.org/licenses/by/4.0/>

Peer reviewed



Muribaculaceae Genomes Assembled from Metagenomes Suggest Genetic Drivers of Differential Response to Acarbose Treatment in Mice

 Byron J. Smith,^a  Richard A. Miller,^b  Thomas M. Schmidt^{c,d}

^aThe Gladstone Institute of Data Science & Biotechnology, San Francisco, California, USA

^bDepartment of Pathology and Geriatrics Center, University of Michigan, Ann Arbor, Michigan, USA

^cDepartment of Ecology & Evolutionary Biology, University of Michigan, Ann Arbor, Michigan, USA

^dDepartment of Internal Medicine, University of Michigan, Ann Arbor, Michigan, USA

ABSTRACT The drug acarbose is used to treat diabetes and, by inhibiting α -amylase in the small intestine, increases the amount of starch entering the lower digestive tract. This results in changes to the composition of the microbiota and their fermentation products. Acarbose also increases longevity in mice, an effect that has been correlated with increased production of the short-chain fatty acids propionate and butyrate. In experiments replicated across three study sites, two distantly related species in the bacterial family *Muribaculaceae* were dramatically more abundant in acarbose-treated mice, distinguishing these responders from other members of the family. Bacteria in the family *Muribaculaceae* are predicted to produce propionate as a fermentation end product and are abundant and diverse in the guts of mice, although few isolates are available. We reconstructed genomes from metagenomes (MAGs) for nine populations of *Muribaculaceae* to examine factors that distinguish species that respond positively to acarbose. We found two closely related MAGs (B1A and B1B) from one responsive species that both contain a polysaccharide utilization locus with a predicted extracellular α -amylase. These genomes also shared a periplasmic neopullulanase with another, distantly related MAG (B2) representative of the only other responsive species. This gene differentiated these three MAGs from MAGs representative of nonresponding species. Differential gene content in B1A and B1B may be associated with the inconsistent response of this species to acarbose across study sites. This work demonstrates the utility of culture-free genomics for inferring the ecological roles of gut bacteria, including their response to pharmaceutical perturbations.

IMPORTANCE The drug acarbose is used to treat diabetes by preventing the breakdown of starch in the small intestine, resulting in dramatic changes in the abundance of some members of the gut microbiome and its fermentation products. In mice, several of the bacteria that respond most positively are classified in the family *Muribaculaceae*, members of which produce propionate as a primary fermentation product. Propionate has been associated with gut health and increased longevity in mice. We found that genomes of the most responsive *Muribaculaceae* showed signs of specialization for starch fermentation, presumably providing them a competitive advantage in the large intestine of animals consuming acarbose. Comparisons among genomes enhance existing models for the ecological niches occupied by members of this family. In addition, genes encoding one type of enzyme known to participate in starch breakdown were found in all three genomes from responding species but none of the other genomes.

KEYWORDS competition, gut microbiome, longevity

Editor Garret Suen, University of Wisconsin—Madison

Copyright © 2021 Smith et al. This is an open-access article distributed under the terms of the [Creative Commons Attribution 4.0 International license](https://creativecommons.org/licenses/by/4.0/).

Address correspondence to Thomas M. Schmidt, schmidt@umich.edu.

The authors declare no conflict of interest.

Received 30 October 2021

Accepted 16 November 2021

Published 1 December 2021

The mammalian gut microbiome is a complex ecological system that influences energy balance (1), pathogen resistance (2), and inflammation (3), among other processes with importance to host health. Understanding how the bacterial inhabitants of the gut respond to pharmaceutical and dietary perturbations is a major step in developing a predictive framework for microbiome-based therapies. Acarbose (ACA) is an α -glucosidase inhibitor prescribed for the treatment of type 2 diabetes mellitus because it reduces the absorption of glucose from starch in the small intestine (4). In rats, ACA has been shown to increase the amount of starch entering the lower digestive system after a meal (5). ACA treatment also changes the composition of the gut microbiota and its fermentation products in many rodents (5–14). Interestingly, long-term treatment with ACA has been shown to substantially increase longevity in male mice and, to a lesser extent, in females (15–17).

Previously we found that the relative abundance of multiple bacterial species as well as the concentrations of propionate and butyrate respond to long-term treatment with ACA (14). That study was notable in being replicated across three sites: The University of Michigan (UM) in Ann Arbor, The University of Texas Health Science Center at San Antonio (UT), and The Jackson Laboratory (TJL) in Bar Harbor, Maine. At UM and TJL, one highly abundant bacterial species was enriched nearly 4-fold in ACA-treated mice. This species, defined at a 97% identity threshold of the 16S rRNA gene V4 region and designated OTU-1, was classified as a member of the family *Muribaculaceae* in the order *Bacteroidales*. OTU-1 was also present and abundant at UT but was not significantly more abundant in ACA-treated mice relative to controls. Instead, a different *Muribaculaceae* species, designated OTU-4, was found to be highly abundant and enriched 4-fold in ACA-treated mice but was nearly absent at UM and TJL. Other *Muribaculaceae* were also identified as being among the most abundant members of the mouse gut microbiota across the three sites, although none of these were found to be enriched during ACA treatment.

Members of the family *Muribaculaceae*, previously referred to as family S24-7 after an early clone (GenBank accession no. [AJ400263.1](#)) (18), or sometimes as “*Candidatus* Homeothermaceae” (19), have only recently been isolated (20–22) despite being a common and abundant inhabitant of the mammalian gut, especially in mice (19, 21). Studies using culture-free methods suggest that the *Muribaculaceae* specialize in the fermentation of complex polysaccharides (19, 21), much like members of the genus *Bacteroides*, which is also a member of the order *Bacteroidales*. Genomic analysis has also suggested that the capacity for propionate production is widespread in the family (19).

Recently, techniques have been developed to reconstruct genomes of uncultivated members of bacterial communities (23, 24). Based on 30 such metagenome assembled genomes (MAGs) they reconstructed, Ormerod and colleagues (19) proposed that the *Muribaculaceae* fall into three distinct carbohydrate utilization guilds, which they describe as specialists on α -glucans, plant glycans, and host glycans, respectively. While it is reasonable to expect that α -glucan specialists would benefit the most from the large influx of starch to the gut resulting from ACA treatment, this prediction has not been tested, and physiological inferences based on the genome content of members of the family have been largely divorced from biological observations.

Experimental perturbations of complex microbial communities present an opportunity to observe ecological features of many bacterial taxa without cultivated members and generate hypotheses about their physiology. Given the observed, dramatically increased relative abundance of OTU-1 and OTU-4 (here referred to as responders) in mice treated with ACA, we hypothesize that these species are capable of robust growth on starch, while the other *Muribaculaceae* found in the study (nonresponders) lack the genomic features necessary for the utilization of polysaccharides that reach the colon in greater quantities following ACA treatment. Alternatively, responders may be resistant to the inhibitory effects of ACA or benefit from elevated levels of intermediate starch degradation products. Since isolates of the *Muribaculaceae* strains in these mice are not available for characterization, we took a comparative genomic approach to explore their functional potential.

Most of the research on the genomic components of polysaccharide degradation in Gram-negative bacteria has been carried out in the genus *Bacteroides*, particularly *B. thetaiotaomicron* (25). Starch utilization in *B. thetaiotaomicron* is dependent on an ensemble of eight proteins, SusRABCDEFG, that enable recognition, binding, hydrolysis, and import of starch and related polysaccharides (26). Homologs of SusC and SusD characterize all known polysaccharide utilization systems in this clade (27), are located in Sus-like genomic regions known as polysaccharide utilization loci (PULs), and are widespread in the phylum *Bacteroidetes* (28). The molecular range of these systems is determined by the carbohydrate-active enzymes and structural proteins they encode, based on the specificity of glycoside hydrolase (GH) and carbohydrate binding module (CBM) domains, which have been extensively cataloged in the CAZy and dbCAN databases (29–32).

Here, MAGs from the feces of mice at UT and UM are analyzed in the context of previously generated MAGs and cultivar genomes to explore two closely related questions about the niche of OTU-1 and OTU-4 in the lower digestive system. First, why do these species each increase in relative abundance with ACA treatment, while other species of *Muribaculaceae* do not? Second, why is the response of OTU-1 site specific? Despite similar patterns of abundance at their respective sites, the two responding species seem to be only distantly related, sharing just 90% of nucleotides in their 16S rRNA gene V4 hypervariable region (14). We nonetheless find genomic evidence that OTU-1 and OTU-4 occupy overlapping niches, specializing in the degradation of α -glucans, a role not held by the other *Muribaculaceae* described in this study. In addition, we identify two distinct genomic variants of OTU-1, referred to as B1A and B1B, which are differentially distributed between UM and UT and have functionally relevant differences in gene content.

Reconstructing genomes from metagenomes allows for the comparison of the functional potential of *Muribaculaceae* at UM and UT. This work demonstrates the utility of culture-free genomics to understand the potential ecological roles of these key members of the mouse gut microbial community and explore several hypotheses that may explain differences in the distribution and response of these bacteria to ACA treatment. Hypotheses derived from this analysis provide a foundation for future physiological studies in recently obtained cultivars. While a large fraction of host-associated bacterial species are without isolated representatives (33), let alone characterized (34), combining experimental data from complex communities with the analysis of reconstructed genomes provides a powerful tool for expanding understanding to these understudied taxa.

RESULTS

Recovered population genomes are of high quality and resemble other *Muribaculaceae* genomes. MAGs were constructed for 9 populations classified as members of the family *Muribaculaceae*, including for two species, OTU-1 and OTU-4, previously shown to respond positively to ACA. All 9 novel MAGs are estimated to be more than 84% complete, and all had less than 2% estimated contamination based on the recovery of ubiquitous, single-copy genes (Table 1). The median N_{50} statistic was approximately 61 kbp, suggesting that assembly was suitable for inferring the genomic context of functional genes. For OTU-1, two closely related genomic variants were recovered, here designated B1A and B1B, possessing 0.56 and 0.31 Mbp of unshared sequence, respectively (Table 2). We designate the MAG constructed for OTU-4 as B2. MAGs obtained from nonresponding species are designated B3 through B8. Estimated genome sizes, %GC, and number of predicted genes are all similar to reference genomes from cultivated members of the family *Muribaculaceae*.

