Glycan complexity dictates microbial resource allocation in the large intestine

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The structure of the human gut microbiota is controlled primarily through the degradation of complex dietary carbohydrates, but the extent to which carbohydrate breakdown products are shared between members of the microbiota is unclear. We show here, using xylan as a model, that sharing the breakdown products of complex carbohydrates by key members of the microbiota, such as Bacteroides ovatus, is dependent on the complexity of the target glycan. Characterization of the extensive xylan degrading apparatus expressed by B. ovatus reveals that the breakdown of the polysaccharide by the human gut microbiota is significantly more complex than previous models suggested, which were based on the deconstruction of xylans containing limited monosaccharide side chains. Our report presents a highly complex and dynamic xylan degrading apparatus that is fine-tuned to recognize the different forms of the polysaccharide presented to the human gut microbiota.

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Genomic and metagenomic data show that the catabolism of complex dietary carbohydrates (CDCs), also known as dietary fibre, play a central role in defining the structure of the human gut microbiota (HGM)\textsuperscript{1–2}. The Bacteroidetes of the HGM, which include the genus Bacteroides, are viewed as glycandegrading generalists capable of utilizing an extensive array of both dietary (plant and animal) and host complex carbohydrates\textsuperscript{3–4}. The other main phyla of the gut, Firmicutes and Actinobacteria (particularly Bifidobacterium spp.), also have the capacity to degrade glycans, but appear more specialized with a preference for plant storage polysaccharides (starch and fructan) and oligosaccharides, such as those derived from arabinoxylans (AXs)\textsuperscript{5–6}. Current strategies deployed to manipulate the composition of the HGM to maximize human health are centred on the use of complex carbohydrates (prebiotics)\textsuperscript{7} to encourage the expansion of beneficial (probiotic) microorganisms\textsuperscript{8}. The effects of these strategies are ultimately dependent on the glycan food web in the HGM\textsuperscript{9,10}.

In contrast, the number and complexity of the side chains on cereal GAXs, exemplified by corn bran xylan (CX) (Fig. 1) explains why these glycans are recalcitrant to breakdown by known xylan-degrading enzyme systems\textsuperscript{11}.

Bacteroides ovatus, a prominent member of the HGM, was previously shown to utilize an extensive range of plant polysaccharides, including xylans\textsuperscript{4}. Here we explored the mechanisms by which this bacterium degrades the different forms of the polysaccharide presented to the HGM. Our data show that B. ovatus has evolved a highly dynamic xylan degrading system that can respond to the different forms of the xylose polymer. The bacterium has recruited several enzymes from glycoside hydrolase (GH) families\textsuperscript{12} not previously associated with xylan degradation. We also show that, although polysaccharide breakdown products (PBPs) are released during degradation of both simple and complex xylans by B. ovatus, the ability of other members of the HGM to use these PBPs is dictated by the complexity of the target xylan.

**Results**

*B. ovatus* contains two discrete loci activated by xylans. The genomes of Bacteroidetes contain numerous polysaccharide utilization loci (PULs) that are optimized to orchestrate the degradation of specific polysaccharides\textsuperscript{4,17–19}. The glycans

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**Figure 1** | **Schematic of the structures of the main classes of xylan.** The monosaccharides and linkages in the main classes of xylan are shown and are represented in their Consortium for Functional Glycomics format\textsuperscript{69}. The xylans used in this study were from birchwood (birch glucuronoxylan; BGX), wheat flour (wheat arabinoxylan; WAX) and corn bran (corn glucuronoorabinoxylan; CX).
targeted by the PULs activate transcription of their cognate loci. *Bacteroides ovatus* strain ATCC 8483 contains two PULs, spanning locus tags bacova_03417-50 (large xylan PUL or PUL-XylL) and bacova_04385-94 (small xylan PUL or PUL-XylS), that are activated when the organism is grown on wheat arabinoxylan (WAX) (Fig. 2a,b). In contrast, linear xylooligosaccharides (preferentially xylotetraose) induce transcription of PUL-XylS alone. Here we show that growth on CX leads to significant activation of PUL-XylL and only very low activation of PUL-XylS, while glucuronoxylan (from birch; BGX) activates only PUL-XylS (Fig. 2c). These data suggest that the degradative system encoded by PUL-XylS targets simple linear or glucurono-substituted xylans, while the products of PUL-XylL drive the metabolism of more complex decorated forms of the hemicellulose. Bacteroidetes spp. are known to express enzymes capable of breaking down simple forms of xylan,20,21; however, the mechanisms by which these microorganisms utilize complex cereal GAXs (for example, CX) are not known. Both PULs encode glucanases belonging to GH families16 that are known to contribute to the degradation of simple xylans (GH3, GH10, GH30, GH43, GH67 and GH115)22 (Fig. 2a,b). Significantly, PUL-XylL also directs the synthesis of enzymes belonging to families (GH31, GH95, GH97 and GH98)4 previously implicated in the deconstruction of the hemicellulose (Fig. 2a).

**Surface acting xylan-degrading enzymes.** Bioinformatic analysis, in conjunction with cell localization studies and whole-cell assays, was used to identify the enzymes encoded by PUL-XylS and PUL-XylL that are displayed on the bacterial surface. The presence of a predicted Type II lipoprotein signal sequence or uncleaved transmembrane (TM) anchor in seven GH family proteins indicated that they were membrane attached. The cell localization of these seven proteins was evaluated experimentally by (i) fluorescence microscopy of whole cells treated with labelled antibodies directed against the appropriate proteins and (ii) western blotting after removal of surface proteins with Proteinase K. The data indicated that BACOVA_03419 (GH3), BACOVA_03421 (GH43), BACOVA_03431 (inactive GH10, see later results), BACOVA_03432 (GH30), BACOVA_03433 (GH98) and BACOVA_04390 (GH10) were presented on the surface of *B. ovatus* (Fig. 3). The only lipoprotein facing the periplasm was the GH43 enzyme BACOVA_04325. These conclusions were further supported by whole-cell assays of the *B. ovatus* mutant lacking PUL-XylS (ΔPUL-XylS) incubated with BGX. The product profile of ΔPUL-XylS was extremely similar to that of recombinant BACOVA_03432 GH30 against the polysaccharide (Supplementary Fig. 1a). Thus, BGX degradation is mediated predominantly by BACOVA_03432 in the ΔPUL-XylS mutant, confirming the extracellular location of the GH30 enzyme. Whole-cell assays also showed that the PUL-XylL encoded GH43 arabinofuranosidase, BACOVA_03421 (see results below), was extracellular, as arabinose was released from WAX by the ΔPUL-XylS mutant (Supplementary Fig. 1b). It is surprising that BACOVA_03419, a GH3 β-xylanosidase, is located on the surface as its obvious role is to contribute to the degradation of the xylooligosaccharides generated in the periplasm (Supplementary Fig. 2a). BACOVA_03419, however, displays only low activity against linear β1,4-xylooligosaccharides, but, unlike the cytoplasmic xylanosidase, BACOVA_04386 (see results below), will

