

Distinct polysaccharide growth profiles of human intestinal *Prevotella copri* isolates

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Summary:

The gut-dwelling *Prevotella copri*, the most prevalent *Prevotella* species in human gut, have been associated with diet as well as disease, but our understanding of their diversity and functional roles remains rudimentary, as studies have been limited to 16S and metagenomic surveys and experiments using a single type strain. Here, we describe the genomic diversity of 83 *P. copri* isolates from 11 donors. We demonstrate that genomically distinct isolates, which can be categorized into different *P. copri* complex clades, utilize defined sets of plant polysaccharides that can be predicted in part from their genomes and from metagenomic data. This study thus reveals both genomic and functional differences in polysaccharide utilization between several novel human intestinal *P. copri* strains.

Keywords:

human microbiome, intestinal commensals, whole genome sequencing, diet

Introduction:

Humans live in close association with a teeming bacterial ecosystem known as the gut microbiome. The diverse inhabitants of this extraordinary environment perform beneficial and remarkable services for their host, out-competing harmful bacteria and digesting the majority of dietary fiber (El Kaoutari et al., 2013; Martens et al., 2008). The relative abundance of two genera, *Bacteroides* and *Prevotella*, is inversely correlated in the intestine (Arumugam et al., 2011; Costea et al., 2018). Although a *Bacteroides*-dominated microbiome is more common, *Prevotella* are present at >10% relative abundance in the stool of 10-25% of healthy American

and European individuals (Koren et al., 2013). Compared to at least 25 species of *Prevotella* in the oral cavity, and at least 17 species of intestinal *Bacteroides*, only five *Prevotella* species have been reported in human intestine, with *Prevotella copri* being the most abundant (Accetto and Avgustin, 2015; Ferrocino et al., 2015; Ibrahim et al., 2017; Li et al., 2009; Lin et al., 2013; Liu et al., 2018). Although mechanistic models of metabolism, growth, and colonization have been established for several intestinal *Bacteroides* and oral *Prevotella*, studies of intestinal *Prevotella* have lacked experimental tools and relied heavily on only two type strains, *Prevotella copri* and *Prevotella stercorea* (Hayashi et al., 2007).

Many researchers have observed *Prevotella*'s presence and abundance by 16S sequencing, but its genomic diversity and biological properties have not been described. Because *Prevotella copri* are highly sensitive to oxygen, and difficult to isolate and cultivate, previous studies have been limited to the single commercially available type strain, DSM 18205, to perform metagenomic comparisons and draw experimental conclusions. *Prevotella* spp. have extensive gene acquisition and loss (Gupta et al., 2015; Zhu et al., 2015). In fact, accessory genes within the *P. copri* DSM 18205 genome account for about 40% of all genes encoded within the genome, the highest proportion of accessory genes identified in a comparison of 11 gut-associated species (Zhu et al., 2015). This estimate should be revised, however, since a metagenomic study (supplemented by whole genome sequencing data from the present study) affirms that *P. copri* is diverse, comprising four genomically distinct clades (Tett et al., 2019). If species can be determined by average nucleotide identity (ANI) difference, these four clades of *P. copri* could be considered four distinct species, as they have >10% ANI difference (Tett et al., 2019). Further improvements in genomic comparisons of *P. copri* may be expected soon, since another study reports a closed genome of *P. copri*, generated from long-read sequencing of whole microbiomes

(Moss and Bhatt, 2018). Whether the genomic diversity in human *P. copri* translates to functional diversity has not yet been fully explored. This is particularly relevant in light of elevated frequency of *P. copri* colonization in some cohorts of treatment-naïve rheumatoid arthritis patients (Maeda et al., 2016; Scher et al., 2013). A better understanding of whether there are functional differences in strains from patients compared to healthy subjects will contribute to determining whether there is a causal relationship between *P. copri* and this autoimmune disease.

Several studies have found associations between diets and the presence of intestinal *Prevotella*. *Prevotella* have been positively associated with diets high in plant fiber and carbohydrates, and negatively associated with fat and amino acid diets (De Filippo et al., 2010; Fragiadakis et al., 2019; Ruengsomwong S, 2016; Smits et al., 2017; Wu et al., 2011). Genes associated with carbohydrate catabolism were identified in *P. copri* metagenomes and correlated with vegan diets (De Filippis et al., 2019). The plant polysaccharide xylan has been used to select for growth of human-associated *Prevotella* (Tan et al., 2018). *Prevotella* isolates from livestock grow in the presence of plant polysaccharides and encode gene products capable of breaking down these polysaccharides (Accetto and Avgustin, 2019). Thus, diet may be an important factor promoting growth of intestinal *Prevotella*. Whether human-associated *P. copri* are uniquely suited to digest dietary components highly abundant in plant-based diets remains unsubstantiated by experimental and genomic evidence.

Intestinal commensal bacteria have a system of genes, often clustered, whose products collaborate to harness energy from carbohydrates present in the host's intestinal lumen. Extensive and careful work in *Bacteroides* species has identified several polysaccharide utilization loci (PULs) that are specific for glycans derived from dietary plant and animal sources, as well as endogenous host sources (Martens et al., 2008; Martens et al., 2011). Upon

exposure to the target polysaccharide, genes encoded in the PUL are upregulated. Bacteria lacking one of several critical genes in any polysaccharide-specific PUL fail to grow when the polysaccharide is provided as the sole carbon source (Martens et al., 2011; Raghavan et al., 2014). PULs have been linked to diet and subsequently exploited in experimental systems. Notably, a seaweed porphyran-utilizing PUL was horizontally transferred from marine bacteria to intestinal bacteria of seaweed-ingesting hosts (Hehemann et al., 2010). This PUL was experimentally introduced into *B. thetaiotaomicron*, and seaweed-selective growth was demonstrated in laboratory mice (Shepherd et al., 2018). Although by far the most extensive work on PULs has been performed in *Bacteroides* species, PULs have been annotated in *Prevotella* type strains and used to predict growth on plant polysaccharides in *Prevotella* isolates from livestock (Accetto and Avgustin, 2015, 2019). Despite their association with plant diet and PUL predictions, the functional capabilities of human-associated intestinal *Prevotella* have not been tested, due to a dearth of cultured isolates.