To confirm the assertion that each of the reconstructed genomes is representative of a previously described *Muribaculaceae* species identified in these mice (14), the median mapping rate of metagenomic reads to protein-coding features for each MAG was compared to the relative abundance of the associated 16S rRNA gene across matched amplicon libraries. Reassuringly, cosine similarities were above 0.88 for all MAGs, suggesting robust

TABLE 1 Summary of novel MAGs, previously described, high-quality MAGs, and cultivar genomes^a

Genome	CheckM completeness (%)	No. of scaffolds	Length (Mbp)	N ₅₀	GC (%)	OTU assignment ^b
B1A ^c	95	286	3.1	23,337	46.6	OTU-1
B1B ^c	94	320	2.8	19,144	47.0	OTU-1
B2 ^c	96	116	2.7	75,014	50.6	OTU-4
B3	93	62	2.7	80,587	55.8	OTU-8
B4	97	30	2.7	127,141	55.4	OTU-5
B5	89	124	2.3	46,359	54.2	OTU-6
B6	90	87	2.3	61,151	54.2	OTU-39
B7	84	217	2.9	78,457	48.0	OTU-30
B8	87	154	2.2	33,525	53.4	OTU-11
Mi	99	1	3.3	3,307,069	50.1	
Dm	99	80	3.3	93,062	50.8	
Df	99	5	3.6	2,271,962	48.5	
Dd	97	5	3.7	3,598,157	47.9	
Pi	98	378	2.9	13,720	53.1	
Ha	99	60	2.8	80,867	50.3	
Ac	90	157	2.4	41,191	51.9	

^a*Muribaculum intestinale* (Mi), *Duncaniella muris* (Dm), *Duncaniella freteri* (Df), *Duncaniella dubosii* (Dd), *Paramuribaculum intestinale* (Pi), *"Candidatus Homeothermus arabinosylisolvans"* (Ha), and *"Candidatus Amulmruptor caecigallinaris"* (Ac).

^bOTU data are from reference 14.

^cMAGs from responder populations.

concordance in coverage between the shotgun metagenomic and amplicon libraries. Correlated coverage statistics can be found in the supplemental results, which are available at <https://doi.org/10.5281/zenodo.4450697> (see build/otu_correlation_and_aca_response.ipynb.html).

Phylogenetics. To better understand the evolutionary relationships between these organisms, a concatenated gene tree (Fig. 1A and Fig. S1) was constructed for the 9 new MAGs along with publicly available MAGs and isolate genomes (19–22). The tree was rooted by five other *Bacteroidales* species: *Bacteroides ovatus* (ATCC-8483), *Bacteroides thetaiotaomicron* VPI-5482, *Porphyromonas gingivalis* (ATCC-33277), *Barnesiella viscericola* (DSM-18177), and *Barnesiella intestinihominis* (YIT-11860). Most internal nodes were supported with high topological confidence (>95% bootstrap support), and the placement of the MAGs derived by Ormerod and colleagues (19) was highly consistent with their published tree. To further check the robustness of our phylogeny, a second, approximate maximum likelihood tree was constructed based on the *rpoB* gene, which is generally not thought to be transmitted horizontally (with exceptions [35]). While *rpoB* was not annotated in a number of the genomes, and some nodes were unable to be resolved, this approach largely confirmed the topology of the concatenated gene tree (see Fig. S2). The estimated phylogeny shows that the newly generated MAGs encompass most of the documented diversity of *Muribaculaceae* (Fig. 1A). While many of the novel MAGs are phylogenetically similar to previously described genomes, two of the MAGs, B3 and B4, are notably diverged from the most closely related taxa. This demonstrates that despite a growing number of *Muribaculaceae* genomes deposited in public repositories, novel taxa remain to be described.

TABLE 2 Summary of variant specific features in two highly similar MAGs

Parameter	B1A		B1B	
	Total	Specific	Total	Specific
Nucleotide length (Mbp)	3.07	0.56	2.82	0.31
Genes (no.)	2,612	307	2,419	291
OPFs (no. unique)	2,221	242	2,096	234
KOs (no. unique)	1,026	64	1,000	44
COGs (no. unique)	539	15	528	8
Pfam domains (no. unique)	3,657	635	3,490	488
CAZy domains (no. unique)	252	35	240	18

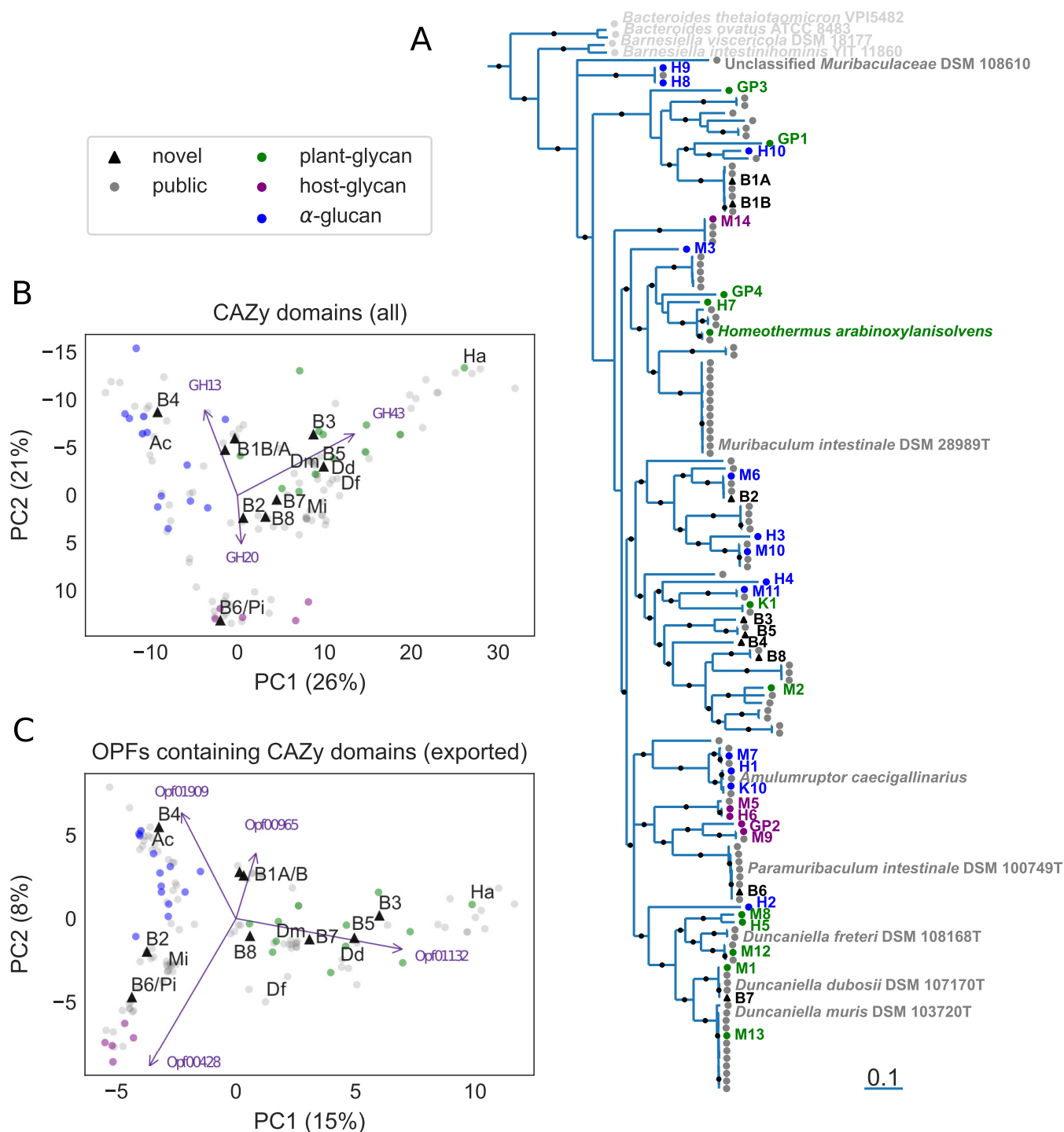


FIG 1 Comparison of novel and previously described *Muribaculaceae* genomes. Novel MAGs (labeled B1A through B8) are combined with publicly available genomes and MAGs as well as 30 MAGs constructed in reference 19 that are hypothesized to reflect three polysaccharide utilization guilds (marker colors). (A) MAGs reconstructed in this study (highlighted in black) are placed in a phylogenetic context using an approximate maximum-likelihood concatenated gene tree based on an amino acid alignment of 36 shared, single-copy genes. The tree is rooted by *Porphyromonas gingivalis* ATCC 33277 (not shown), and four additional *Bacteroidales* genomes are included as an outgroup. Nodes with less than 70% confidence are collapsed into polytomies, and topological support greater than 95% is indicated (black dots on internal branches). Branch lengths indicate an estimate of expected substitutions per site. A version of this panel with GenBank accessions for all publicly available genomes is available as Fig. S1 at <https://doi.org/10.5281/zenodo.4450697>. (B and C) Functional comparisons are visualized by plotting the first two principal components (PCs) of an ordination based on counts of predicted proteins annotated with GH and CBM domains either aggregated by CAZy family (B) or possessing a signal peptide (C) and aggregated by OPF. Purple arrows indicate the contributions of the labeled features, and axes are labeled with the fraction of all variation accounted for by that PC. Novel MAGs (black triangles) are labeled, as are 7 previously described cultivar genomes and high-quality MAGs: “*Candidatus Homeothermus arabinoxylanisolvens*” (Ha), *Muribaculum intestinale* (Mi), *Duncaniella muris* (Dm), *Duncaniella freteri* (Df), *Duncaniella dubosii* (Dd), *Paramuribaculum intestinale* (Pi), “*Candidatus Amuluruptor caecigallinaris*” (Ac).

Novel protein families. Annotations based on alignment to a database of previously characterized sequences may provide only limited insight into the function of gene products, particularly for genes from largely unstudied families of bacteria. In order to identify previously uncharacterized orthologous groups of genes, *de novo* clustering (36) was carried out based on amino acid similarity of all 348,908 putative protein coding genes found in the 9 novel MAGs, 30 MAGs previously generated by Ormerod and colleagues (19), all 98 *Muribaculaceae* genome assemblies available from GenBank, as of this work, and 5 reference genomes from other members of the order *Bacteroidales*. The resulting 16,859 clusters with more than one member contain 315,581 sequences (90%) and are referred to here as operational protein families (OPFs). While a fraction of these clusters may be due to spurious sequence similarity and are without biological relevance, 12,876 have representatives in at least three genomes, increasing the likelihood that these reflect evolutionarily conserved protein sequences. Of this higher confidence set, only 4,448 have members annotated with any COG, KO, or putative function. The remaining 8,528 unannotated OPFs encompass 111,177 predicted protein sequences across the 142 genomes.

To better understand the relationship between our *de novo* clusters and previously described annotations, we inspected the concordance between known GH13 subfamilies, which possess α -amylase activities on different starches (37), and the 20 OPFs with these domains (see build/gh13_families_to_opfs_mapping.ipynb.html at <https://doi.org/10.5281/zenodo.4450697>). We find that most OPFs are composed of members with just one subfamily, suggesting that our clusters are consistent with these known groupings. Several OPFs are dominated by members without a more specific subfamily, and while outside the scope of this study, these may offer hypotheses for additional divisions within GH13. We provide an estimated phylogenetic tree of all GH13 domains, their OPF assignments, and their predicted localization in supplemental material (build/gh13_tree.nwk). Detailed annotations of predicted genes in MAG and reference genomes with OPFs, COGs, KOs, and Pfam and CAZy domains are available as Table S1.