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**Figure 2 | Schematic of the *B. ovatus* xylan PULs and differential expression during growth on different xylans.** (a,b) schematic of PUL-XylL and PUL-XylS, respectively. Each gene is drawn to scale as a rectangle with its orientation indicated by the arrow head. The numbers below each gene is its locus tag (bacova_XXXXX). Genes encoding known or predicted functionalities are colour-coded and, where appropriate, are also annotated according to their CAZy family number: glycoside hydrolase (GH; green), carbohydrate esterase (CE; purple), carbohydrate-binding module (CBM). Surface located proteins are marked with an asterisk. SGBP = surface glycan binding proteins. **(c)** Cells were grown on minimal media with polysaccharide as the sole carbon source, and levels of different susC transcripts (locus tags shown; used as a proxy for expression of the whole PUL) from each PUL were quantified by qRT-PCR. The y-axis shows the Log fold-change relative to a minimal media-glucose reference; x-axis labels indicate the xylans used. WAX = wheat arabinoxylan, CX = corn xylan, BGX = birchwood glucuronoxylan. CX is a highly complex xylan compared to WAX and BGX. All data were analysed by one-way ANOVA followed by Tukey’s multiple comparison test (*** = P ≤ 0.001).
cleave linkages in which the leaving group xylose is decorated (Supplementary Table 1 and Supplementary Fig. 2b,c). While the roles of BACOVA_03419 and BACOVA_03421 in xylan degradation are unclear, we speculate that the enzymes contribute to the surface ‘trimming’ of complex xylans.

Three of the surface enzymes (BACOVA_04390 GH10, BACOVA_03432 GH30 and BACOVA_03433 GH98) exhibited endo activity and therefore are likely involved in cleaving the xylan backbone before importing into the periplasm. Studies on recombinant BACOVA_04390 GH10 showed that the endo-xylanase displayed a catalytic efficiency against xylan ~30-fold lower than typical GH10 enzymes\(^\text{23,24}\) (Supplementary Table 2). The limit products against sparsely decorated xylan contains oligosaccharides with a degree of polymerization (d.p.) \(>4\) (Supplementary Fig. 3a). Typical GH10 xylosidases\(^\text{33,24}\), exemplified by the periplasmic enzyme BACOVA_04387 generate xylose and xylobiose as the final products (Supplementary Fig. 3b). These data suggest that BACOVA_04390 contains an extended substrate binding cleft that requires a high degree of occupancy for catalysis to occur. Substrate depletion data using xylooligosaccharides confirmed that the enzyme had very poor activity against substrates with a d.p. \(\leq 5\) compared with typical GH10 enzymes (Supplementary Table 2). BACOVA_04390 has an unusual primary sequence in which the canonical GH10 (\(\beta/\alpha\)_\(\text{c}\)-fold barrel is interrupted by two copies of a non-catalytic carbohydrate-binding module (CBM) that is distantly related to CBM family 4, suggesting that the two CBMs contribute to an extended substrate binding cleft\(^\text{21,25}\). Indeed a recent study has shown that the internal CBMs from a related \(B.\) intestinalis GH10 bind xylans and long xylooligosaccharides\(^\text{21}\). This view is consistent with the observation that endo-acting enzymes, which specifically target large polymers, contain additional subsite(s) likely compensating for relatively weak binding at subsites proximal to the active site\(^\text{26,27}\).

The GH30 enzyme BACOVA_03432 produced a range of large oligosaccharides from BGX (Supplementary Fig. 1a and Supplementary Table 2), but was not active on WAX or CX. This specificity is typical of GH30 glucuronoxylanases and is driven by recognition of the (Me)GlcA side chain at the -2 subsite of these enzymes\(^\text{28}\). While the lack of activity against CX is somewhat surprising as the enzyme is part of PUL-XyIL, it may be that less heavily decorated GAXs are the target substrates for this enzyme.

The surface GH98 enzyme BACOVA_03433 also generates very large oligosaccharides but exclusively from CX (Fig. 4a,b). As BACOVA_03433 is a member of GH98, a family that hitherto contained only endo-\(\beta\)-\(\text{d}\)-galactosidases\(^\text{29}\), it was formally possible that the enzyme targets rare \(\beta\)-\(\text{d}\)-Gal residues in the backbone of CX. Converting the reducing end aldehyde of the products generated by BACOVA_03433 to the aldito by acid hydrolysis revealed the presence of xylitol but not galactitol (Fig. 4c). These data show that BACOVA_03433 is an endo-xylosidase that hydrolyses xylosidic bonds in a context found only in very complex xylans. It is possible that the enzyme targets an unusual linkage in the xylan backbone; however, this is unlikely as the enzyme was not active against \(\beta\)-\(1\,\text{,}\,4\)-\(\text{b}\)-\(1\,\text{,}\,4\) and \(\beta\)-\(1\,\text{,}\,3\)-\(\text{b}\)-\(1\,\text{,}\,4\) linked xylans and CX is reported to contain a backbone comprising exclusively \(\beta\)-\(1\,\text{,}\,4\)-\(\text{Xylo}\) units\(^\text{13}\). To explore the specificity determinants of the GH98 enzyme further, CX was pre-treated with cocktails of side-chain cleaving hydrolases (arabinofuranosidases, xylosidases, glucuronidases and galactosidases) encoded by PUL-XyIL, and the debranched forms of the xylan were then incubated with the GH98 xylanase. Only GH consortia lacking BACOVA_03417 generated a variant of CX that was hydrolysed by the GH98 enzyme (Fig. 4d). As described below BACOVA_03417 is an \(\alpha\)-\(L\)-arabinofuranosidase that targets the O3 linked Ary/ on backbone Xyl residues decorated at both O2 and O3 with \(\alpha\)-\(L\)-Araf. It would appear, therefore, that the double Ary/ decoration is a component of the specificity determinant for BACOVA_03433. Wheat arabinoxylan, however, also contains this double arabinose decoration and is not a substrate for the GH98 xylanase (Fig. 4a). Thus, the specificity of BACOVA_03433 is more complex than simply Xyl units linked O2 and O3 to arabinose. Interestingly, the CX backbone contains low levels of doubly substituted residues with Xylp as the O3 linked side chain (see Fig. 1 and results below). This structure is also a substrate for BACOVA_03417 (Supplementary Fig. 4a), suggesting Xylp at O3 (and Araf at O2) may be the critical specificity determinant of the GH98 enzyme. These Xylp decorations are absent in WAX (Supplementary Fig. 4a–c), providing a possible explanation for the lack of activity of the GH98 enzyme against this substrate.