Previous studies have been limited to analysis and experimentation with the available genomes and strains of *Prevotella* species found in human, two of which were isolated from the same individual (Hayashi et al., 2007). Here, we report whole genome sequencing of 83 *Prevotella copri* isolates from the stool of eleven individuals. We observe extensive genomic diversity of *P. copri* isolates within and between hosts. This genomic variation was exemplified by specific gene clusters and linked to functional growth of isolates on predicted plant polysaccharides, explaining the association of colonization of *Prevotella* with a plant-rich diet.

Results:

Isolation of *Prevotella* from human stool

To investigate the genomic diversity of *Prevotella copri* and also identify strains for functional analysis, we developed a qPCR screening and isolation strategy for *P. copri* from human stool (**Methods**). *P. copri* was identified in 31/63 individuals by qPCR (**Figure S1A**). 16S rRNA gene sequencing of the hypervariable V3-V4 region found *P. copri* in 41/63 individuals, with 18/63 individuals harboring *P. copri* at levels >10% relative abundance (**Figure S1B**). From the V3-V4 16S rRNA gene data, five *P. copri* 16S variants were identified (**Figure S1B, Supplemental file 1**). The *P. copri* 16S variants displayed 2-6 single-nucleotide variants (SNVs) within the 253 bp region, corresponding to >97% sequence-sequence identity (Callahan et al., 2016). *P. copri* detection by qPCR positively correlated with *P. copri* detection by 16S sequencing (**Figure S1C**). We collected 83 isolates from eleven individuals harboring *P. copri* (**Figure S1B**), one of whom had <1% *P. copri* (**Methods**).

Variability in *P. copri* isolates

Our earlier study indicated differences in the *P. copri* genomes from new onset rheumatoid arthritis and healthy individuals (Scher et al., 2013). Other studies have since described genetic variation between strains from healthy individuals (De Filippis et al., 2019; Zhao et al., 2019). We assessed whether genomic diversity existed between *P. copri* isolates from different individuals, or, furthermore, between isolates from the same individual, by sequencing the full genomes of the isolates. Alignment of filtered isolate sequencing reads to the *P. copri* PanPhlAn database revealed that fewer than 60% of reads mapped to the reference genome, indicating unexpected and unexplored diversity of *P. copri* genomes in our isolates (**Figure 1A**) (Scholz et al., 2016).

Evaluation of the *de novo* assembled genomes revealed low contamination and similar genome size among all isolates (**Table S1**) (Bankevich et al., 2012; Parks et al., 2015). Phylogenetic analysis based on the 16S rRNA gene revealed that most of the isolates clustered with *Prevotella copri* DSM 18205 (**Figure 1B**). One isolate clustered with *Paraprevotella*, and three isolates clustered with other species of *Prevotella*; these four isolates were excluded from the rest of the analysis. A separate phylogenetic analysis based on 30 conserved ribosomal proteins further confirmed that the isolates clustered with *P. copri*, and also revealed distinct clades (**Figure 1C**).¹ Comparison with a catalog of *P. copri* metagenomes confirmed that most isolates were in a main clade (clade A), four isolates were in another clade (clade C), and one isolate each was in two additional clades (clades B and D) (Tett et al., 2019). Hosts AQ, A, and AA harbored isolates belonging to more than one clade (**Figure 1C1**). Enzyme activity and acidification assays revealed functional differences between select isolates from different clades and within clade A (**Table 1**).

***P. copri* genomic diversity between hosts**

Despite close phylogenetic placement, most isolates exhibited striking genomic dissimilarities to the *P. copri* reference strain (**Figure 1A**). To better understand the differences in the *P. copri* genomes, we compared the presence and absence of predicted ORFs in all the isolates (**Figure 1D**). This analysis revealed a core genome of approximately 1750 genes, suggesting accessory genomes ranging from approximately 1250-2250 genes per isolate.

Clustering of the isolates by presence of predicted ORFs revealed that isolates often clustered by donor. When isolates from the same donor fell into the same clade, accessory genomes were often unique to isolates from individual hosts, as visualized by blocks of predicted ORFs in the

accessory genomes in **Figure 1D**. Host AK had isolates with distinct accessory genomes within the same clade (**Figures 1C and 1D**). Annotation of the genomes revealed that, among other functions, many of the host-specific genes were involved in polysaccharide transport, as indicated by the frequency of *susC* genes (**Figure 1D**).

Polysaccharide utilization prediction

SusC is the bacterial transmembrane protein that transports polysaccharides into the periplasm. SusC collaborates with the outer-membrane glycan-binding protein SusD, as well as other proteins, for uptake and breakdown of polysaccharides. Conveniently, the genes for this polysaccharide utilization machinery usually occur grouped in close proximity to one another in the genome in polysaccharide utilization loci (PULs) in Bacteroidetes, although occasionally genes encoding polysaccharide catabolism enzymes are located elsewhere in the genome (Sonnenburg et al., 2010). We hypothesized that the genetic diversity of *susC* genes might indicate functional diversity in polysaccharide utilization by the isolates. We surveyed the *susC* and surrounding annotated genes in the isolate genomes to predict the repertoire of PULs and, by extension, candidate polysaccharide substrates for the *P. copri* isolates.

By clustering *susC* genes at 90% identity, and examining the neighboring gene products within each genome, we identified 125 *susC* gene clusters with a neighboring *susD* gene, which we called predicted PULs (pPULs) (**Methods, Table S2**). 87 pPULs could be categorized based on the presence of annotated genes encoding glycan-degrading functions flanking the *susC* and *susD* in each pPUL (**Figure 2A**). Due to a high frequency of genes encoding hypothetical proteins, we were unable to categorize or predict candidate substrates for 38 pPULs. Genes annotated to encode similar carbohydrate-active enzymes were found in pPULs with distinct

susC clusters in different isolate genomes. For example, xylanases were found in close proximity to *susC* clusters 2, 65, and the adjacent clusters 100 and 140. *SusC* cluster 2 was only in isolate genomes from individual BU, *susC* cluster 65 was only in the genome of isolate A622, and *susC* clusters 100 and 140 were only in isolate genomes from individual AA. In contrast with several *Bacteroides* species, which encode PULs that catabolize polysaccharides from both animal and plant dietary glycans, all annotated enzymes in the *P. copri* isolate pPULs were predicted to catabolize dietary plant glycans and endogenous host mucin, but none were predicted to catabolize dietary animal glycans (Martens et al., 2011; Salyers AA, 1982). Consistent with this prediction, the V1-V3 16S sequences of select isolates were most closely related to two previously reported non-omnivore-associated *P. copri* 16S sequences, P10 and P19 (De Filippis et al., 2016) (**Figure S2A, Supplemental file 2**).