Ordination of gene content. To better understand the polysaccharide utilization potential encoded by the novel MAGs and other available *Muribaculaceae* genomes, we performed an ordination of the counts of genes with predicted homology to GH or CBM domains (Fig. 1B). This holistic analysis of genes that could plausibly be involved in polysaccharide degradation recapitulates the three clusters previously described by Ormerod and colleagues for the first 30 MAGs from the family (19), including associations of the hypothesized α -glucan, plant glycan, and host glycan guilds with GH13, GH43, and GH20 domains, respectively. However, given the ambiguous, intermediate placement of many newer genomes, it also suggests exceptions to the three-guild model. Notably, none of B1A, B1B, or B2 MAGs representative of responding species were placed cleanly within the α -glucan guild, as we had hypothesized.

It is likely that a more nuanced approach to comparing *Muribaculaceae* genomes will yield enhanced predictions of functional capacities. To better reflect *a priori* assumptions about the cellular mechanisms of polysaccharide utilization in the family, the same analysis was repeated, but with two modifications. First, since proteins must be localized to the cell envelope to participate in the breakdown of large, extracellular polysaccharides, genes were only counted if they include an N-terminal signal peptide sequence enabling export from the cytoplasm. Second, given the potential for OPFs to reflect orthologous functions better than domain-level homology, the OPF designations were tallied for predicted proteins with homology to GHs and CBMs rather than the domains themselves. While this more targeted analysis illustrates trends similar to those described before, the distinction between the three clusters appears visually more defined (Fig. 1C). Interestingly, in this ordination B1A, B1B, and several closely related genomes now occupy a space proximate to but distinct from the previously hypothesized α -glucan guild.

Reassuringly, OPFs driving the separation between clusters possess domains matching the original guild descriptions (19). For instance, Opf00428 is associated with MAGs

previously assigned to the host glycan guild, and all but one member of this protein family include regions homologous to GH20. Likewise, Opf01132 parallels the plant glycan guild, and all predicted proteins in this family include a GH43 domain. Surprisingly, not all protein families sharing the same domains are equivalent. Although Opf01909 and Opf00965 both possess a predicted GH13 domain and are the first and fourth most positively weighted features in PC2, indicating an association with the α -glucan guild, the latter is also enriched in the plant glycan guild while the former is not; of the 12 MAGs originally classified to the plant glycan guild, 11 have at least one copy of Opf00965 with a signal peptide versus just one MAG with Opf01909. Based on annotation by the dbCAN2 meta server (31), the GH13 domain in Opf01909 members is classified to the recently defined subfamily 42 (38); however, Opf00965 is not assigned to a more precise subfamily (see [build/gh13_families_to_opfs_mapping.ipynb.html](https://doi.org/10.5281/zenodo.4450697) at <https://doi.org/10.5281/zenodo.4450697>). While a detailed enrichment analysis is outside the scope of this work, this suggests that OPFs indeed provide more functional resolution than annotations based on existing reference databases.

Analysis of MAGs from species responsive to ACA treatment suggests genes involved in starch utilization. Based on the characterization of genes and genomic regions with a role in starch utilization in the closely related genus *Bacteroides*, it is plausible that an α -amylase localized to the outer membrane (OM) is common to starch-utilizing bacteria in the order *Bacteroidales* (39). Indeed, B1A and B1B both have three genes predicted to code for GH13-containing, OM-localized lipoproteins (B1A_01660, B1A_01692, and B1A_02267 in B1A and B1B_01504, B1B_01538, and B1B_02118 in B1B), each in a separate PUL (Fig. 2). While it also includes members without this activity, GH13 is the most common family of α -amylases (40). These genomic regions also possess additional genes with carbohydrate-active domains that are expected to interact with α -glucans.

Besides B1A and B1B, B3 is the only other MAG to possess a putative PUL coding for a full complement of predicted starch-active proteins. This PUL has a large degree of synteny with one of the three matching PULs found in both B1A and B1B and also includes two OPFs with members also found in the *Sus* operon of *B. thetaiotaomicron*, suggesting shared function. However, while B3 also has a GH13-containing lipoprotein (B3_00513), its predicted localization is to the inner membrane, not the OM. It is unclear whether this explains B3's nonresponse in ACA-treated mice. Only two other IM-localized GH13-containing proteins are found across all of the genomes analyzed here (see [build/gh13_tree.nwk](#) in the supplemental materials). Plausible OM-localized, GH13-containing proteins are not found in any nonresponders. While this characteristic does not seem to perfectly discriminate responders from nonresponders (B2 also lacks such a gene), it nonetheless demonstrates concordance between inferred genomic features and observed population dynamics of the corresponding species.

Despite the absence of a GH13 domain on the OM, it is plausible that B2 is capable of degrading starch using other enzymatic mechanisms. We speculate about one putative locus (Fig. 2), which is highly syntenic with characterized dextran PULs in *B. thetaiotaomicron* and *B. ovatus* (41–43). While the only predicted, OM-localized GH contains a GH66 domain (B2_02191), other proximal genes have domains with known affinity for starch, including GH97, CBM20, and CBM69.

To expand the search for relevant genetic features, we identified the subset of *de novo* protein clusters that had members in B1A and B2, and in at most one of the nonresponders, B3 to B8. Of these 52 OPFs, two stood out as particularly relevant. Opf01405 includes *SusR*, the regulator of transcription of the starch utilization system in *B. thetaiotaomicron*, as well as its homolog in *B. ovatus*. It is an apparent subcluster of the larger family defined by K21557 and, in many cases, is located directly upstream of *susC* in putative PULs that are considered likely to have affinity for α -glucans. In each of B1A and B1B, one of the three putative starch PULs encodes a member of Opf01405, and it is similarly located in PULs with starch-active CBM and GH domains in B2 and B3. Interestingly, this OPF is not found in any of the other novel MAGs. In addition, of the seven MAGs constructed by

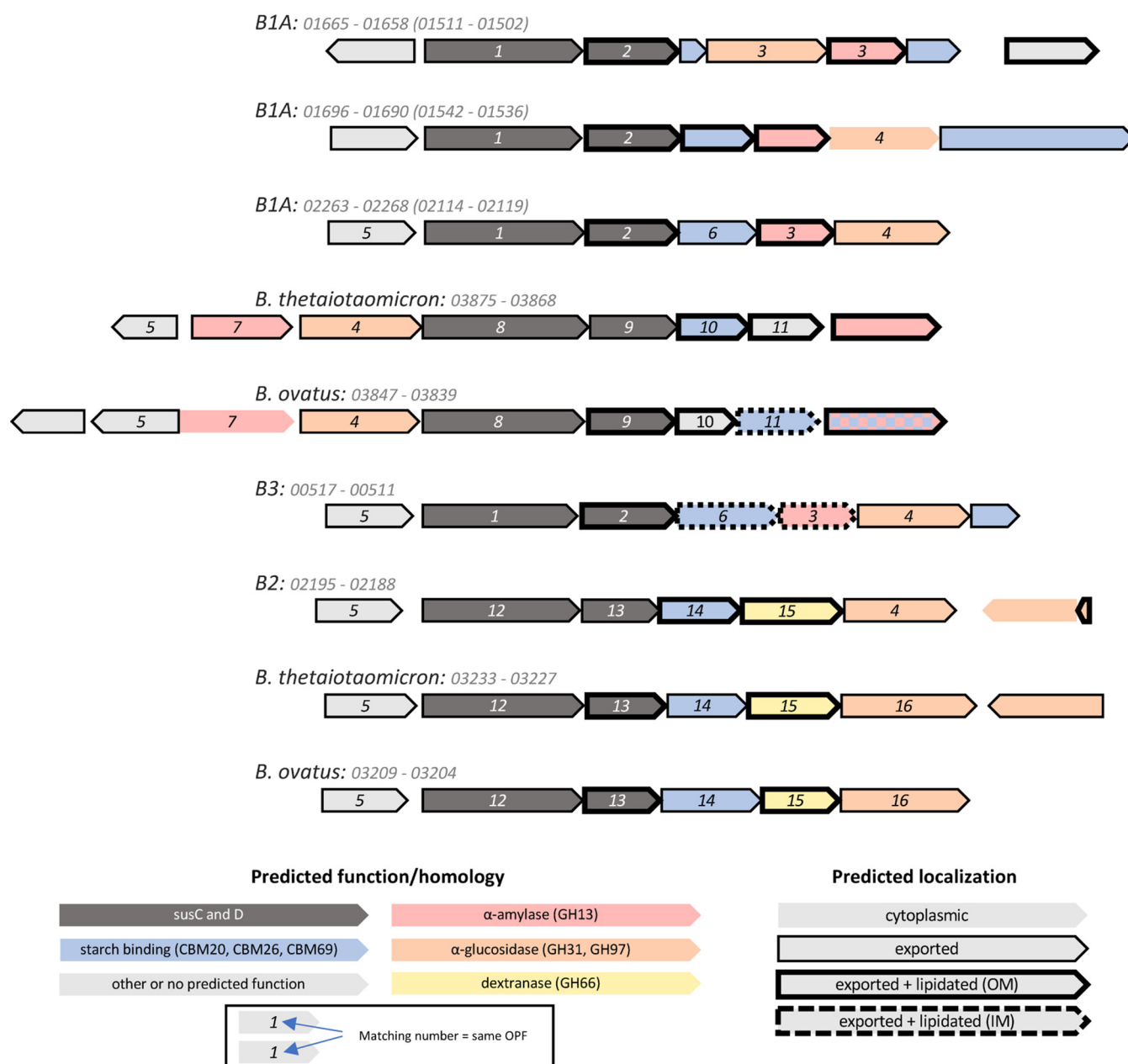


FIG 2 Diagrams of PULs plausibly reflecting activity on starch and/or dextran in B1A, B2, B3, *B. thetaiotaomicron*, and *B. ovatus*. Regions are labeled with the genome name and the interval of gene numbers. For B1A PULs, the matching gene numbers in B1B are noted in parentheses. ORFs are depicted as arrows pointed 5' to 3' along the coding sequence, and colors indicate homology to genes and domains known to participate in either starch or dextran utilization. ORF outlines indicate predicted localization based on the presence of an N-terminal signal peptide and nearby residues. Matching numbers indicate homology based on OPF clustering and are arbitrarily assigned (1, Opf01209; 2, Opf02007; 3, Opf02000; 4, Opf00042; 5, Opf01405; 6, Opf02584; 7, Opf01765; 8, Opf00431; 9, Opf09589; 10, Opf15294; 11, Opf14773; 12, Opf01209; 13, Opf03138; 14, Opf04347; 15, Opf04327; 16, Opf16791).

Ormerod et al. that encode a member of this cluster, five of them are classified to the α -glucan guild. It is plausible that members of Opf01405 share a functional role regulating transcriptional responses to α -glucans.