The large products generated by all three surface xylanases indicate that these enzymes recognize infrequent specificity determinants in the xylan, which, as discussed below, likely
contributes to the capacity of B. ovatus to efficiently utilize the oligosaccharides generated by these GHs. It is particularly intriguing that the specificities of the three endo-acting enzymes are adapted to recognize the variable decorations appended to the backbone of different xylans, likely reflecting the multiple forms of the polysaccharide presented to the human large intestine.

Surface xylan-binding proteins. PUL-XylS and PUL-XylL encode one (BACOVA_04392) and two (BACOVA_03427 and BACOVA_03429) SusD-like proteins, respectively. SusD-like proteins appear to be involved in binding and delivering oligosaccharides to their cognate SusC-like transporter for import into the periplasm. Isothermal titration calorimetry showed that BACOVA_04392 bound to the relatively simple xylans WAX and BGX with moderate affinity ($K_d \approx 0.3-0.4$ mM), but did not interact with highly decorated CX (Supplementary Table 3). The protein bound xylotriose, but not shorter xylooligosaccharides, consistent with the long products generated by the surface GH10 xylanase encoded by PUL-XylS, BACOVA_04390 (Supplementary Fig. 3a). The SusD-like protein, derived from PUL-XylL, BACOVA_03427, bound tightly to BGX and CX ($K_d \approx 7-12$ mM), but not WAX or linear xylooligosaccharides, although the precise target ligand for this protein is unclear (Supplementary Table 3). The other PUL-XylL SusD-like protein, BACOVA_03429, could not be expressed in a recombinant form. In addition to the SusD-like proteins, PULs often encode surface-located glycan binding proteins (SGBPs), which are functionally analogous to SusE and F from the prototypic starch utilization system. The capacity of three non-catalytic

Figure 4 | The activity of the GH98 enzyme BACOVA_03433. (a) Different xylans at 1% (w/v) were incubated with 1 $\mu$M of the GH98 enzyme in PBS for 16 h at 37°C. The reaction products were subjected to thin layer chromatography and compared with xylooligosaccharide standards with a d.p. ranging from 1 to 5 (X1-X5). (b) HPAEC-PAD trace showing the high molecular weight oligosaccharide products of CX digestion by the GH98 endo-xylanase, separated using a sodium acetate gradient of 0-500 mM. The reaction conditions used are the same as in panel a Control reactions of substrate and enzyme only are shown. (c) CX was incubated with 5 $\mu$M of the GH98 enzyme for 24 h in 50 mM sodium phosphate buffer, pH 7.0, at 37°C. The reaction products, and CX that was not incubated with the GH98 enzyme, were then treated with sodium borohydride, to convert the reducing end monosaccharide unit into its alditol, followed by acid hydrolysis to convert the oligosaccharides (or in the case of control CX alone) into their monosaccharide constituents. The hydrolysed products were subjected to HPAEC and compared to the migration of standard monosaccharides as well as the alditol of xylose (xylitol) and galactose (galactitol). (d) The GH98 enzyme was incubated with CX (lane CX vs GH98), or CX that had been pre-treated with all the side chain cleaving enzymes encoded by PUL-XylL (lane CX vs MIX), or CX that had been pre-treated with all the side chain cleaving enzymes encoded by PUL-XylL, except the GH43 enzyme BACOVA_03417 that removes O3 linked arabinose from double substituted xylose (lane CX vs MIX no double abfase).
Novel xylan debranching enzymes. As discussed above the two xylan PULs encode debranching enzymes located in GH families typically associated with xylan degradation (GH43, GH67 and GH115) that cleave α-L-arabinofuranosyl and α-D-glucuronolxylosyl linkages. In addition, PUL-XylL encodes three periplasmic proteins that belong to GH31, GH97 and GH95, families that have not previously been implicated in the deconstruction of the polysaccharide. Assays showed that the GH31 enzyme, BACOVA_03422, released xylose from CX but displayed no activity against a range of other xylans (Supplementary Fig. 6). As GH31 enzymes target α-D-glycosides, BACOVA_03422 is likely an α-xylanosidase, consistent with the presence of α-D-Xyl in CX13. BACOVA_03422 is ~3,000-fold more active against CX than a xyloglucan oligosaccharide (Supplementary Table 6), demonstrating that the enzyme is a xylan-specific α-xylanosidase. In contrast, a previously characterized GH31 α-xylanosidase, BACOVA_03421,40, exhibited minimal activity against CX, strongly preferring xyloglucan oligosaccharide (Supplementary Table 6). These data show that the xylan specificity displayed by the BACOVA_03422 enzyme is not a general property of GH31 enzymes. The GH97 enzyme, BACOVA_03423, was active against 4-nitrophenyl-α-D-galactopyranoside, consistent with the presence of α-D-Gal in some highly complex xylans13. The GH95 enzyme BACOVA_03438, although located in an “α-L-fucosidase specific family”42, released exclusively 1-Gal from CX, displaying only trace activity against oligosaccharides containing terminal α-L-Fuc residues (Fig. 5a-c and Supplementary Table 7). Conversely, a previously characterized GH95 α-1,2-L-fucosidase (Afα)42 exhibited minimal α-L-galactosidase activity against CX (Supplementary Table 7), showing that GH95 enzymes can be
either L-Gal or L-Fuc specific. Although these data provide an initial example of a glycosidase specific for L-galactose, it should be noted that the GH117 family contains enzymes that target the 3,6-anhydro form of the sugar.43