Consistent with the 17 predicted PULs for *P. copri* DSM 18205, the number of predicted PULs per isolate ranged from 14-28, with a median of 23 (Terrapon et al., 2018). Intestinal *Prevotella* have been reported to utilize xylan and, as expected, pPULs containing xylanases were ubiquitously represented throughout the isolate genomes (**Figure 2A**) (Tan et al., 2018). A survey of the pPULs containing annotated xylan catabolism enzymes revealed additional features: all isolates contained pPULs with xylan catabolism enzymes; each genome contained 4-5 xylan catabolism enzymes that were present in pPULs (**Figure S2B**); and *SusC/D* pairs in pPULs containing xylan catabolism enzymes often occurred in tandem.

Polysaccharide utilization *in vitro*

Based on the pPUL analysis, we predicted that *P. copri* isolates would grow in the presence of plant polysaccharides targeted by the enzymes encoded by the annotated genes in the pPULs

(candidate substrates in **Figure 2A**). We therefore tested the ability of *P. copri* isolates to grow when plant polysaccharides were provided as the sole carbon source *in vitro* (**Methods**). Growth on individual polysaccharide substrates revealed functional diversity between isolates (**Figure 3A, 3B, Figure S3A**). For example, isolate AA108 reached a high OD when pullulan was provided as the sole carbon source, but isolates P54 and T2112 did not. Interestingly, isolates from the same individual grew in the presence of different polysaccharides: compared to isolate AK1212, isolate AK2718 grew to a higher OD in the presence of PGA and rhamnogalacturonan I, and a lower OD in the presence of beta-glucan and arabinan. None of the isolates grew in the presence of animal polysaccharides chondroitin or heparin or the proteoglycan mucin (**Figure 3A, Figure S3A**), indicating that *Prevotella* isolates could be cultured in the presence of glycans from dietary plant sources, but not dietary animal or endogenous host sources. Growth (or lack of growth) on particular polysaccharides distinguished isolates from each clade (**Figure 3B, Table S3**). In particular, arabinan, levan, inulin, xyloglucan, beta-glucan, and glucomannan growth distinguished isolates from each clade. All isolates except AQ1173 (which fell into clade B) grew on arabinoxylan and arabinan (**Figures 3A and 3B**). Isolate AQ1173 also failed to grow on xyloglucan and glucomannan. Isolate A622 (which fell into clade D) failed to grow on both levan and inulin. Tested isolates from clade C failed to grow on beta-glucan, xyloglucan, glucomannan, and rhamnogalacturonan I. Tested isolates from clade A grew well on most substrates, although there was a range of growth among clade A isolates on several plant polysaccharide sources.

Isolates AQ1173 and A622 were the only isolates that fell into clades B and D, respectively. Since growth data from single isolates limited our ability to generalize to those clades, we asked whether pPUL-containing enzymes targeting relevant polysaccharides were present or absent in

these clades using *P. copri* metagenome-assembled genomes (MAGs) reconstructed from metagenomic datasets (Tett et al., 2019) (**Methods, Figure 3C**). Consistent with our observation that isolate AQ1173 (clade B) lacked pPULs containing enzymes that target arabinan, arabinoxylan, and glucomannan and also did not grow on those polysaccharides (**Figures 2A, 3A, and 3B**), the clade B *P. copri* MAGS lacked pPULs with CAZymes capable of degrading those polysaccharides (**Figure 3C**). The clade B MAGs contained pPULs with CAZymes capable of degrading beta-glucan, however, consistent with our pPUL analysis and growth observations. pPULs with CAZymes capable of degrading glucomannan and arabinan were found in the clade D *P. copri* MAGs consistent with our predictions and growth observations for isolate A622 (Clade D).

In order to determine which pPUL respond to specific polysaccharides, we asked which *susC* genes were upregulated in *P. copri* DSM 18205 during growth on arabinan, xyloglucan, corn xylan, beechwood xylan, and arabinoxylan. A screen for all 22 *susC* genes in *P. copri* DSM 18205 revealed upregulation of distinct *susC* genes in each polysaccharide growth condition (**Figures 3D and S3B**). Upregulation of the *susC* gene is considered a proxy for expression of the whole PUL, therefore, the pPULs associated with the *susC* gene likely target the tested polysaccharides. Consistent with the presence of arabinosidase genes in close proximity to the *susC* gene, *susC* cluster 55 was upregulated in the presence of arabinan. Similarly, the tandem *susC* clusters 135 and 83, which are flanked by genes encoding xylanases, were upregulated in the presence of both corn and beechwood xylans, and to a lesser extent in the arabinoxylan condition. Analysis of pPUL structure revealed synteny between genes flanking *susC* clusters 83, 135, 100, 140, 71, and 138 (**Figure 2B**). Surprisingly, *susC* cluster 76 was upregulated in the presence of xyloglucan. *SusC* cluster 76 was flanked by genes encoding two mannosidases and

several hypothetical proteins in all isolates. Further examination of the locus revealed a hypothetical protein with homology to GH5. GH5 has endo-beta-glucanase activity, which could potentially target xyloglucan. Synteny was also observed between susC clusters 76 and 73, and surrounding genes, suggesting that the cluster 73 pPUL may also target xyloglucan (**Figure S3C**).