Finally, we find one example of a GH13-containing OPF that is found in MAGs representing the responder, B1A, B1B, and B2 but none of the other MAGs generated in this study. Opf01765, which recapitulates K21575, includes SusA, the periplasmic neopullulanase of *B. thetaiotaomicron*, and an important component of starch utilization in that organism (44). What's more, the OPF is found in 12 of the 13 α -glucan and a minority of the plant glycan guild members. Interestingly, although it is encoded by the Sus

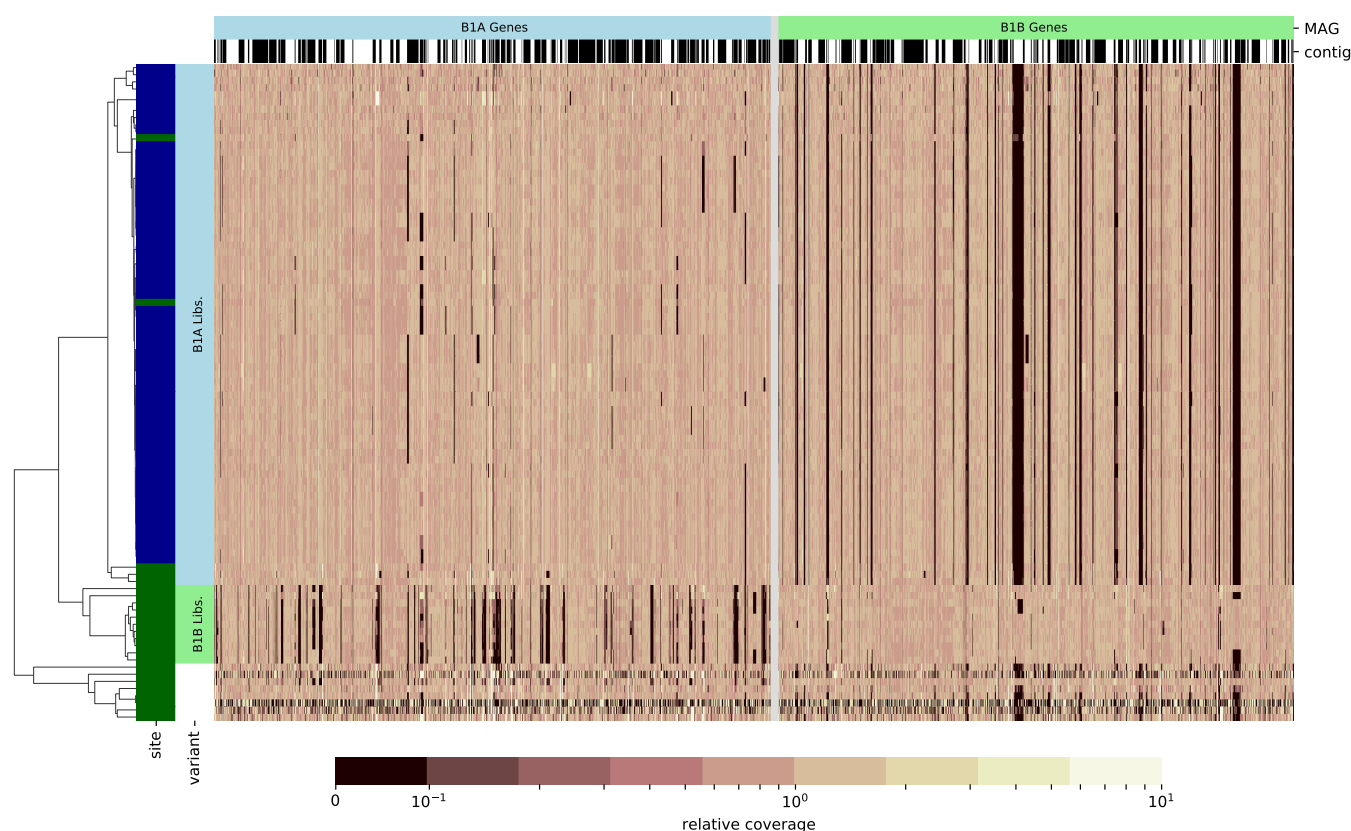


FIG 3 Visualization of differential gene content in two OTU-1 populations. Heatmaps depict relative mapping coverage of metagenomes against putative protein-coding genes in MAGs B1A (left of gray line) or B1B (right). Rows represent one or more pooled libraries for each mouse included in the study, and columns represent individual genes. Alternating black and white spans over heatmap columns indicate contig boundaries in each MAG; the orientation and order of contigs is arbitrary. All coverage values are normalized to the median coverage of that genome's features within each mouse. The site at which each mouse was housed is indicated by colored spans on the left (UT, dark green; UM, dark blue), and mice identified as unambiguous representations of each population are indicated (B1A, light blue; B1B, light green; uncertain, white). Rows are ordered based on a hierarchical clustering by cosine distance, depicted in the tree on the left.

operon in *B. thetaiotaomicron* and its homologous locus in *B. ovatus*, in the *Muribaculaceae* members of Opf01765 are not encoded by PULs, with only 7 of 44 members within 25 kbp of the closest PUL *susC* homolog, and all of these are either upstream of *susC* or on the opposite strand.

Unshared gene content in B1A and B1B. Two distinct genomic variants were associated with OTU-1, with one found in a majority of the UT mouse metagenomes and the other ubiquitous at UM. Using the QAST package (45), 18.1% of the B1A MAG sequence and 11.0% of B1B were found to not align to the other (Table 2). While these hundreds of kilobase pairs may in part reflect errors in genome recovery, much of the unaligned length suggests differences in gene content between these populations of OTU-1. This observation was confirmed by assessing the mapping of metagenomic reads against predicted protein-coding genes in each variant (Fig. 3). For each pairing of metagenomic read library to genomic variant, gene coverage was normalized by the median gene coverage to identify genes with conspicuously fewer reads in particular subsets of the mice. Metagenomic libraries were manually chosen as unambiguous representatives of either B1A or B1B based on concordance in their coverage profiles (Fig. 3), and these were used to systematically identify genes differentiating the two populations.

The median normalized mapping depths in each set of libraries against predicted genes in each MAG were compared, providing a measure of the relative enrichment or depletion of genomic sequences between the two populations of OTU-1. Libraries specific to each variant have low coverage over large portions of either the B1A or B1B MAG, suggesting that mice are primarily colonized by one of the two variants and that a portion of genes are variant specific. At a 5-fold depletion cutoff, selected as a compromise between

sensitivity and specificity in identifying genomic regions that differentiate the two, we found 11.7% of predicted genes in B1A were depleted in B1B populations, and 12.0% of B1B genes were depleted in B1A populations (Table 2). Coverage ratios over all genes in B1A and B1B can be found in Table S2 at <https://doi.org/10.5281/zenodo.4450697>. While the observed coverage ratios could indicate variation in copy number, differential gene content between variants is a more parsimonious explanation for most loci. These predicted genes reflect 6.2% of unique KOs in B1A and 4.4% in B1B. Interestingly, the fraction of variant-specific OPFs is greater, 10.8% and 11.1%, respectively, suggesting that *de novo* clustering is more sensitive to potential differences in physiology.

Given the observation that the relative abundance of OTU-1 was dramatically increased with ACA treatment at UM while not being significantly affected at UT and that B1B was not found in metagenomes at UM, we hypothesized that differences in the genomic potential of B1A and B1B explain the different responses to ACA at the two sites.

Genomic regions apparently specific to B1A, defined as an at least 5-fold enrichment in B1A-specific libraries relative to B1B-specific libraries, include just one PUL (SusC homolog encoded by B1A_02041). This locus includes a predicted OM-localized, GH30-containing protein. Proteins that contain a GH30 domain have β -glucosylceramidase, β -1,6-glucanase, or β -xylosidase activity (46). Given that this PUL also encodes a periplasmic, GH3-containing protein, it appears unlikely that it has specificity for starch. Both variants also possess numerous phage insertions not seen in the other. Likewise, a CRISPR operon (Cas9 encoded by B1B_00401) appears to be specific to B1B.

Most strikingly, a 15-kbp region specific to B1A (from B1A_01550 to B1A_01566) was found to contain numerous genes with homology to cell capsule and exopolysaccharide synthesizing enzymes. Based on annotations with KEGG orthologous groups, these include homologs of *tuaG* (K16698), *tagE* (K00712), *gmhB* (K03273), *gmhA* (K03271), *hddA* (K07031), *exoO* (K16555), *waaH* (K19354), and *tagF* (K09809). Interestingly, the B1B MAG contains a different region of about 6.2 kbp (B1B_00746 to B1B_00751) with *wfeD* (K21364), *pglI* (K17248), and *epsH* (K19425). For each, several of the OPFs in the respective regions were not found anywhere in the opposing genome, suggesting that the makeup of each variant's exterior surface is distinctly different.

DISCUSSION

Mice are a key model system for the study of the mammalian gut microbiome, with an outsized importance in testing mechanistic hypotheses about the roles of this community in host health (47). The generalizability of observations made in mice is a constant concern (47), in part due to extensive difference in taxonomic composition compared to humans (20). Bacteria classified in the family *Muribaculaceae* are abundant in the murine gut microbiome (19). While these bacteria are also found in humans (albeit at lower abundance), only a few members of this clade have been cultivated and described (20–22). As a result, the ecological roles of these bacteria have not yet been characterized, limiting the value of the mouse as a model system. Better understanding the ecology of *Muribaculaceae* in the murine gut will increase the transferability of microbiome findings from mice to humans. Attempts to study these organisms make use of genomes reconstructed from metagenomic reads and have suggested, in the absence of experimental data, that members of the family consume a diversity of polysaccharides in the lower gut.

Here, we have extended that approach to nine new genomes and associated those with species for which changes in relative abundance in response to ACA treatment have been experimentally assessed. This enabled us to explore why two responding species, represented by MAGs B1A, B1B, and B2, increase with ACA treatment while the other species of *Muribaculaceae* do not. Annotations of reconstructed genomes suggest that the responders possess starch degradation capabilities absent from the nonresponders.

By including otherwise unannotated genes, genomic comparisons based on OPFs may reflect shared functional potential better than applying previously defined orthologies. Besides the identification of novel gene families, *de novo* homology clustering

(36) also enables differentiation of subgroups not captured by standard annotations. For instance, hypothetical genes annotated as homologs of SusC, SusD, and SusEF were clustered into 98, 187, and 191 different OPFs, respectively. It is plausible that this subclustering captures differences in protein structure with importance in oligo- and polysaccharide recognition, import, and binding. Combined with annotation of characterized functional domains, we hypothesize that these clusters more narrowly predict the polysaccharide utilization ranges of uncultured organisms. Testing these predictions will require characterization of the metabolic potential of these genes after obtaining cultivars or through heterologous expression in appropriate hosts.