Crystal structure of the GH95 α-L-galactosidase. The crystal structure of the α-L-galactosidase, BACOVA_03438, displays a central (α/α)6-helical barrel domain flanked by β-sandwich domains at the N- and C-termini, respectively. In the structure of the protein-product complex, β-L-Gal (GH95s are inverting enzymes), which adopts a 1C4 geometry typical of L-mono-saccharides, is buried in the central cavity of the helical barrel domain, (Fig. 5d–g). Structural analyses in combination with mutational experiments revealed that the highly conserved Glu501 acts as the general acid catalyst in a single displacement mechanism.
(inverting) mechanism (Fig. 5f and Supplementary Table 7). No carboxylic acid residue was found at the appropriate position for a general base catalyst, similar to the Bifidobacterium bifidum GH95 \(\alpha\)-l-fucosidase, AfcA. It was proposed that in AfcA two asparagine residues activate a water molecule that is suitably located to perform a nucleophilic attack on the C1 atom of \(\alpha\)-fucose in the active site\(^{42}\), which is in sharp contrast with typical GHS where Asp or Glu act as the catalytic base\(^{44}\). The proposed catalytic asparagines in AfcA (Asn421 and Asn423) are structurally conserved in the GH95 \(\alpha\)-l-galactosidase (Asn372 and Asn374), suggesting a similar role in BACOVA_03438 (Fig. 5f,g). Almost all of the residues that interact with the pyranose ring of \(\alpha\)-Gal in BACOVA_03438 are conserved in the AfcA fucosidase, although, intriguingly, Thr370 in the \(\alpha\)-l-galactosidase is replaced with His419 in the fucosidase (Fig. 5f,g). In AfcA His419 contributes to the hydrophobic sheath in which the C6 methyl group of the Fuc is housed, while in BACOVA_03438 the \(\gamma\)O of Thr370 makes a direct polar contact with the O6 of \(\alpha\)-Gal, the only part of the monosaccharide that differs from \(\alpha\)-Fuc. The importance of the Thr370 in BACOVA_03438 is demonstrated by the complete loss of activity of the T370H mutant against CX (Supplementary Table 7). Thr370 in BACOVA_03438 is demonstrated by the complete loss of activity of the T370H mutant against CX (Supplementary Table 7). While the Thr/His polymorphism may be a specificity determinant for galactosidase and fucosidase activity within GH95, the paucity of enzymes characterized in this family prevents a definitive conclusion regarding its functional significance. Indeed, the observation that the T370H mutation does not increase fucosidase activity suggests that \(\alpha\)-Gal to \(\alpha\)-Fuc specificity is driven by more than just the \(-1\) site (Supplementary Table 7). Significantly, the leaving group subsite (+1 subsite) is not conserved in the two enzymes (inferred in the case of BACOVA_03438), and it is possible that substrate binding in this region contributes to enzyme specificity. A key role for the +1 subsite in the two enzymes is consistent with their lack of activity against 4-nitrophenyl-\(\alpha\)-l-fucopyranoside (AfcA) and 4-nitrophenyl-\(\alpha\)-l-galactopyranoside (BACOVA_03438).

**Hydrolysis of undecorated xylooligosaccharides.** Breakdown of debranched xylooligosaccharides, derived through the action of the exo-acting enzymes described above, is mediated by enzymes encoded by PUL-XylS. A typical GH10 endo-acting xylanase (BACOVA_04387) rapidly generates oligosaccharides in the periplasm with a d.p. of primarily 1 to 4 (Supplementary Fig. 3b and Supplementary Table 2). These small xylooligosaccharides are substrates for the cytoplasmic GH43 \(\beta\)-xylosidase, BACOVA_04386 (Supplementary Table 1 and Supplementary Table 2). The spatial separation of the \(\beta\)-xylosidase from the enzymes that generate its substrate may reflect a selection advantage as less energy would be required to transport a small number of xylooligosaccharides across the inner membrane (likely via the PUL-encoded MFS class transporter, BACOVA_04388, Fig. 2b), compared with a larger number of xylose units if the \(\beta\)-xylosidase was periplasmic. It is also possible that the different cellular locations of the xylosidase and xylanase maximizes the signal concentration for the PUL-XylS HTCS sensor-regulator (BACOVA_04394), which recognizes short xylooligosaccharides\(^9\).

The function of the two xylan PULs in *B. ovatus*. The role of PUL-XylS and PUL-XylL in xylan degradation is illustrated by the growth profiles of mutant strains. Deletion of PUL-XylS (\(\Delta\)PUL-XylS) did not influence the growth of *B. ovatus* on complex xylans (GAXs) from corn, rice and sorghum, but prevented growth on simpler xylans (BGX or WAX) (Fig. 6). By contrast, inactivation of PUL-XylL (APUL-XylL) prevented growth on the three complex GAXs but did not significantly affect the utilization of simpler BGX or WAX (Fig. 6). The somewhat higher final density observed for the wild-type bacterium on WAX vs APUL-XylL is likely due to the inability of this mutant to remove the arabinose decorations from this substrate. The growth profile of the mutants are consistent with the biochemical data presented above showing that PUL-XylL encodes the only endo-acting enzyme (GH98) able to cleave the backbone of CX (and likely other complex GAXs), a prerequisite for the utilization of these polysaccharides (see below). Similarly, the inability of the PUL-XylS mutant to utilize simple xylans such as GXs and AXs, where the xylan backbone is the major component, is consistent with the locus encoding the key endo-acting surface GH10 (BACOVA_04390) that hydrolyses the backbone of sparsely