Polysaccharide growth predictions

Using a combination of genomic predictions, gene expression data, and homology, we evaluated how well the pPULs analysis predicted *in vitro* growth on candidate polysaccharides: 180 isolate/polysaccharide conditions matched predictions while 56 did not (**Figure S3D**). We observed 21 instances in which we did not predict particular polysaccharide utilization by the isolate, yet we observed growth. For 18 of these cases of unpredicted growth, we found genes encoding relevant polysaccharide breakdown enzymes elsewhere in the genome. Perhaps the genes are part of fragmented PULs (as has been observed in *Bacteroides* fructan PULs (Sonnenburg et al., 2010)), or are simply located on a contig separate from the rest of the PUL. More puzzlingly, we observed isolates that failed to grow on a particular substrate, despite presence of a corresponding intact pPUL in the genome. For example, we predicted PULs with arabinogalactan catabolism genes in most of genomes of clade A, yet most failed to grow on this substrate, and those that did, particularly *P. copri* isolate BVe41210, grew poorly. Perhaps *P. copri* isolates prefer a different form or source of arabinogalactan. Additionally, substrate predictions were based on gene annotations that may be incomplete or inaccurate. Overall, the pPUL analysis performed well in predicting growth. Welch's t-test on predicted growth or unpredicted growth had a *p*-value < 0.0001 (**Methods**).

Discussion:

In this study, we observed genomic and functional diversity of human intestinal *P. copri* isolates. Whole genome sequencing exposed diversity of *P. copri* isolates, demonstrated by the low percentage of isolate whole genome reads that mapped to the *P. copri* PanPhlAn database. Within-species genomic variation, both between individuals and within individuals, was demonstrated by genetic diversity of *susC* genes and PULs and linked to functional growth of isolates on plant polysaccharides. PULs predicted to target several polysaccharides in large metagenomic datasets support our predictions and growth data for representative isolates, indicating that the results of this study may be generalizable beyond the isolates shown here.

Variability in *P. copri* genomes

Despite 97% sequence identity in the 16S rRNA gene, and conservation of ribosomal protein sequences, the *P. copri* isolates vary in their genomes, enzyme activities, and polysaccharide utilization profiles, suggesting that they could be considered distinct species by some definitions and the same species by others. ORF presence/absence among isolates revealed host-specific genes in the accessory genome, several of which were *susC* transport protein genes. The variability of the *susC* genes associated with *P. copri* strains in different hosts is intriguing, given their common function of transporting glycans across the outer membrane, and could be the result of several factors, including selective pressures from the environment. SusC proteins from different isolates could also transport slightly different substrates, as has been observed in two closely related *B. thetaiotaomicron* strains (Joglekar et al., 2018). Isolates from the same host but with different genomes suggest a distinct niche for each type of isolate. Hosts AQ, A, and AA

harbored isolates from different clades. Isolates from host AK all fell into the same clade and possessed two distinct accessory genomes. Perhaps one type of isolate inhabits the cecum, the other the colon; alternatively, polysaccharide cross-feeding may occur between *P. copri* isolates, as was demonstrated *in vitro* with *Bacteroides* species (Rakoff-Nahoum et al., 2014). In hosts with more than one strain of *P. copri*, representative isolates often complemented each other to grow in the presence of more polysaccharides than any individual isolate. This point was illustrated best by isolates from donor AQ: isolates AQ1173 and AQ1179 did not grow on xyloglucan or glucomannan, but AQ1149 grew on both polysaccharides (**Figure 4A**). These results suggest that the total *P. copri* population in hosts with multiple strains of *P. copri* was capable of catabolizing a greater diversity of polysaccharides than any individual *P. copri* strain. Indeed, a separate study found that non-Western microbiomes contained multiple *P. copri* clades, and that some carbohydrate active enzymes were differentially represented between clades (Tett et al., 2019). Whether microbiomes of individuals with only one *P. copri* strain completely lack the ability to break down additional polysaccharides or rely on other bacterial species to maintain a full complement of polysaccharide digestion should be the subject of future metagenomic studies.

***P. copri* presence in the human intestine may be diet-dependent**

In humans, differences in *P. copri* 16S sequences and metagenomes have been correlated with host diet (De Filippis et al., 2019; De Filippis et al., 2016; Fragiadakis et al., 2019; Smits et al., 2017). Although our study lacks host diet information, the pPUL analysis predicted and *in vitro* growth experiments confirmed that the *P. copri* isolates used only plant polysaccharides. A previous study found that particular *P. copri* V1-V3 16S sequences were associated with diet (De

Filippis et al., 2016). The V1-V3 sequences of select isolates from different clades in this study were most closely related to two non-omnivore-associated sequences, consistent with their growth on plant polysaccharides (**Figure S2A**). Although the dynamics of dietary and intestinal growth of *P. copri* within a complex microbiome require experimental validation, *in vitro* data presented here suggest that dietary plant polysaccharides promote *P. copri* growth, but dietary animal polysaccharides do not. These data provide a possible mechanism for the correlation between vegetable-rich diets and abundance of intestinal *Prevotella*: perhaps continuous availability of dietary plant polysaccharides is required to maintain *Prevotella* presence in the gut, and their absence may result in elimination of *Prevotella* from the human intestinal microbiome.

Prevotella polysaccharide utilization

Although the pPUL analysis predicted growth surprisingly well, incorrect predictions may indicate interesting biology or areas in need of technical improvements. In cases in which PULs were predicted but growth on a corresponding candidate polysaccharide was not observed, perhaps mutations in the genes encoding the enzymes or regulators have rendered the gene products non-functional. Future studies should aim to evaluate how often and how rapidly such mutations might occur in intestinal commensals. Open genomes may account for some cases in which polysaccharide utilization was not predicted by the presence of a PUL, yet growth was observed. If a PUL were split across contigs, it would be impossible to predict a candidate substrate with current approaches. Recent closure of a *P. copri* genome from metagenomic analysis may help reduce such instances in future studies (Moss and Bhatt, 2018).

Utilization of xylan, which is found in cereal grains, has been repeatedly established in *Prevotella* species and in *P. copri* specifically (Accetto and Avgustin, 2019; da Silva et al., 2012; Dodd et al., 2011; Tan et al., 2018). Two distinct PULs encoding enzymes not previously associated with xylan catabolism were identified for corn xylan, birchwood xylan, and arabinoxylan in multiple *B. ovatus* and *B. xylanisolvens* strains (Rogowski et al., 2015). The numerous xylan-degrading enzymes identified in pPULs in *P. copri* isolates in this study suggest that *P. copri* may have an expanded xylan degrading enzyme repertoire, and possibly a superior ability to target xylans compared to other intestinal bacteria.