We examine the three-guild model proposed by Ormerod and colleagues (19) by extending their dimensional reduction approach to the much larger set of *Muribaculaceae* genomes now available. In this analysis, annotations of B1A, B1B, and B2 are not clearly colocated with members of the α -glucan guild, complicating this simple interpretation. Interestingly, a more nuanced analysis involving OPFs and predicted export indicates that B1A and B1B have polysaccharide utilization potentials distinct from the α -glucan guild previously described. The improved resolution presented by OPF clusters suggests that this more detailed examination can identify specific functions that discriminate responders from nonresponders. The approach is bolstered by the phylogenetic and genomic distinction between B2 and both B1A and B1B, reducing the confounding effects of shared evolutionary history.

A detailed analysis of PULs identified multiple loci shared in both B1A and B1B that appear to be adapted to the degradation of starch or related carbohydrates due to the presence of an OM-localized GH13-containing protein (48). Counterintuitively, B2 had no such PUL, suggesting that its response to ACA results from other enzymatic capabilities. Of particular interest is a PUL encoding proteins with GH97, CBM20, and CBM69 domains, all of which have documented activity on starch (49, 50). While the only OM-localized hydrolase in this PUL is a GH66, and members of this family have characterized activity on the α -1,6 linkages between glucose monomers in dextran (51), it is plausible that this PUL can be repurposed and confers some ability to grow on starch.

Most compellingly, a gene family encoding a SusA homolog was identified in B1A, B1B, and B2 but in none of the nonresponders, presenting the best case for a single enzyme that can confer a growth advantage in ACA-treated mice. While it is unclear how expression of this important component of starch utilization might be regulated, given that it is not located in a PUL in any of the responding populations, SusA is important for growth on amylopectin in *B. thetaiotaomicron* (44). Since inhibition by ACA is variable across enzymes (52), it is possible that ACA treatment results in elevated production of dextrin and maltooligosaccharides in the lower guts of mice due to residual α -amylase activity, even at levels sufficient to prohibit host digestion. Periplasmic hydrolysis of these starch breakdown products may be sufficient for increased abundance of these species in ACA-treated mice.

It is notable that of the closely related variants associated with OTU-1, B1B is found at UT and not UM. We previously observed site specificity of the ACA response of this species in which OTU-1 did not have a significantly increased abundance in treated mice at UT, while it was the most dramatic change at UM (14). Differences in the functional potential due to differences in gene content of populations found at each of the sites is one possible explanation for this pattern. Alternatively, differences in the occurrence or gene content of other microbial community members could lead to the differential response of OTU-1 across these sites, for instance, by contributing to the partial breakdown of larger starch molecules or through resource competition. Notably, ACA appears to be less inhibitory to members of the *Firmicutes* than the *Bacteroidetes* (53). Intriguingly, while we do not conjecture a mechanistic link, an ACA-by-site interaction effect on longevity has been previously observed in the mouse colonies sampled here, with male mice at UT showing a larger increase in median longevity with ACA treatment than those at UM (15, 17).

Despite evidence that large differences in gene content can be found between even closely related populations (54, 55), studies reconstructing genomes from

metagenomes have just started to consider these pangenome dynamics (56–60). Improvements to software (e.g., binning) and sequencing technologies (e.g., long reads) will increase the accuracy of physiological inferences based on homology and synteny. The discovery of two populations of OTU-1 therefore demonstrates the value of considering pangenome dynamics and presents a potential explanation for the observed site-specific response of that taxon. The finding that both variants have the same complement of three PULs apparently specializing in starch utilization and the same *SusA* homolog does not support the hypothesis that differences in starch utilization potential account for these abundance patterns. We did, however, identify numerous differences in the gene content of B1A and B1B, including variant-specific loci that may influence the structure and function of the outer surface of the cell. Given the size of these regions, their parallel physiological roles, and their plausible effect on interactions with the host or other microbes (61), we speculate that these variable loci could result in relevant, functional differences between the two populations.

While these results do not establish a mechanistic explanation for differences in the response of B1A and B1B at UM and UT or conclusively identify the pathways that enable starch utilization in B2, they do suggest a number of genomic features that likely contribute to previously observed patterns in taxon abundance. Future studies utilizing metatranscriptomic analysis might demonstrate active expression of these genes or differential expression in mice treated with ACA compared to controls. Likewise, even in the absence of a B2 cultivar, the potential role of its dextran PUL in enrichment under ACA treatment could be tested using available cultivars, like *B. thetaiotaomicron*, that possess a homologous gene cluster.

Conclusions. In this study, we have reconstructed and described genomes representing 8 species in the family *Muribaculaceae* from the mouse fecal microbiome and have found features that differentiate those bacterial species that respond positively to ACA treatment from those that do not. This analysis suggests that utilization of starch and related polysaccharides enables increased population size in mice treated with ACA, an α -amylase inhibitor. In addition, two distinct genomic variants of one species were identified that differ in functional gene content, potentially explaining site-specific differences in response. By combining observed changes in relative abundance during experimental manipulation with inferred functional gene content, we are able to study mammalian symbionts in the absence of cultured representatives. This sequence-based approach is broadly applicable in microbial ecology and enables improved understanding of *in situ* dynamics within complex microbial communities.

MATERIALS AND METHODS

Mouse treatment, sample collection, extraction, and sequencing. Mice were bred, housed, and treated as described previously (15). Briefly, genetically heterogeneous UM-HET3 mice at each study site were produced by a four-way cross between (BALB/cByJ \times C57BL/6J) F1 mothers and (C3H/HeJ \times DBA.2J) F1 fathers, as detailed previously (62). Mice were fed LabDiet 5LG6 (TestDiet Inc.) from weaning onwards. Starting at 8 months of age, mice randomly assigned to treatment were fed chow with 1,000 ppm ACA (Spectrum Chemical Manufacturing Corporation). Mice were housed 3 males or 4 females to a cage. Colonies were assessed for infectious agents every 3 months, and all tests were negative.

Individual fecal pellets were collected from a single mouse per cage. 16S rRNA gene libraries and metabolite analyses of these samples were as described previously (14). From this collection, a subset of samples was nonrandomly selected for metagenomic sequencing to test several unrelated hypotheses about SCFA production. Samples were from 54 mice, with at least six treated and control representatives of both males and females at each site.

Fecal samples were slurried with nuclease-free water at a 1:10 (wt/vol) ratio, and most samples were spiked with *Sphingopyxis alaskensis* RB2256 prepared as described previously (14) before DNA extraction and sequencing. Based on alignment to the reference genome, sequenced reads from *S. alaskensis* can be distinguished from all endogenous bacteria in mouse feces. This spike was added as an internal standard to quantify total 16S rRNA gene abundance in a separate study (14). A small number of these were split for both spiked and unspiked samples, which we used to validate this procedure. For each, 150 μ l of this sample was transferred for extraction using the MoBio PowerMag microbiome kit. Metagenomic libraries were prepared using standard procedures and sequenced on the Illumina HiSeq 400 platform using v4 paired-end 2 \times 150 bp.

Assembly, binning, and MAG refinement. Bioinformatic processing of metagenomes was implemented as a Snakemake pipeline (63) run with version 5.18.1. Code and metadata to run this pipeline can be obtained at <https://doi.org/10.5281/zenodo.4450697>.

Raw metagenomic reads were deduplicated using FastUniq (64) version 1.1, adapter trimmed using Scythe (65) version 0.991, and quality trimmed using Sickle (66) version 1.33 to produce processed reads for all downstream analyses. The resulting paired-end reads from all samples were coassembled into primary contigs using MEGAHIT (67) version 1.2.9. Reads were then mapped back to these contigs with Bowtie2 (68) version 2.4.1, and per-library coverage was estimated for each contig.

For all contigs >1,000 bp in length, dimensional reductions built into CONCOCT (69) version 1.1.0 were applied to produce input data for a Gaussian mixture model (GMM) similar to the procedure used by that program for binning. However, unlike CONCOCT, due to computational limitations, the model was trained on just 10% of the input data, sampled randomly, before assigning bins to all contigs. While this may have reduced the accuracy of the binning procedure, we believe that subsequent refinement steps mitigated the impact of this decision.

Using 16S rRNA gene libraries described in reference 14 and processed as described in that reference, operational taxonomic units (OTUs) were classified taxonomically and relative abundance was calculated. Bins were then recruited to one or more OTUs by calculating a canonical partial least square between OTU abundance and bin coverage as implemented in the scikit-learn machine learning library (70) version 0.23.2. For bins recruited to OTUs classified as *Muribaculaceae*, contigs were reclustered based on coverage across samples. First, trusted contigs that correlated closely with OTU abundance were manually selected. The mean coverage of these was used to normalize the per-library coverage of all other contigs. Next, using a GMM, groups of contigs were clustered such that the normalized coverage across samples was consistent. These groups were used to inform the manual assignment of contigs to MAGs. Libraries in which MAGs had nonnegligible coverage were identified manually and used in subsequent refinements. While clustering contigs were associated with OTU-1, a number of groups containing on the order of 10^5 bp were found with a bimodal coverage distribution, low normalized coverage in a subset of libraries, and normal coverage in others. By this criterion, contigs in these variable groups were partitioned into two MAG variants, A and B, with the nonvariable contig groups shared by both. To avoid challenges associated with admixture, only libraries that appeared on further inspection to have coverage over just one of the two variants were considered in downstream refinement steps. The mice matching these libraries are highlighted in Fig. 3. None of the other *Muribaculaceae* OTUs described in this study were found during this manual curation process to have genomic variants comparable to B1A and B1B.

For each MAG, the list of contigs and relevant libraries identified in the previous step, as well as the per-library coverage of trusted contigs, were used in a final refinement procedure. All read pairs from the relevant libraries with at least one read mapping to the contigs were extracted and used in a single pass of the Pilon assembly refinement tool (71) version 1.23. Regions of these improved contigs were then excised where they had low cosine similarity to the trusted contig coverage, using a 0, 0.5, 0.6, 0.7, 0.8, and 0.9 cosine similarity threshold. Genome completeness and contamination estimates were calculated based on ubiquitous single-copy genes using the program CheckM (72) version 1.1.2. Based on these results, the final refined assembly with the highest completeness and with contamination of <2% was selected; ties were broken using the N_{50} statistic.

Reference genomes. To compare our 9 novel MAGs to previously reconstructed genomes, we downloaded a total of 98 draft genomes from GenBank, both isolates and MAGs, representing all genomes taxonomically assigned to the *Muribaculaceae* as of September 2020, along with the 30 MAGs described in reference 19 (BioProject no. [PRJNA313232](https://www.ncbi.nlm.nih.gov/bioproject/PRJNA313232)). For comparison, nucleotide sequences for *Bacteroides thetaiotaomicron* VPI-5482 ([GCA_900624795.1](https://www.ncbi.nlm.nih.gov/nuccore/GCA_900624795.1)), *Bacteroides ovatus* ATCC-8483 ([GCA_000154125.1](https://www.ncbi.nlm.nih.gov/nuccore/GCA_000154125.1)), *Barnesiella viscericola* DSM-18177 ([GCA_000512915.1](https://www.ncbi.nlm.nih.gov/nuccore/GCA_000512915.1)), *Barnesiella intestinhominis* YIT-11860 ([GCA_000296465.1](https://www.ncbi.nlm.nih.gov/nuccore/GCA_000296465.1)), and *Porphyromonas gingivalis* ATCC-33277 ([GCA_000010505.1](https://www.ncbi.nlm.nih.gov/nuccore/GCA_000010505.1)) were also downloaded.