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Figure 6 | Growth profile of wild type and mutants of *B. ovatus* on xylans. WT and PUL deletion mutants were cultured in minimal media containing different xylans at 0.5% (w/v) as the sole carbon source. Birchwood glucuronoxylan (BGX; (a)) and wheat arabinoxylan (WAX; (b)) are relatively simple, sparsely decorated structures, while corn (c), sorghum (d) and rice (e) xylans are more complex heavily decorated glucuronoarabinoxylans (GAXs). The high starting OD for BGX is due to the background turbidity of the polysaccharide. A total of 6 replicate cultures were monitored at 20 min intervals for each substrate and used to generate the average growth curves shown.
decorated xylans. It is somewhat surprising, however, that ΔPUL-XylS is able to efficiently utilize GAX even though the mutant is unable to fully depolymerize the xylan backbone. This likely reflects the extensive side chains present in CX and other complex GAXs, where the xylose backbone constitutes only a relatively minor component of the total monosaccharide units in these polysaccharides.

To summarize, using the expression, localization and biochemical data presented above Fig. 7 shows the proposed models for the acquisition and breakdown by B. ovatus of the different classes of xylan presented to the colonic microbiota.

The conservation of PUL-XylS and PUL-XylL in Bacteroidetes. To identify PULs similar to PUL-XylS and -XylL in other genomes, we examined not only the gene family composition but also gene synteny in 260 Bacteroidetes species (Supplementary Table 8). Comparative genomics revealed a high degree of conservation of PUL-XylL in B. ovatus and B. xylanisolvens strains. Indeed, PULs with a high level of synteny with PUL-XylL was observed in 22 of the 23 strains examined. In these PULs, 17 genes show strict conservation with those present in PUL-XylL, which are arranged into three syntenic blocks (Fig. 8a). These 22 PUL-XylL homologues can be subdivided into two groups that

Figure 7 | Models for the degradation of different forms of xylan by B. ovatus. In the upper part of each panel the monosaccharides and linkages in the main classes of xylan are shown and are represented in their Consortium for Functional Glycomics format66. The lower parts of each panel show the model for degradation of complex GAXs like CX (a) AXs like WAX (b) and GXS like BGX (c). The red asterisk next to BACOVA_03425 GH43 indicates that the single specific arabinofuranosidase acts after BACOVA_03417 (α,1 double-specific arabinofuranosidase) to remove the remaining α,1,2 linked L-arabinose. The black arrows indicate examples of the linkages cleaved by the enzymes, shown in green ovals and identified by their glycoside hydrolase family and locus tag. In panel a, the dotted arrows from the GH98 enzyme identify possibly linkages hydrolysed by the xylanase. Although not shown on the CX model, the PUL-XylS apparatus will be present, albeit at low levels (see Fig. 2c). A schematic of the crystal structures, colour-ramped from blue (N-terminus) to red (C-terminus), of two of the xylan degrading enzymes, determined in this study (BACOVA_03438 GH95; PDB accession code 4UFC) and in a recent report67 (BACOVA_03449 GH11S; PDB accession code 4C91) are shown. The red arrow heads indicate glycans transport between cellular locations, while the yellow arrow heads show the ligand (or potential ligand in case of the question mark) that binds and activates the hybrid two component system (HTCS) encoded by PUL-XylS (BACOVA_04394, light blue HTCS; activating ligand xylooligosaccharide68 or PUL-XylL (BACOVA_03437, dark blue HTCS; activating ligand likely an arabinoxylooligosaccharide). MFS is a member of the Major Facilitator Superfamily of transport proteins, and SGBP signifies the two surface xylan binding proteins encoded by the xylan PULs, BACOVA_03431 (inactive GH10, PUL-XylL) and BACOVA_04391 (PUL-XylS). A schematic of the crystal structure of BACOVA_03439 (PDB accession code 3ORJ), colour-ramped from blue (N-terminus) to red (C-terminus), is shown. Note only one of the two SusC/D pairs from PUL-XylL is shown for clarity.
Figure 8 | Conservation of the B. ovatus ATCC 8483 xylan PULs in other Bacteroides species. CAZymes (green, CBM modules were omitted for clarity), susC-like (yellow), susD-like (orange) and the HTCS sensor-regulator (blue) genes are colour-coded. Genes encoding proteins of unknown function or non-CAZymes (for example, surface glycan binding protein and MFS carbohydrate importer in PUL-XylS) are shown in grey. (a) PUL-Xyl. Vertical dotted lines indicate split gene models, while missing gene models are shown in red. Microsyntenic segments are depicted by grey connectors, and include adjacent conserved non-PUL members (shown with dotted lines). Eighteen genes are conserved across all strains, and 17 of these form three highly conserved blocks indicated on top. The major difference between the two groups of the GH98 for GH5_21 swap is indicated. Inset shows a TLC of the activity of the recombinant B. xylanisolvens XBA1 GH5_21 enzyme (BXY_29320; boxed in red) against different xylans (1%), WAX vs GH5, CX vs GH5, X5 vs X4 vs X3 vs X2 vs X1). PUL-XylL displays activity on CX (Fig. 8 inset), similar to the B. ovatus GH98 enzyme BACOVA_03433. This suggests that both forms of PUL-Xyl retain the capacity to hydrolyse the backbone of complex GAXs without the need for extensive debranching. The high synteny described above for PUL-XylL in the B. ovatus and B. xylanisolvens strains was not found in any of the other 260 Bacteroidetes genomes analysed. In particular, the distribution of the GH98-encoding gene is extremely narrow, being present in only five of the 260 genomes analysed.
With respect to PUL-XylS there is perfect conservation of the synteny of the genes encoding the GH67 α-glucuronidase, GH43 β-xylosidase and periplasmic GH10 xylanase in all 23 B. ovatus and B. xylanisolvens strains (Fig. 8b). In 11 of these strains, however, the genes encoding the surface GH10 xylanase, SusD-like, SusC-like and HTCS proteins were absent. The loss of these critical components in these strains is likely to render the PULs non-functional. No other candidate PUL with complete conservation of all PUL-XylS components could be found in the other Bacteroidetes that we examined. The closest candidate PULs were found in B. intestinalis, B. gallinarum, B. oleicaplens and Proteiniphilum acetatigenes. Interestingly, some of these PULs are adjacent to gene families that are found in PUL-XylL that encode GH5_21, GH10, GH31, GH43, GH95 or GH115 enzymes (the degree of synteny is uncertain due to limited scaffold assemblies), suggesting a possible common evolutionary origin for the B. ovatus XylL- and XylS- PULs. Overall, of the 23 B. ovatus and B. xylanisolvens strains, only five have both XylL- and XylS-PULs, illustrating the dynamic acquisition (and possibly loss and/or reshuffling) of PULs by Bacteroidetes.