Xyloglucan is an abundant polysaccharide in vascularized plants, seeds, and food additives, yet xyloglucan utilization by known intestinal microbes is relatively rare, demonstrated only in a low proportion of *Bacteroides* species (Larsbrink et al., 2014; Nishinari et al., 2007). Despite its abundance in the human diet, xyloglucan utilization was difficult to predict from *P. copri* genome sequence alone (**Figure S3C, 2A, Table S2**). Gene expression analysis of *susC* genes in *P. copri* grown on xyloglucan suggests that gene products in the pPUL catabolize xyloglucan. Isolates with synteny in PULs grew in the presence of xyloglucan. Given the rarity of xyloglucan catabolism genes reported to date in *Bacteroides*, perhaps *P. copri* is responsible for xyloglucan degradation in hosts lacking *Bacteroides* species possessing xyloglucan-breakdown genes.

Implications

The collection of *P. copri* isolates and their corresponding polysaccharide utilization repertoire described herein enable future experimentation *in vitro* and *in vivo* with isolates whose cultivation was previously elusive perhaps because of the absence of their required specific plant polysaccharides. The human genome encodes enzymes that break down only starch, sucrose, and

lactose (El Kaoutari et al., 2013). Carbohydrates make up 45-65% of typical human diets (Board., 2005). Therefore, the ability of intestinal microbes to break down polysaccharides is critical for human survival. Whether *P. copri* strain-type and presence is meaningful to human disease, or simply an indication of diet, requires further investigation. This study was underpowered to assess differences between healthy and disease *P. copri* genomes. Future studies that aim to assess human microbiome interactions with disease will require dietary intervention or, at minimum, collection of patient dietary habits or records. To avoid confounding effects of diet, future metagenomic analyses may better illuminate disease-associated genes by excluding genes known to interact with host diet. If *P. copri* does exacerbate or ameliorate disease in specific populations, diet may be readily manipulated to decrease or increase *P. copri* growth. Establishment of polysaccharide substrate preferences for genetically diverse human *P. copri* isolates allows for informed experimentation *in vitro* and *in vivo*, as well as clinical study design and interpretation.

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Author contributions:

HF-P, VR, and CM designed the study with input from DRL and RB. HF-P isolated *Prevotella* and designed and performed all experiments, with isolation strategy by LMC. JUS provided stool samples from NORA patients. CM analyzed 16S and genome sequencing data. VR predicted PULs from the isolate genomes and helped design growth experiments. AT and NS identified *susC* clusters in metagenomic data. CG provided technical assistance for growth experiments and analysis of PULs. AW and JDW-G helped analyze genome and growth data. HF-P wrote the manuscript with contributions from VR, CM, and DRL, and input from all other authors.

Declaration of interests:

The authors declare no competing interests.

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KEY RESOURCES TABLE**

REAGENT or RESOURCE	SOURCE	IDENTIFIER
Biological Samples		
Fecal samples from new onset rheumatoid arthritis patients and healthy controls	This study	N/A
Critical commercial assays		
PowerSoil DNA isolation kit	Qiagen	Cat No./ID: 12888-100
TruSeq DNA PCR-free Library Prep Kit	Illumina, California, USA	20015962
Wheaton™ CryoELITE™ Cryogenic Storage Vials	Fisher Scientific	02-912-737
FALCON 14 ml Polypropylene Round-Bottom Tubes	Fisher Scientific	352059
Titertube® Micro Test Tubes	Bio-Rad	2239391
RNeasy Mini Kit	Qiagen	74104
RNA Protect Bacteria Reagent	Qiagen	76506
SuperScript™ IV First-Strand Synthesis System	ThermoFisher	18091050
API® ZYM	BioMerieux	25200
API® Rapid ID 32 A Microbial Identification Kit	BioMerieux	32300
API® 20 A Microbial Identification Kit	BioMerieux	20300
Deposited Data		
<i>P. copri</i> isolate genomes	This study	NCBI-BioProject: PRJNA559898 and PATRIC: https://www.patricbrc.org/
16S sequencing	This study	NCBI-BioProject: PRJNA559898
Experimental Models: Organisms/Strains		
<i>Prevotella copri</i> DSM 18205	DSMZ	DSM 18205
<i>Bacteroides ovatus</i> ATCC8482	Michael Fischbach	
<i>Bacteroides thetaiotaomicron</i> VPI-5482	Michael Fischbach	N/A
<i>P. copri</i> isolates	This study	N/A
Oligonucleotides		
Universal 16S Primer 340F: ACTCCTACGGGAGGCAGCAGT	Scher et al., 2013	N/A
Universal 16S Primer 515R: ATTACCGCGGCTGCTGGC	Scher et al., 2013	N/A
P_copri_screen_1_F: CCGGACTCCTGCCCTGCAA	Scher et al., 2013	N/A
P_copri_screen_1_R: GTTGCGCCAGGCACTGCGAT	Scher et al., 2013	N/A

P_copri_screen_2_F: AATAGCTTTGGTGGCGCTTT	This study	N/A
P_copri_screen_2_R: AGGTTCCCTCACCGCATACA	This study	N/A
P_copri_screen_3_F: TTGGGTTCGCTGATGTTACC	This study	N/A
P_copri_screen_3_R: TCTTCAGCAGAAGCGTTGGT	This study	N/A
P_copri_screen_4_F: AGGACGTTGACCAGCAGAAG	This study	N/A
P_copri_screen_4_R: GCATAGCGGTTTTGGCTGAG	This study	N/A
P_copri_screen_5_F: GAGACGGGCTGGAGTGATTT	This study	N/A
P_copri_screen_5_R: TCCGTGTGGCTTTCATGGAG	This study	N/A
P_copri_screen_6_F: ACCGTAAGGAGCGTGTCTG	This study	N/A
P_copri_screen_6_R: GTCGCGATTCTCTACTCCCG	This study	N/A
P_copri_screen_16S_7_F: TTTGGAGACAATGACGCCCT	This study	N/A
P_copri_screen_16S_7_R: ACTTAAGCCGACACCTCAG	This study	N/A
P_copri_screen_16S_8_F: CACRGTAACGATGGATGCC	Scher et al., 2013	N/A
P_copri_screen_16S_8_R: GGTCGGGTTGCAGACC	Scher et al., 2013	N/A
susC_123_PUL1116_F_1: CAAGTAAAGCCCGCAAGGAC	This study	N/A
susC_123_PUL1116_R_1: TTACCCTGAAGTGCTACCGC	This study	N/A
susC_129_PUL1802_F_1: TGGCTCTGTCAAGCGTAAGG	This study	N/A
susC_129_PUL1802_R_1: TGCCTTTCCCTGCATACCTG	This study	N/A
susC_130_PUL762_F_1: TGCTCGATGAGGTGGTTGTC	This study	N/A
susC_130_PUL762_R_1: TAACACC CGCAACCTTACCC	This study	N/A
susC_131_PUL1281_F_1: CGTGGCATTGTTTCGTCATCG	This study	N/A
susC_131_PUL1281_R_1: TCAGAGCCTCACCATTCTGC	This study	N/A
susC_132_PUL1287_F_1: TGCTGAAGACAACGAACCGA	This study	N/A