Genome annotation. All genomes were initially annotated with Prokka (73) version 1.14.6, which uses Prodigal (74) version 2.6.3 for gene finding. Putative protein sequences were additionally annotated with domains from both the dbCAN database (29) release 6 of carbohydrate-active domains and Pfam (75) release 31.0, using HMMER3 (76, 77) version 3.3. To maximize the sensitivity of this analysis, these annotations used a bitscore cutoff of 5.0 and no cutoff (any hit). Protein sequences were also annotated with KO numbers by BLAST using the KEGG database as of March 2018 as the reference and taking the best hit with a maximum E value of $1e-10$.

Lipoproteins were predicted using LipoP (78) version 1.0a, a score cutoff of 5, and a margin cutoff of 2. Lipoproteins with an aspartate at position +2 relative to the cleavage site were labeled as localized to the inner membrane (79, 80). Periplasmic proteins were identified with SignalP (81) version 4.1. Predicted protein sequences from all annotated genomes were first dereplicated using CD-HIT (82, 83) version 4.8.1 at a similarity threshold of 0.99 and then locally all-by-all aligned using the DIAMOND (84) version 0.9.31 implementation of the BLASTp algorithm. Each pair was then assigned a similarity value as the bitscore of their best local alignment normalized by the greater of the two self-alignments. This results in a matrix of pairwise scores reflecting the proximity to perfect homology. Scores of less than 0.2 were replaced with 0. Clusters were formed using the MCL algorithm (85) version 14-137 with an inflation parameter of 5.

SusCDEF homologs were identified based on relatively relaxed criteria, harnessing OPF assignments, domain predictions, and Prokka annotations to avoid false negatives while maintaining specificity. For each OPF, all KOs assigned to members were collected as plausible KOs for the cluster. Protein sequences in OPF clusters that included K21572 were flagged as putative SusC homologs, as were sequences directly annotated as such by Prokka. Using a similar approach, proteins in clusters tagged with K21571

or with any of domains PF12771, PF14322, PF12741, and PF07980 were identified as putative SusD. Proteins in clusters tagged with K21571, or with either PF14292 or PF16411, were considered SusEF homologs. Putative PULs were identified by tandem *susCD*-like pairs: a SusC homolog within 5 kbp of a SusD homolog on the same strand, similar to that described in reference 86. Of the 121 putative PULs identified across the nine novel MAGs in this study, 103 of them have at least one gene within 10 kbp with homology to a CAZy domain family, suggesting that the majority of our identified PULs indeed have activity on polysaccharides.

Annotations were compared across all genomes using dimensional reduction (Fig. 1B and C). To reduce the effects of redundancy on ordination, highly similar genomes were first clustered by complete linkage at a 0.8 cosine similarity threshold. Clusters were then replaced by their mean counts before principal component analysis. Finally, the original genomes were projected onto these principal components and visualized.

Phylogenetics. Predicted amino acid sequences for open reading frames from all MAGs and all reference genomes were searched for homology to the TIGRFAM protein clusters (87) release 14.0 using HMMER3. Hits were filtered at the trusted-cutoff score threshold, defined separately for each protein model. Sequences found in no more than one copy and in at least 95% of genomes were used as taxonomic marker genes for phylogenetic analysis. Marker gene sequences were aligned to their respective models using HMMER3, dropping unaligned residues. Aligned markers were then concatenated for each genome. The concatenated alignment was masked using Gblocks (Castresana2000) version 0.91b using a wrapper script provided at https://github.com/mattb112885/clusterDbAnalysis/blob/master/src/Gblocks_wrapper.py, with default parameters, before estimating an approximate maximum likelihood phylogeny using the FastTree software (88) version 2.1.10 with default parameters.

Data availability. MAGs generated for this study were deposited in GenBank under accession numbers JAEKDF000000000 (B1A), JAEKDG000000000 (B1B), JAEKDH000000000 (B2), JAEKDI000000000 (B3), JAEKDJ000000000 (B4), JAEKDK000000000 (B5), JAEKDL000000000 (B6), JAEKDM000000000 (B7), and JAEKDN000000000 (B8). Metagenomic libraries are also available under BioProject no. PRJNA448009. Code and additional data/metadata needed to reproduce this analysis, as well as all supplemental materials, are available at <https://doi.org/10.5281/zenodo.4450697>.

ACKNOWLEDGMENTS

We acknowledge technical support from Randy Strong and others at The University of Texas Health Science Center at San Antonio for initial sample collection. We also thank Nicole Koropatkin for feedback on the manuscript.

Funding for this work was provided in part by a grant from the Glenn Foundation for Medical Research. B.J.S. received support from the University of Michigan ITiMS Training Grant.

REFERENCES

- Turnbaugh PJ, Ley RE, Mahowald MA, Magrini V, Mardis ER, Gordon JI. 2006. An obesity-associated gut microbiome with increased capacity for energy harvest. *Nature* 444:1027–1031. <https://doi.org/10.1038/nature05414>.
- Britton RA, Young VB. 2012. Interaction between the intestinal microbiota and host in *Clostridium difficile* colonization resistance. *Trends Microbiol* 20:313–319. <https://doi.org/10.1016/j.tim.2012.04.001>.
- Syal G, Kashani A, Shih DQ. 2018. Fecal microbiota transplantation in inflammatory bowel disease: a primer for internists. *Am J Med* 131:1017–1024. <https://doi.org/10.1016/j.amjmed.2018.03.010>.
- Hiele M, Ghooys Y, Rutgeerts P, Vantrappen G. 1992. Effects of acarbose on starch hydrolysis. *Dig Dis Sci* 37:1057–1064. <https://doi.org/10.1007/BF01300287>.
- Dehghan-Kooshkghazi M, Mathers JC. 2004. Starch digestion, large-bowel fermentation and intestinal mucosal cell proliferation in rats treated with the α -glucosidase inhibitor acarbose. *Br J Nutr* 91:357–365. <https://doi.org/10.1079/BJN20031063>.
- Zhao L, Zhang F, Ding X, Wu G, Lam YY, Wang X, Fu H, Xue X, Lu C, Ma J, Yu L, Xu C, Ren Z, Xu Y, Xu S, Shen H, Zhu X, Shi Y, Shen Q, Dong W, Liu R, Ling Y, Zeng Y, Wang X, Zhang Q, Wang J, Wang L, Wu Y, Zeng B, Wei H, Zhang M, Peng Y, Zhang C. 2018. Gut bacteria selectively promoted by dietary fibers alleviate type 2 diabetes. *Science* 359:1151–1156. <https://doi.org/10.1126/science.aao5774>.
- Holt PR, Atillasoy E, Lindenbaum J, Ho SB, Lupton JR, McMahon D, Moss SF. 1996. Effects of acarbose on fecal nutrients, colonic pH, and short-chain fatty acids and rectal proliferative indices. *Metabolism* 45:1179–1187. [https://doi.org/10.1016/s0026-0495\(96\)90020-7](https://doi.org/10.1016/s0026-0495(96)90020-7).
- Wolever TMS, Chiasson JL. 2000. Acarbose raises serum butyrate in human subjects with impaired glucose tolerance. *Br J Nutr* 84:57–61. <https://doi.org/10.1017/S0007114500001239>.
- Weaver GA, Tangel CT, Krause JA, Parfitt MM, Jenkins PL, Rader JM, Lewis BA, Miller TL, Wolin MJ. 1997. Acarbose enhances human colonic butyrate production. *J Nutr* 127:717–723. <https://doi.org/10.1093/jn/127.5.717>.
- Weaver GA, Tangel CT, Krause JA, Parfitt MM, Stragand JJ, Jenkins PL, Erb TA, Davidson RH, Alpern HD, Guiney WB, Higgins PJ. 2000. Biomarkers of human colonic cell growth are influenced differently by a history of colonic neoplasia and the consumption of acarbose. *J Nutr* 130:2718–2725. <https://doi.org/10.1093/jn/130.11.2718>.
- Wolin MJ, Miller TL, Yerry S, Bank S, Weaver GA, Zhang Y. 1999. Changes of fermentation pathways of fecal microbial communities associated with a drug treatment that increases dietary starch in the human colon changes of fermentation pathways of fecal microbial communities associated with a drug treatment that increases dietary starch in the human colon. *Appl Environ Microbiol* 65:2807–2812. <https://doi.org/10.1128/AEM.65.7.2807-2812.1999>.
- Zhang X, Fang Z, Zhang C, Xia H, Jie Z, Han X, Chen Y, Ji L. 2017. Effects of acarbose on the gut microbiota of prediabetic patients: a randomized, double-blind, controlled crossover trial. *Diabetes Ther* 8:293–307. <https://doi.org/10.1007/s13300-017-0226-y>.
- Baxter NT, Lesniak NA, Sinani H, Schloss PD, Koropatkin NM. 2019. The glucoamylase inhibitor acarbose has a diet-dependent and reversible effect on the murine gut microbiome. *mSphere* 4:e00528-18. <https://doi.org/10.1128/mSphere.00528-18>.
- Smith BJ, Miller RA, Ericsson AC, Harrison DC, Strong R, Schmidt TM. 2019. Changes in the gut microbiome and fermentation products concurrent with enhanced longevity in acarbose-treated mice. *BMC Microbiol* 19:130. <https://doi.org/10.1186/s12866-019-1494-7>.
- Harrison DE, Strong R, Allison DB, Ames BN, Astle CM, Atamna H, Fernandez E, Flurkey K, Javors MA, Nadon NL, Nelson JF, Pletcher S, Simpkins JW, Smith DL, Wilkinson JE, Miller RA. 2014. Acarbose, 17-