Loss of xylan degradation products from the cell surface. Similar to other characterized Bacteroides polysaccharide degrading systems, the xylan degrading apparatus of B. ovatus appears to be optimized to maximize intracellular breakdown, which may indicate that the bacterium adopts a ‘selfish’ strategy when deconstructing the hemicellulose17–19. In contrast with this selfish hypothesis is the observation that on simple xylans, such as BGX and WAX, B. ovatus was able to support the growth of Bifidobacterium adolescentis strain ATCC 15703, which utilizes simple linear and arabinono-xylooligosaccharides but not xylans47 (Fig. 9a,b and Supplementary Fig. 7). The Bacteroides, however, was unable to promote the growth of the Bifidobacterium on highly complex xylans such as CX (Fig. 9c). It could be argued that this differential capacity of B. ovatus to support the growth of B. adolescentis reflects the extent to which oligosaccharides from complex (CX) and simple (BGX and WAX) xylans are released into the culture medium and are thus available to the Bifidobacterium. It should be emphasized, however, that as we show here, CX breakdown requires a suite of enzymes not associated with canonical xylan degrading systems. Indeed only a limited number of closely related species of Bacteroides appear to contain these additional enzymes (see above and Fig. 8a). Thus, the inability of B. ovatus to support growth of B. adolescentis on CX may simply reflect the fact that the Bifidobacterium lacks the necessary apparatus to utilize this complex xylan. To explore this possibility a BACOVA_03433 GH98 mutant was constructed in which the key catalytic residues of the GH98 enzyme (Glu361 and Asp467) had been replaced with alanine. The mutant (ΔGH98) lacked an active surface GH98 xylanase and was unable to cleave the backbone of CX, preventing growth on the complex GAX (Fig. 9e). The B. ovatus variant, however, retained the other components of the xylan degrading apparatus and was thus able to grow well on CX that had been pre-digested with recombinant GH98 (Fig. 9e). Co-culture of ΔGH98 with wild-type B. ovatus on intact CX supported the growth of both strains, demonstrating that PBPs are indeed released into the culture medium by wild-type B. ovatus during growth on CX (Fig. 9f). Thus, the inability of B. adolescentis to grow on CX in the presence of B. ovatus does not reflect the release of an inadequate supply of xylan-derived PBPs, but indicates that the Bifidobacterium lacks the apparatus required to utilize these highly complex oligosaccharides. This is consistent with the finding that B. adolescentis is unable to grow on CX that had been pre-digested with either the GH98 alone or all the surface located enzymes encoded by PUL-XylL (Supplementary Fig. 7).

Figure 9 | The ability of other members of the microbiota to use PBPs released by B. ovatus during growth on xylan is determined by the complexity of the polysaccharide. Bacteroides ovatus (Bo) and Bifidobacterium adolescentis were co-cultured on BGX (a) WAX (b) and CX (c) and the number of CFUs of each determined at different points on the growth curve. (d) Shows the CFU ml⁻¹ of B. ovatus alone at different phases of growth. Note, when the Bifidobacterium alone was grown on digested xylans (see Supplementary Fig. 8), the CFU ml⁻¹ at late exponential phase (OD₆₀₀ ~1.2) was ~8.0 × 10⁸, indicating that ~25% of the total xylan is used by B. adolescentis. (e) Growth of B. ovatus wild type and the ΔGH98 mutant (BACOVA_03433) on CX (0.5% w/v). Pre-digested indicates the CX was digested to completion with the GH98 xylanase prior to addition to the media. (f) Tagged strains of wild-type B. ovatus and the ΔGH98 mutant were co-cultured on CX (see panel e for growth curve). Samples were taken at different time points and qPCR with primers unique to each strain was used to determine the ratio of each in the culture. Each data point is the average and s.d. of triplicate growths.
Currently the degree to which glycan degradation is mediated by synergistic interactions between different members of the gut microbiota is unclear. It has been proposed that there are keystone organisms belonging to the Firmicute phylum that make recalcitrant glycans, such as resistant starch, available to other members of the ecosystem.\(^4^\) Similarly, a recent study showed that the products released during degradation of highly accessible glycans (branched starch and fructans) by Bacteroides spp. were shared with recipients that alone are unable to utilize these polysaccharides.\(^5^\) In contrast, B. thetaiotaomicron utilizes complex yeast mannans in a selfish manner as breakdown occurs almost entirely intracellularly, with no evidence of sharing of PBPs with other Bacteroides spp.\(^6^\) This study extends our understanding of resource allocation in the microbiota by showing that selfish glycan utilization is not dependent solely on an ability to retain PBPs within the cell, but can be determined by the complexity of the enzyme systems required to access the extracellular oligosaccharides generated.

Discussion
This report provides a model for the microbial utilization of xylan, a key polysaccharide present in the human diet. The data demonstrate that current understanding of xylan degradation, based primarily on the breakdown of relatively simple xyains by soil microbial ecosystems, is inadequate when considering the complex GAXs found in the human diet.\(^2^\) It is evident that B. ovatus has acquired an extensive range of enzymes, several from GH families that had not previously been implicated in xylan breakdown, to assemble an efficient catalytic apparatus capable of deconstructing highly complex forms of the polysaccharide. It is significant that the genetic basis for xylan degradation is organized into two distinct loci regulated by different signals. This provides B. ovatus with a flexible strategy to degrade the hemicellulose. When presented with a simple linear xylan or GX a minimum enzyme cocktail is assembled (that is, PUL-XylS), while PUL-XylL is activated only in response to complex GAXs (Fig. 2c). It is likely, therefore, that the distribution of these two PULs within human microbiomes provides insight into the nature of the diet consumed by the respective host.