susC_132_PUL1287 R_1: TTTCACAACGTGCTTCTGCG	This study	N/A
susC_133_PUL484 F_1: TCGAGGGTGGCGACAATAAC	This study	N/A
susC_133_PUL484 R_1: CCATTAGAGGCACGGCTACC	This study	N/A
susC_135_PUL1745 F_1: TACGTGGGTGGTTGGAATGG	This study	N/A
susC_135_PUL1745 R_1: TCTCGCTAGGCATCACGTTG	This study	N/A
susC_102_PUL1218 F_2: GGTGCTACGCTCGTGATTC	This study	N/A
susC_102_PUL1218 R_2: TCACCACCCACATTGGCTAC	This study	N/A
susC_112_PUL792 F_1: TGACATGCAGAGTAGCGGTC	This study	N/A
susC_112_PUL792 R_1: GATGACACCATTGGCAGCAC	This study	N/A
susC_147_PUL2366 F_1: GTGCAGCCCGCTATATGGAC	This study	N/A
susC_147_PUL2366 R_1: CCACTGGGTTTTGTGTAGCG	This study	N/A
susC_156_PUL610 F_1: CTCAGAGCGTTGACCCTAGC	This study	N/A
susC_156_PUL610 R_1: TAATGCCCACTGCCTTACCG	This study	N/A
susC_1_PUL1911 F_1: GATAATGGAGCTGCCATCCG	This study	N/A
susC_1_PUL1911 R_1: CAGGCACTACAACAGAAGCG	This study	N/A
susC_21_PUL1668 F_1: AGCTCTTGACGAGGTGCGTTG	This study	N/A
susC_21_PUL1668 R_1: TGGGTGATGTTGACACCAGG	This study	N/A
susC_41_PUL1672 F_1: AAGGAGCTTGCTGAATCCC	This study	N/A
susC_41_PUL1672R_1: TGATGTTTACACCGGCAAGC	This study	N/A
susC_49_PUL1214 F_1: ATTTACCGTTCTGCAGCCG	This study	N/A
susC_49_PUL1214 R_1: CCAAGCGTCTGTTACGTTGG	This study	N/A
susC_55_PUL621 F_1: TCTTGGCTACAAGGAGCGTG	This study	N/A
susC_55_PUL621 R_1: ACAGAACCGGTAAGGTCTGC	This study	N/A

susC_67_PUL1810 F_1: CTGGCGAAACCTTGATTGGC	This study	N/A
susC_67_PUL1810 R_1: GCCATTAGCAGCTTTACGG	This study	N/A
susC_76_PUL296 F_1: TGTCTCAATGCTCAGACGG	This study	N/A
susC_76_PUL296 R_1: CTGCACCATGACACTTGCAC	This study	N/A
susC_83_PUL1747 F_1: TCTTGAAGACGACAACGCCA	This study	N/A
susC_83_PUL1747 R_1: CCTGTGGTCTGCGCAATAGA	This study	N/A
susC_90_PUL499 F_1: GCGCTCTGTA CTGTTGCAC	This study	N/A
susC_90_PUL499 R_1: AATGACTGTCTCACCGGCAG	This study	N/A
susC_165_PUL2751 F_1: TCGTGAAAGGTACCGTGCTG	This study	N/A
susC_165_PUL2751 R_1: CCTCTTGTGCCACCATACCG	This study	N/A
susC_166_PUL485 F_1: GAACCGGTTATTGGCGCATC	This study	N/A
susC_166_PUL485 R_1: CTTGGCATTTCCACTAGGAGC	This study	N/A
Software and Algorithms		
DADA2	Callahan et al., 2016	https://benjjneb.github.io/dada2/
Trimmomatic	Bolger et al., 2014	http://www.usadellab.org/cms/?page=trimmomatic
SPAdes	Bankevich et al., 2012	http://cab.spbu.ru/software/spades/
PanPhlAn	Scholz et al., 2016	https://bitbucket.org/CibioCM/panphlan/src/default/
CheckM	Parks et al., 2015	https://ecogenomics.github.io/CheckM/
PATRIC	(Wattam et al., 2014)	https://www.patricbrc.org/
MUSCLE	Edgar, 2004	https://www.drive5.com/muscle/
RAXML	Stamatakis, 2014	https://cme.h-its.org/exelixis/web/software/raxml/index.html
Silva SINA aligner	(Pruesse et al., 2012)	https://www.arb-silva.de/aligner/
Other		
Anaerobic Tissue Transport Medium Surgery Pack – ATTMSP	Anaerobe Systems	AS-915
Brucella Blood Agar - BRU	Anaerobe Systems	AS-141
Laked Brucella Blood Agar w/ Kanamycin and Vancomycin – LKV	Anaerobe Systems	AS-112