- α -estradiol, and nordihydroguaiaretic acid extend mouse lifespan preferentially in males. *Aging Cell* 13:273–282. <https://doi.org/10.1111/acer.12170>.
16. Strong R, Miller RA, Antebi A, Astle CM, Bogue M, Denzel MS, Fernandez E, Flurkey K, Hamilton KL, Lamming DW, Javors MA, de Magalhães JP, Martinez PA, McCord JM, Miller BF, Müller M, Nelson JF, Ndukum J, Rainger GE, Richardson A, Sabatini DM, Salmon AB, Simpkins JW, Steegenga WT, Nadon NL, Harrison DE. 2016. Longer lifespan in male mice treated with a weakly estrogenic agonist, an antioxidant, an α -glucosidase inhibitor or a Nrf2-inducer. *Aging Cell* 15:872–884. <https://doi.org/10.1111/acer.12496>.
 17. Harrison DE, Strong R, Alavez S, Astle CM, DiGiovanni J, Fernandez E, Flurkey K, Garratt M, Gelfond JAL, Javors MA, Levi M, Lithgow GJ, Macchiarini F, Nelson JF, Sukoff Rizzo SJ, Slaga TJ, Stearns T, Wilkinson JE, Miller RA. 2019. Acarbose improves health and lifespan in aging HET3 mice. *Aging Cell* 18:e12898. <https://doi.org/10.1111/acer.12898>.
 18. Salzmann KL, de Jong H, Paterson Y, Harmsen HJM, Welling GW, Bos NA. 2002. Analysis of 16S libraries of mouse gastrointestinal microflora reveals a large new group of mouse intestinal bacteria. *Microbiology (Reading)* 148:3651–3660. <https://doi.org/10.1099/00221287-148-11-3651>.
 19. Ormerod KL, Wood DLA, Lachner N, Gellatly SL, Daly JN, Parsons JD, Dal'Molin CGO, Palfreyman RW, Nielsen LK, Cooper MA, Morrison M, Hansbro PM, Hugenholtz P. 2016. Genomic characterization of the uncultured Bacteroidales family S24-7 inhabiting the guts of homeothermic animals. *Microbiome* 4:36. <https://doi.org/10.1186/s40168-016-0181-2>.
 20. Lagkouvardos I, Pukall R, Abt B, Foesel BU, Meier-Kolthoff JP, Kumar N, Bresciani A, Martinez I, Just S, Ziegler C, Brugiroux S, Garzetti D, Wenning M, Bui TPN, Wang J, Hugenholtz F, Plugge CM, Peterson DA, Hornef MW, Baines JF, Smidt H, Walter J, Kristiansen K, Nielsen HB, Haller D, Overmann J, Stecher B, Clavel T. 2016. The Mouse Intestinal Bacterial Collection (miBC) provides host-specific insight into cultured diversity and functional potential of the gut microbiota. *Nat Microbiol* 1:16131. <https://doi.org/10.1038/nmicrobiol.2016.131>.
 21. Lagkouvardos I, Lesker TR, Hitch TCA, Gálvez EJC, Smit N, Neuhaus K, Wang J, Baines JF, Abt B, Stecher B, Overmann J, Strowig T, Clavel T. 2019. Sequence and cultivation study of *Muribaculaceae* reveals novel species, host preference, and functional potential of this yet undescribed family. *Microbiome* 7:28. <https://doi.org/10.1186/s40168-019-0637-2>.
 22. Miyake S, Ding Y, Soh M, Low A, Seedorf H. 2020. Cultivation and description of *Duncaniella dubosii* sp. nov., *Duncaniella freteri* sp. nov. and emended description of the species *Duncaniella muris*. *Int J Syst Evol Microbiol* <https://doi.org/10.1099/ijsem.0.004137>.
 23. Parks DH, Rinke C, Chuvochina M, Chaumeil PA, Woodcroft BJ, Evans PN, Hugenholtz P, Tyson GW. 2017. Recovery of nearly 8,000 metagenome-assembled genomes substantially expands the tree of life. *Nat Microbiol* 2:1533–1542. <https://doi.org/10.1038/s41564-017-0012-7>.
 24. Lee STM, Kahn SA, Delmont TO, Shaiber A, Esen Özcan C, Hubert NA, Morrison HG, Antonopoulos DA, Rubin DT, Eren AM. 2017. Tracking microbial colonization in fecal microbiota transplantation experiments via genome-resolved metagenomics. *Microbiome* 5:10. <https://doi.org/10.1186/s40168-017-0270-x>.
 25. Martens EC, Koropatkin NM, Smith TJ, Gordon JL. 2009. Complex glycan catabolism by the human gut microbiota: the Bacteroidetes Sus-like paradigm. *J Biol Chem* 284:24673–24677. <https://doi.org/10.1074/jbc.R109.022848>.
 26. Foley MH, Cockburn DW, Koropatkin NM. 2016. The Sus operon: a model system for starch uptake by the human gut Bacteroidetes. *Cell Mol Life Sci* 73:2603–2617. <https://doi.org/10.1007/s00018-016-2242-x>.
 27. Grondin JM, Tamura K, Déjean G, Abbott DW, Brumer H. 2017. Polysaccharide utilization loci: fueling microbial communities. *J Bacteriol* 199:e00860–16. <https://doi.org/10.1128/JB.00860-16>.
 28. Fernández-Gómez B, Richter M, Schüller M, Pinhassi J, Acinas SG, González JM, Pedrós-Alió C. 2013. Ecology of marine Bacteroidetes: a comparative genomics approach. *ISME J* 7:1026–1037. <https://doi.org/10.1038/ismej.2012.169>.
 29. Yin Y, Mao X, Yang J, Chen X, Mao F, Xu Y. 2012. dbCAN: a web resource for automated carbohydrate-active enzyme annotation. *Nucleic Acids Res* 40:445–451. <https://doi.org/10.1093/nar/gks479>.
 30. Lombard V, Golaconda Ramulu H, Drula E, Coutinho PM, Henrissat B. 2014. The carbohydrate-active enzymes database (CAZy) in 2013. *Nucleic Acids Res* 42:D490–D495. <https://doi.org/10.1093/nar/gkt1178>.
 31. Zhang H, Yohe T, Huang L, Entwistle S, Wu P, Yang Z, Busk PK, Xu Y, Yin Y. 2018. dbCAN2: a meta server for automated carbohydrate-active enzyme annotation. *Nucleic Acids Res* 46:W95–W101. <https://doi.org/10.1093/nar/gky418>.
 32. The CAZypedia Consortium. 2018. Ten years of CAZypedia: a living encyclopedia of carbohydrate-active enzymes. *Glycobiology* 28:3–8. <https://doi.org/10.1093/glycob/cwx089>.
 33. Lagkouvardos I, Overmann J, Clavel T. 2017. Cultured microbes represent a substantial fraction of the human and mouse gut microbiota. *Gut Microbes* 8:493–503. <https://doi.org/10.1080/19490976.2017.1320468>.
 34. Stewart EJ. 2012. Growing unculturable bacteria. *J Bacteriol* 194:4151–4160. <https://doi.org/10.1128/JB.00345-12>.
 35. Kim BJ, Hong SH, Kook YH, Kim BJ. 2013. Molecular evidence of lateral gene transfer in *rpoB* gene of *Mycobacterium yongonense* strains via multilocus sequence analysis. *PLoS One* 8:e51846. <https://doi.org/10.1371/journal.pone.0051846>.
 36. Schloss PD, Handelsman J. 2008. A statistical toolbox for metagenomics: assessing functional diversity in microbial communities. *BMC Bioinformatics* 9:34–45. <https://doi.org/10.1186/1471-2105-9-34>.
 37. Stam MR, Danchin EGJ, Rancurel C, Coutinho PM, Henrissat B. 2006. Dividing the large glycoside hydrolase family 13 into subfamilies: towards improved functional annotations of α -amylase-related proteins. *Protein Eng Des Sel* 19:555–562. <https://doi.org/10.1093/protein/gz1044>.
 38. Valk V, van der Kaaij RM, Dijkhuizen L. 2016. Characterization of the starch-acting MaAmyB enzyme from *Microbacterium aurum* B8.A representing the novel subfamily GH13_42 with an unusual, multi-domain organization. *Sci Rep* 6:36100. <https://doi.org/10.1038/srep36100>.
 39. Shipman JA, Cho KH, Siegel HA, Salyers AA. 1999. Physiological characterization of SusG, an outer membrane protein essential for starch utilization by *Bacteroides thetaiotaomicron*. *J Bacteriol* 181:7206–7211. <https://doi.org/10.1128/JB.181.23.7206-7211.1999>.
 40. Janeček Š, Svensson B, MacGregor EA. 2014. α -Amylase: an enzyme specificity found in various families of glycoside hydrolases. *Cell Mol Life Sci* 71:1149–1170. <https://doi.org/10.1007/s00018-013-1388-z>.
 41. Ravcheev DA, Godzik A, Osterman AL, Rodionov DA. 2013. Polysaccharides utilization in human gut bacterium *Bacteroides thetaiotaomicron*: comparative genomics reconstruction of metabolic and regulatory networks. *BMC Genomics* 14:873. <https://doi.org/10.1186/1471-2164-14-873>.
 42. Rogers TE, Pudlo NA, Koropatkin NM, Bell JSK, Moya Balasch M, Jasker K, Martens EC. 2013. Dynamic responses of *Bacteroides thetaiotaomicron* during growth on glycan mixtures. *Mol Microbiol* 88:876–890. <https://doi.org/10.1111/mmi.12228>.
 43. Lammerts van Bueren A, Saraf A, Martens EC, Dijkhuizen L. 2015. Differential metabolism of exopolysaccharides from probiotic lactobacilli by the human gut symbiont *Bacteroides thetaiotaomicron*. *Appl Environ Microbiol* 81:3973–3983. <https://doi.org/10.1128/AEM.00149-15>.
 44. D'Elia JN, Salyers AA. 1996. Contribution of a neopullulanase, a pullulanase, and an α -glucosidase to growth of *Bacteroides thetaiotaomicron* on starch. *J Bacteriol* 178:7173–7179. <https://doi.org/10.1128/jb.178.24.7173-7179.1996>.
 45. Mikheenko A, Prijbelski A, Savelyev V, Antipov D, Gurevich A. 2018. Versatile genome assembly evaluation with QUAST-LG. *Bioinformatics* 34:i142–i150. <https://doi.org/10.1093/bioinformatics/bty266>.
 46. St John FJ, González JM, Pozharski E. 2010. Consolidation of glycosyl hydrolase family 30: a dual domain 4/7 hydrolase family consisting of two structurally distinct groups. *FEBS Lett* 584:4435–4441. <https://doi.org/10.1016/j.febslet.2010.09.051>.
 47. Nguyen TLA, Vieira-Silva S, Liston A, Raes J. 2015. How informative is the mouse for human gut microbiota research? *Dis Model Mech* 8:1–16. <https://doi.org/10.1242/dmm.017400>.
 48. Koropatkin NM, Smith TJ. 2010. SusG: a unique cell-membrane-associated α -amylase from a prominent human gut symbiont targets complex starch molecules. *Structure* 18:200–215. <https://doi.org/10.1016/j.str.2009.12.010>.
 49. Naumoff DG. 2005. GH97 is a new family of glycoside hydrolases, which is related to the α -galactosidase superfamily. *BMC Genomics* 6:112. <https://doi.org/10.1186/1471-2164-6-112>.
 50. Boraston AB, Bolam DN, Gilbert HJ, Davies GJ. 2004. Carbohydrate-binding modules: fine-tuning polysaccharide recognition. *Biochem J* 382:769–781. <https://doi.org/10.1042/BJ20040892>.
 51. Kim YM, Yamamoto E, Kang MS, Nakai H, Saburi W, Okuyama M, Mori H, Funane K, Momma M, Fujimoto Z, Kobayashi M, Kim D, Kimura A. 2012. *Bacteroides thetaiotaomicron* VPI-5482 glycoside hydrolase family 66 homolog catalyzes dextranolytic and cyclization reactions. *FEBS J* 279:3185–3191. <https://doi.org/10.1111/j.1742-4658.2012.08698.x>.
 52. Kim MJ, Lee SB, Lee HS, Lee SY, Baek JS, Kim D, Moon TW, Robyt JF, Park KH. 1999. Comparative study of the inhibition of α -glucosidase, α -amylase, and cyclomaltodextrin glucanotransferase by acarbose,