Using xyains that contain variable decorations but a conserved core backbone, we have shown that it is the complexity of the target polysaccharide that determines whether other members of the HGM can utilize the PBPs released by B. ovatus during growth on different xyains. Thus, the relatively simple oligosaccharides that are released during growth on WAX and BGX can be used by B. adolescentis, while the more complex molecules released from CX cannot be accessed by the Bifidobacterium. Indeed, our analysis of Bacteroidetes genomes suggests that very few organisms, outwith B. ovatus and B. xylanisolvens, have a sufficiently complex xylan degrading system to utilize the PBPs released from CX by these two Bacteroides species. It is possible that these complex glycans are only used in a nutritional crisis when competition for resources is intense within the microbiota. Recently, it has been shown that Bacteroides thetaiotaomicron displays a degradative hierarchy in which some simple glycans are prioritized above more complex polysaccharides.\(^3^\) Understanding the mechanisms by which different glyamins are degraded by the microbiota, specifically whether they are public-good products or are used exclusively by the degrading organism, is critical to developing nutraceutical strategies to manipulate the structure of the microbiota to maximize human health. For example, complex GAXs will favour the growth of B. ovatus, which produces propionate, a molecule that reduces lipogenesis in the host.\(^2^\) In contrast, the degradation of simple xyains by B. ovatus will also promote the growth of Bifidobacteria, which can lead to the production of butyrate, a molecule that is known to maintain the health of the intestinal epithelium.\(^2^\) Thus, it may be possible to develop bespoke xylan-based diets that are designed to have different beneficial effects on human health.

Methods
Isolation of corn xylan. Oven-dried corn fibre sample was kindly provided by ADM Research (North America), which was further oven-dried and ground to a 20 mesh particle size using a Wiley mill.\(^\text{5}\) The ground corn fibre sample was extracted with hexane to remove oil and treated with thermostable z-amylase (a gift from Novo Nordisk Bioindustrials, Danbury, CT) at 90 °C to hydrolyse the starch present in the fibre.\(^\text{6,5}\) CX was isolated from de-oiled and de-starched corn following the alkaline hydrogen peroxide technology of Yadav et al.\(^\text{5}\) with the following modifications. De-oiled and de-starched corn fibre (50 g) was sufficiently complex xylan degrading system to utilize the PBPs with other Bacteroides spp.\(^2^\) and B. adolescentis growth on different xylans. Thus, the relatively simple oligosaccharides that are degraded by the microbiota, specifically whether they are prioritized above more complex polysaccharides.\(^\text{5}\)

Cloning, expression and purification of recombinant proteins. The open-reading frames encoding mature forms of the proteins used in the study were amplified from Bacteroides ovatus ATCC 8483 genomic DNA by PCR using appropriate primers (Supplementary Table 9), which introduce BamHI/HindIII and Ndel/Xhol sites to the flanks of the target genes. The amplified DNA was cloned into Escherichia coli expression vectors (see Supplementary Table 9) such that the encoded recombinant proteins contain either an N- or C-terminal His-tag. Site-directed mutants were made using the Quikchange kit (Agilent), using primers shown in Supplementary Table 10. The recombinant proteins were produced in E. coli BL21 (DE3) or Tuner (Novagen) cells cultured in LB broth containing ampicillin (100 µg ml\(^{-1}\)) or kanamycin (50 µg ml\(^{-1}\); Sigma-Aldrich) as appropriate. The E. coli cells were grown to mid-exponential phase (OD\(_{600}\) = 0.6), at which point isopropyl \(\beta\)-D-thiogalactopyranoside was added to a final concentration of 1 mM (BL21) or 0.2 mM (Tuner), and the cultures were incubated for a further 16 h at 16 °C. The cells were harvested by centrifugation, sonicated and His-tagged recombinant protein was purified from cell-free extracts by immobilized metal ion affinity chromatography using Talon resin (Clontech) and standard methodology as described previously.\(^\text{5}\) For crystallographic studies BACOVA_03438 was further purified by size exclusion chromatography using a Superdex 75 column (GE Healthcare Life Sciences). All proteins were purified to electrophoretic homogeneity as judged by SDS-PAGE and their concentrations determined from their calculated molar extinction coefficient at 280 nm using a NanoDrop 2000c (Thermo Scientific).

Growth of anaerobic bacteria. Bacteroides ovatus ATCC 8483 was routinely grown in tryptone-yeast extract-glucose (TYG) medium or on brain-heart infusion (BHI; Beckton Dickinson) agar plus porcine hematin (1.2 µg ml\(^{-1}\); Sigma-Aldrich)\(^\text{5,7}\). Bifidobacterium adolescentis ATCC 15703 was grown on clostridial growth medium (CNM; Sigma-Aldrich)\(^\text{7}\) plus hematin. Antibiotics were added as appropriate: erythromycin (25 µg ml\(^{-1}\)) and gentamicin (200 µg ml\(^{-1}\)). Bacteroides minimal medium (MM) was formulated as described previously\(^\text{5}\) as Bifidobacterium MM\(^\text{2}\). Cells were grown in an anaerobic cabinet (Whitley A35 Workstation; Don Whitley, UK) or incubated in an anaerobic glove box (Birnbaum, USA) for 1 week. Growth of B. ovatus wild type and PU-knockout mutants was measured on various xylan preparations by monitoring the increase in absorbance at 420 nm when grown in 5 ml of minimal medium containing 0.5% (w/v) carbon source.