Peptone Yeast Broth with Glucose - PYG	Anaerobe Systems	AS-822
Yeast Casitone Fatty Acids Broth with Carbohydrates – YCFAC Broth	Anaerobe Systems	AS-680
Amylopectin from maize	Sigma	10120-250G
Arabinan (Sugar Beet)	Megazyme	P-ARAB
Arabinogalactan (Larch Wood)	Megazyme	P-ARGAL
Arabinoxylan (Wheat Flour; Low Viscosity ~ 10 cSt)	Megazyme	P-WAXYL
Beta-glucan (Barley; High Viscosity)	Megazyme	P-BGBH
Chondroitin sulfate sodium salt (from shark cartilage)	Sigma	C4384-5G
Glucomannan (Konjac; Low Viscosity)	Megazyme	P-GLCML
Heparin sodium salt (from porcine intestinal mucosa)	Sigma	H3393-100KU
Inulin (chicory)	Megazyme	P-INUL
Levan (from Erwinia herbicola)	Sigma	L8647-1G
Pectin (from apple)	Sigma	93854-100G
Polygalacturonic Acid (PGA)	Megazyme	P-PGACT
Pullulan (from Aureobasidium pullulans)	Sigma	P4516-1G
Rhamnogalacturonan (Soy Bean)	Megazyme	P-RHAGN
Starch	VWR	1.01252.0100
Xylan (Beechwood; purified)	Megazyme	P-XYLNBE-10G
Xylan (from Corn Core, TCI America™)	Fisher	X007825G
Xyloglucan (Tamarind)	Megazyme	P-XYGLN
Mucin from porcine stomach, Type III, bound sialic acid 0.5-1.5 %, partially purified powder (100g)	Sigma	M1778-100G

LEAD CONTACT AND MATERIALS AVAILABILITY

Further information and requests for resources and reagents should be directed to and will be fulfilled by the Lead Contact, Dan R. Littman (Dan.Littman@med.nyu.edu).

EXPERIMENTAL MODEL AND SUBJECT DETAILS

Subjects enrolled in this study included new onset rheumatoid arthritis patients and healthy controls, as described in STAR methods. Ethical approval was granted by the Institutional Review Board of New York University School of Medicine.

METHOD DETAILS

Sample collection

Stool was collected into anaerobic collection tubes (Anaerobe Systems) from healthy controls and new onset rheumatoid arthritis patients, deidentified, and frozen at -80°C. New onset rheumatoid arthritis patients were untreated, presenting with symptoms, seropositive for anti-citrullinated protein antibodies, and not on antibiotics in the previous six months.

qPCR Screen for *Prevotella*

Genomic fecal DNA was extracted from fecal samples using the MoBio PowerSoil DNA extraction kit. Fecal DNA was screened for the presence of *P. copri* using qPCR primers designed for six conserved regions of the *P. copri* genome (Scher et al., 2013), as well as two conserved regions of the *Prevotella* 16S gene (see **Key Resources Table**). Universal primers specific for the 16S rRNA gene were used to quantify total bacterial load in each sample. qPCR was performed with the SYBR green master mix on the Roche Lightcycler 480, with the following cycle conditions: 90°C for 5 minutes, then 40 cycles of 95°C for 10 seconds, 60°C for 30 seconds, and 72°C for 30 seconds. Genomic DNA from *P. copri* DSM 18205 was used to generate a standard curve. The standard curve was used to determine the absolute quantity of *Prevotella* and 16S in a sample. Fold change was calculated by dividing absolute quantity of *Prevotella* by absolute quantity of 16S. Feces with a fold change > 0.1 for any *Prevotella*-specific primer set were considered positive and used to isolate *P. copri*, with the exception of sample AU, which had a lower fold change.

Isolation of *Prevotella*

P. copri was isolated from *Prevotella*-positive feces by anaerobic quadrant streaking of PBS-diluted and undiluted stool on BRU and LKV plates (Anaerobe Systems), then incubated

anaerobically at 37°C for 24-48 h. Isolate colonies were picked, enumerated, streaked on a fresh plate, and the inoculation loop dipped into PCR grade water. The loop-dipped PCR water was used with the qPCR primer sets (above) to screen for *P. copri*. *Prevotella*-positive-screened isolates were confirmed by Sanger sequencing of the V3-V4 hypervariable region of the 16S rRNA gene, and finding >97% identity to known *Prevotella* species. Glycerol stocks of *Prevotella* isolates were frozen at -80°C.

16S rRNA gene sequencing

Genomic fecal DNA was extracted from fecal samples using the MoBio PowerSoil DNA extraction kit. The V3-V4 region of the 16S rRNA gene was amplified, purified, and sequenced on Illumina MiSeq (Caporaso et al., 2012). Sequences generated from amplicon sequencing were trimmed (maxN=0, truncQ=2) and denoised using DADA2 (Callahan et al., 2016). Sequences were annotated using the Silva nr v128 database.

Whole genome sequencing

Isolates were grown on BRU plates for 24-48 hours, until a healthy lawn was present. The lawn was suspended in PYG media (Anaerobe Systems), pelleted, and DNA was isolated with the MoBio PowerSoil DNA isolation kit. Libraries were prepared for sequencing with the TruSeq PCR-free library preparation kit and sheared to 500bp fragment length. Prepared libraries were sequenced on Illumina HiSeq2500 2x100 bp. Paired-end sequences were trimmed using Trimmomatic (Bolger et al., 2014), then whole genomes were assembled using SPAdes (Bankevich et al., 2012). Trimmed reads were also mapped to the panphlan_pcopri_16 database (<https://bitbucket.org/CibioCM/panphlan/wiki/Pangenome%20databases>) using PanPhlAn

(default parameters, **Figure 2A**) (Scholz et al., 2016). Quality and completeness of the assemblies were evaluated using CheckM (Parks et al., 2015). Genomes were annotated in PATRIC and are available at <https://www.patricbrc.org/> and under BioProject **PRJNA559898**. Presence and absence of predicted proteins can be visualized at https://flatironinstitute.github.io/genome_cluster_array_visualization/.

Phylogenetic Trees

Representative genomes for the *Prevotella* and *Bacteroides* genera were downloaded from NCBI. For each genome, a set of 30 ribosomal protein genes (LSU ribosomal proteins L1, L2, L3, L4, L5, L6, L10, L13, L14, L15, L18, L22, L23, L24, L29, and SSU ribosomal proteins S2, S3, S4, S5, S7, S8, S9, S10, S11, S12, S13, S14, S15, S17, S19) were identified and aligned using MUSCLE (Edgar, 2004). A maximum-likelihood tree was calculated using RAxML (-m GTRCAT; (Stamatakis, 2014)) from a concatenated alignment of the 30 ribosomal protein genes. A 16S rRNA gene tree was also estimated from the alignment of full length 16S rRNA gene sequences using Silva's SINA aligner (<https://www.arb-silva.de/aligner/>) and RAxML (-m GTRCAT).