- isoaccharose, and acarviosine-glucose. *Arch Biochem Biophys* 371: 277–283. <https://doi.org/10.1006/abbi.1999.1423>.
53. Santilli AD, Dawson EM, Whitehead KJ, Whitehead DC. 2018. Nonmicrobical small molecule inhibition of polysaccharide metabolism in human gut microbes: a potential therapeutic avenue. *ACS Chem Biol* 13:1165–1172. <https://doi.org/10.1021/acscchembio.8b00309>.
 54. Rasko DA, Rosovitz MJ, Myers GSA, Mongodin EF, Fricke WF, Gajer P, Crabtree J, Sebaihia M, Thomson NR, Chaudhuri R, Henderson IR, Sperandio V, Ravel J. 2008. The pangenome structure of *Escherichia coli*: comparative genomic analysis of *E. coli* commensal and pathogenic isolates. *J Bacteriol* 190:6881–6893. <https://doi.org/10.1128/JB.00619-08>.
 55. Medini D, Donati C, Tettelin H, Massignani V, Rappuoli R. 2005. The microbial pan-genome. *Curr Opin Genet Dev* 15:589–594. <https://doi.org/10.1016/j.gde.2005.09.006>.
 56. Scholz M, Ward DV, Pasolli E, Tolio T, Zolfo M, Asnicar F, Truong DT, Tett A, Morrow AL, Segata N. 2016. Strain-level microbial epidemiology and population genomics from shotgun metagenomics. *Nat Methods* 13: 435–438. <https://doi.org/10.1038/nmeth.3802>.
 57. Truong DT, Tett A, Pasolli E, Huttenhower C, Segata N. 2017. Microbial strain-level population structure and genetic diversity from metagenomes. *Genome Res* 27:626–638. <https://doi.org/10.1101/gr.216242.116>.
 58. Delmont TO, Eren AM. 2018. Linking pangenomes and metagenomes: the *Prochlorococcus* metapangenome. *PeerJ* 6:e4320. <https://doi.org/10.7717/peerj.4320>.
 59. Segata N. 2018. On the road to strain-resolved comparative metagenomics. *mSystems* 3:e00190-17. <https://doi.org/10.1128/mSystems.00190-17>.
 60. Quince C, Delmont TO, Raguideau S, Alneberg J, Darling AE, Collins G, Eren AM. 2017. DESMAN: a new tool for de novo extraction of strains from metagenomes. *Genome Biol* 18:1–22. <https://doi.org/10.1186/s13059-017-1309-9>.
 61. Merino S, Tomás JM. 2015. Bacterial capsules and evasion of immune responses. *eLS* <https://doi.org/10.1002/9780470015902.a0000957.pub4>.
 62. Miller RA, Harrison DE, Astle CM, Baur JA, Boyd AR, de Cabo R, Fernandez E, Flurkey K, Javors M, Nelson JF, Orihuela CJ, Pletcher S, Sharp ZD, Sinclair DA, Starnes JW, Wilkinson JE, Nadon NL, Strong R. 2011. Rapamycin, but not resveratrol or simvastatin, extends life span of genetically heterogeneous mice. *J Gerontol A Biol Sci Med Sci* 66:191–201. <https://doi.org/10.1093/gerona/glq178>.
 63. Köster J, Rahmann S. 2012. Snakemake scalable bioinformatics workflow engine. *Bioinformatics* 28:2520–2522. <https://doi.org/10.1093/bioinformatics/bts480>.
 64. Xu H, Luo X, Qian J, Pang X, Song J, Qian G, Chen J, Chen S. 2012. FastUniq: a fast *de novo* duplicates removal tool for paired short reads. *PLoS One* 7:e52249. <https://doi.org/10.1371/journal.pone.0052249>.
 65. Buffalo V. 2018. Scythe—a Bayesian adapter trimmer. <https://github.com/vsbuffalo/scythe>.
 66. Joshi NA, Fass JN. 2011. Sickle: a sliding-window, adaptive, quality-based trimming tool for FastQ files (version 1.33). <https://github.com/najoshi/sickle>.
 67. Li D, Liu CM, Luo R, Sadakane K, Lam TW. 2015. MEGAHIT: an ultra-fast single-node solution for large and complex metagenomics assembly via succinct de Bruijn graph. *Bioinformatics* 31:1674–1676. <https://doi.org/10.1093/bioinformatics/btv033>.
 68. Langmead B, Salzberg SL. 2012. Fast gapped-read alignment with Bowtie 2. *Nat Methods* 9:357–359. <https://doi.org/10.1038/nmeth.1923>.
 69. Alneberg J, Bjarnason BS, De Bruijn I, Schirmer M, Quick J, Ijaz UZ, Lahti L, Loman NJ, Andersson AF, Quince C. 2014. Binning metagenomic contigs by coverage and composition. *Nat Methods* 11:1144–1146. <https://doi.org/10.1038/nmeth.3103>.
 70. Pedregosa F, Varoquaux G, Gramfort A, Michel V, Thirion B, Grisel O, Blondel M, Prettenhofer P, Weiss R, Dubourg V, Vanderplas J, Passos A, Cournapeau D, Brucher M, Perrot M, Duchesnay É. 2011. Scikit-learn: machine learning in Python. *J Mach Learn Res* 12:2825–2830. <http://www.jmlr.org/papers/v12/pedregosa11a.html>.
 71. Walker BJ, Abeel T, Shea T, Priest M, Abouelliel A, Sakthikumar S, Cuomo CA, Zeng Q, Wortman J, Young SK, Earl AM. 2014. Pilon: an integrated tool for comprehensive microbial variant detection and genome assembly improvement. *PLoS One* 9:e112963. <https://doi.org/10.1371/journal.pone.0112963>.
 72. Parks DH, Imelfort M, Skennerton CT, Hugenholtz P, Tyson GW. 2015. CheckM: assessing the quality of microbial genomes recovered from isolates, single cells, and metagenomes. *Genome Res* 25:1043–1055. <https://doi.org/10.1101/gr.186072.114>.
 73. Seemann T. 2014. Prokka: rapid prokaryotic genome annotation. *Bioinformatics* 30:2068–2069. <https://doi.org/10.1093/bioinformatics/btu153>.
 74. Hyatt D, Chen G-L, LoCascio PF, Land ML, Larimer FW, Hauser LJ. 2010. Prodigal: prokaryotic gene recognition and translation initiation site identification. *BMC Bioinformatics* 11:119. <https://doi.org/10.1186/1471-2105-11-119>.
 75. Punta M, Coghill PC, Eberhardt RY, Mistry J, Tate J, Boursnell C, Pang N, Forslund K, Ceric G, Clements J, Heger A, Holm L, Sonnhammer ELL, Eddy SR, Bateman A, Finn RD. 2012. The Pfam protein families database. *Nucleic Acids Res* 40:D290–D301. <https://doi.org/10.1093/nar/gkr1065>.
 76. Eddy SR. 2011. Accelerated profile HMM searches. *PLoS Comput Biol* 7: e1002195. <https://doi.org/10.1371/journal.pcbi.1002195>.
 77. Eddy SR. 2009. A new generation of homology search tools based on probabilistic inference. *Genome Inform* 23:205–211.
 78. Juncker AS, Willenbrock H, von Heijne G, Brunak S, Nielsen H, Krogh A. 2003. Prediction of lipoprotein signal peptides in Gram-negative bacteria. *Protein Sci* 12:1652–1662. <https://doi.org/10.1110/ps.0303703>.
 79. Seydel A, Gounon P, Pugsley AP. 1999. Testing the “+2 rule” for lipoprotein sorting in the *Escherichia coli* cell envelope with a new genetic selection. *Mol Microbiol* 34:810–821. <https://doi.org/10.1046/j.1365-2958.1999.01647.x>.
 80. Rahman O, Cummings SP, Harrington DJ, Sutcliffe IC. 2008. Methods for the bioinformatic identification of bacterial lipoproteins encoded in the genomes of Gram-positive bacteria. *World J Microbiol Biotechnol* 24: 2377–2382. <https://doi.org/10.1007/s11274-008-9795-2>.
 81. Petersen TN, Brunak S, Von Heijne G, Nielsen H. 2011. SignalP 4.0: discriminating signal peptides from transmembrane regions. *Nat Methods* 8: 785–786. <https://doi.org/10.1038/nmeth.1701>.
 82. Li W, Godzik A. 2006. Cd-hit: a fast program for clustering and comparing large sets of protein or nucleotide sequences. *Bioinformatics* 22:1658–1659. <https://doi.org/10.1093/bioinformatics/btl158>.
 83. Fu L, Niu B, Zhu Z, Wu S, Li W. 2012. CD-HIT: accelerated for clustering the next-generation sequencing data. *Bioinformatics* 28:3150–3152. <https://doi.org/10.1093/bioinformatics/bts565>.
 84. Buchfink B, Xie C, Huson DH. 2015. Fast and sensitive protein alignment using DIAMOND. *Nat Methods* 12:59–60. <https://doi.org/10.1038/nmeth.3176>.
 85. Enright AJ, Van Dongen S, Ouzounis CA. 2002. An efficient algorithm for large-scale detection of protein families. *Nucleic Acids Res* 30:1575–1584. <https://doi.org/10.1093/nar/30.7.1575>.
 86. Terrapon N, Lombard V, Gilbert HJ, Henrissat B. 2015. Automatic prediction of polysaccharide utilization loci in *Bacteroidetes* species. *Bioinformatics* 31:647–655. <https://doi.org/10.1093/bioinformatics/btv176>.
 87. Selengut JD, Haft DH, Davidsen T, Ganapathy A, Gwinn-Giglio M, Nelson WC, Richter AR, White O. 2007. TIGRFAMs and Genome Properties: tools for the assignment of molecular function and biological process in prokaryotic genomes. *Nucleic Acids Res* 35:D260–D264. <https://doi.org/10.1093/nar/gkl1043>.
 88. Price MN, Dehal PS, Arkin AP. 2010. FastTree 2—approximately maximum-likelihood trees for large alignments. *PLoS One* 5:e9490. <https://doi.org/10.1371/journal.pone.0009490>.