Enzyme activity and acid production assays

Isolates and type strains were grown anaerobically for 18-24, harvested, and plated as per manufacturer's instructions on API® ZYM, API® Rapid ID 32 A Microbial Identification Kit, and API® 20 A Microbial Identification Kit test strips (BioMerieux). Test strips were incubated at 37 degrees C. Enzyme reactions were developed after 4.5 h, and acid production was assessed after 24h.

SusC analysis and polysaccharide utilization prediction

All annotated *susC* genes in the isolate genomes were clustered at >90% identity. This analysis revealed 168 *susC* gene clusters throughout all the genomes (**Figure S2**). For each *susC* gene cluster, ten flanking genes on either side were pulled from the genome and annotated (**Table S3**). Flanking genes were examined and compared to known gene activities in *Bacteroides* species to predict PUL substrates. *SusC* clusters lacking a companion *susD* gene, as well as those comprised of genes unrelated to polysaccharide catabolism, were excluded from the pPUL repertoire analysis.

In vitro polysaccharide growth assays

B. thetaiotaomicron, *B. ovatus* (both from Michael Fischbach), *Prevotella copri* DSM 18205 (DSMZ), and *P. copri* isolates were grown anaerobically on BRU plates (Anaerobe Systems) for 48h at 37 degrees C. Strains were passaged to YCFAC liquid media (Anaerobe Systems). 48h later, cultures were passaged to pre-reduced YCFA +/- 0.5% (w/v) individual polysaccharides or mucin proteoglycan (**Key Resources Table**) at a dilution of 1:100 in a total volume of 1ml (Browne et al., 2016). Cultures were incubated anaerobically in sterile polypropylene tubes at 37 degrees C. Importantly, *P. copri* grew in Wheaton™ CryoELITE™ Cryogenic Storage Vials, FALCON 14 ml Polypropylene Round-Bottom Tubes, and Titertube® Micro Test Tubes (see **Key Resources Table**), but not in Axygen™ Storage Microplates from Fisher Scientific (Catalog no. 14-222-224). Every 24h, 100 µl of each culture was transferred to a flat-bottom 96-well plate, and OD600 was measured with the Envision plate reader. OD600 for each culture condition was subtracted from the average blank OD600 for the corresponding polysaccharide. Cultures were grown in duplicate or triplicate. For each condition, the maximum average OD600

from the 24h, 48h, or 72h time point was determined and duplicate or triplicate values were used in **Figure 3A**, **3B**, and **Figure S3A**.

susC gene expression analysis

P. copri DSM 18205 was grown anaerobically on BRU plates and passaged to YCFAC liquid media (Anaerobe Systems), as for in vitro polysaccharide growth assays (above). 48h later, cultures were passaged to pre-reduced YCFA +/- 0.5% (w/v) individual polysaccharides (**Key Resources Table**) or glucose as a negative control at a dilution of 1:100 in a total volume of 50 ml. The OD of cultures was monitored and when it indicated mid-log phase, 10 ml of each culture was harvested, pelleted, stabilized in RNA Protect Bacteria Reagent (Qiagen), and frozen at -80 degrees C. RNA was extracted with the RNeasy kit (Qiagen), and cDNA was generated with the Superscript IV kit. qPCR was performed with the SYBR green master mix on the Roche Lightcycler 480, with the following cycle conditions: 90°C for 5 minutes, then 40 cycles of 95°C for 10 seconds, 60°C for 30 seconds, and 72°C for 30 seconds. mRNA levels of *susC* genes were measured with primers specific for each *susC* gene (see **Key Resources Table**), and normalized to the 16S rRNA gene. Genomic DNA from *P. copri* DSM 18205 was used to generate a standard curve.

PUL prediction in *P. copri* MAGs

SusC and *susD* homologs in *P. copri* MAGs (Tett et al., 2019) were identified by blastp (Identity>75% , Bitscore > 300) using *susC* and *susD* predicted proteins from the *P. copri* isolate genomes. Only MAG contigs containing both *susC* and *susD* homologs within 10 gene products were considered further. For each *susC* centroid, the PUL operon was extended to include the 10

gene products upstream and downstream of the first and last occurrence of a *susC* or *SusD* homolog. Each PUL was checked for CAZy annotations (Lombard et al., 2014) predicted using HMMSEARCH (version 3.1b2) (Eddy, 2011) against dbCAN HMMs v6 (Yin et al., 2012) with an E-value < 1e-18 and coverage > 0.35. PULs were predicted by searching for the presence of CAZymes known to act on known polysaccharides. Predictions are presented as the proportion of *P. copri* MAGs containing at least one pPUL out of all *P. copri* MAGs within each *P. copri* complex clade.

QUANTIFICATION AND STATISTICAL ANALYSIS

Prediction analysis

For each isolate used in the *in vitro* polysaccharide growth assay, a one-way ANOVA was performed on the maximum OD for each polysaccharide. A post hoc Dunnett's test was performed with the H₂O condition (no carbon source) as the negative control. Isolate-polysaccharide conditions with OD values that were statistically significant compared to the H₂O control were considered positive for growth. Isolate-polysaccharide conditions that were not significant were considered negative for growth.

Statistical analysis of prediction

Isolate-polysaccharide conditions were separated according to whether growth on a candidate polysaccharide was predicted by *susC* gene expression or genome sequence. Isolate pPULs sharing synteny with *P. copri* DSM 18205 pPULs confirmed by *susC* gene expression were grouped together. If a pPUL was present in the genome of an isolate, the isolate-polysaccharide condition was considered "growth predicted." If a pPUL was absent in the genome of an isolate,

the isolate-polysaccharide condition was considered “growth not predicted.” For xylan, arabinan, and xyloglucan, for which there were both gene expression or genome sequence predictions, gene expression predictions were used in instances of discrepancies between the predictions. Isolate-polysaccharide conditions that were statistically significant as determined by a one-way ANOVA with a post hoc Dunnet’s test using the H₂O condition (no carbon source) as the negative control were given a score of 1 and isolate-polysaccharide conditions that were not significant were given a score of 0. Welch’s t test was performed on the data.

DATA AVAILABILITY

All 16S sequencing data and whole genome sequencing data are available under BioProject **PRJNA559898**.

Supplemental Information:

In figure legend document now

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