Quantitative RT-PCR (qPCR). Comparison of the levels of susC transcript expression from each of the xylan PULs was performed by qPCR. Previous studies have shown that the susC genes are a good proxy for expression of their cognate PUL.\(^\text{4}\) B. ovatus was cultured in 5 ml of minimal media containing 0.5% (w/v) carbon source and 50 µg ml\(^{-1}\) of ampicillin for 2 days before harvesting by centrifugation at 6,000 × g. Cells were harvested by centrifugation at 10,000 × g for 30 min and discarded. Two volumes of ethanol (2.0) were gradually added to the supernatant (1.0), obtained after Hemi. A removal, with stirring to precipitate the major arabinoolxylan fraction (CX). The CX was allowed to settle out as a white flocculent precipitate at the bottom of the beaker for 10–15 min. The clear alcohol/water mixture above the precipitate was removed by decantation. The white flocculent precipitate was collected by filtration and dried in a vacuum oven at 50 °C overnight.

The other xyains used in this study were low viscosity wheat arabinoxylan (WAX; Megazyme International) and birchwood glucuronoxylan (BGX; Sigma Aldrich).
source, as described above. Triplicate bacterial cultures were harvested at mid-log phase (OD600 ~ 0.8) and placed in RNAProtect (Qiagen), then stored at ~ 80 °C overnight, before purification with RNasew kit (Qiagen). RNA purity was assessed spectrophotometrically, and 1 μg of RNA was used immediately for reverse transcription (QuantiTect Reverse Transcription kit, Qiagen). Quantitative RT-PCR was performed in a 96-well plate on a LightCycler 480 System (Roche) with FastStart Essential DNA Green Master (Roche) using the primers shown in Supplementary Table 1. Reactions were carried out in 5 μL SYBR Green mix, 20 ng of cDNA, and 1 μM (sense/genes) or 0.125 μM (16 S RNA) primer mix. Reaction conditions were 95°C 600 s, followed by 45 cycles of 95°C for 10 s, 55°C for 10 s, 72°C for 10 s. Cq values were calculated using LightCycler 480 Software. Data were normalized to 16 S RNA transcript levels, and in change level calculated as fold-change compared with minimal media, glucose cultures.

Cell localization studies. Potential surface-located enzymes (containing either a predicted lipoprotein signal sequence or uncharged TM anchor) encoded by the B. ovatus xylan PULs were initially identified using the Lipop1.0 (http://www.cbs.dtu.dk/services/Lipop1/), and SigP 4.1 (http://www.cbs.dtu.dk/services/SigP/) servers. Note, the database N-termine for BACOVA_03425 GH43 is mis-annotated; the correct N-terminus actually starts 24 aa downstream at the sequence MKN.

Protease K treatment. Cultures of B. ovatus (50 ml) were grown in minimal media on WAX (0.5% w/v) as a sole carbon source, to mid-exponential growth phase (OD600 ~ 0.8). Cells were harvested by centrifugation and washed twice in 10 ml phosphate buffered saline pH 7.2 (PBS), before being resuspended in 2.5 ml PBS, pH 7.2. Cells were incubated at 4°C for up to 16 h, the cells were then pelleted, washed in 5 ml PBS, pH 7.2, before being resuspended in 500 μl PBS. The cells were then pelleted, washed and resuspended in 2.5 ml PBS and remaining Protease K precipitated by the addition of 200 μl trichloroacetic acid and incubation on ice for 30 min. The cells were pelleted by centrifugation and washed four times in 1 ml ice cold acetone. The cell pellets were resuspended in 250 μl Laemmli buffer and subjected to SDS-PAGE. Proteins were transferred to Whatman PTFE BA 85 nitrocellulose membrane using a wet transfer system (BioRad Mini Trans-Blot). Proteins of interest were detected using anti-sera raised in rats (Eurogentec) against the corresponding recombinant form of the protein, diluted 1/1,000 or 1/2,000 in 1% milk powder in TBS buffer. The secondary antibody used was a chicken anti-rat conjugated to horseradish peroxidase (Santa Cruz) diluted 1/500 in 1/5,000 in 1% milk powder in TBS buffer. Antibodies were detected by chemi-luminescence using Biorad Clarity Western ECL Substrate.

Immuno-fluorescence microscopy. B. ovatus suspensions of mid-exponential phase cells (0.5 ml in PBS, produced as for Protease K treatment above) were fixed with an equal volume of 2 × formalin (9% formaldehyde in PBS), and rocked for 90 min at 25°C in Eppendorf tubes. The cells were then pelleted by centrifugation for 3 min at 7,000 g and washed twice with 1 ml of PBS. The bacterial cell pellet was resuspended in 1 ml of blocking solution (2% goat serum, 0.02% NaN3 in PBS) and resuspended in 50 μl of antibody solution. The cells were incubated separately without the enzyme as a control. Both samples were then heated at 100°C for 10 min to inactivate the enzyme, and the amount of reducing sugar was quantified using the dinitrosalicylic acid method. The rye arabinoxylan used to assay the arabinofuranosidases was obtained from a pharmaceutical grade source, as described previously. The xyloglucan heptasaccharide and linear xylooligosaccharides were used as standards.

Glycoside hydrolase assays. Spectrophotometric assays. The activity of the GH31 β-xylanase and GH3 and GH43 β-xylosidases against xylans and xylooligosaccharides were assayed using a 1-xylose detection kit (Megazyme International). The reaction was carried out at 37 °C in 50 mM sodium phosphate buffer, pH 7.0 containing 2 mM MgCl2, 1 mM ATP and 1 mM NAD+, excess concentrations of linked enzymes (hexokinase, xylan mutarotase and β-xylose dehydrogenase) and 1 mg ml−1 BSA. The amount of free 2-chloro-4-nitrophenol released and NADH levels were monitored at 340 nm using an extinction coefficient of 6.220 M−1 cm−1. The xyloglucan heptasaccharide and linear xylooligosaccharide substrates used in this assay were purchased from Megazyme International. The glucurono-xylooligosaccharides, which were re-glycosylated by the remaining Proteinase K, were assayed against BGX, WAX and CX at 37°C and 100°C, respectively. To measure the relative activity of the GH10 enzymes against specific xylooligosaccharides.

To measure the activity of the GH95 enzymes, the 1-fose and 1-galactose products were incubated from 2′-fucose (Carbohydryl and CX), respectively, were monitored by HPAEC-PAD. The reactions were carried out in 20 mM sodium phosphate buffer, pH 7.0 at 37°C. In brief substrates were incubated with an appropriate concentration of each enzyme (10 nM to 5 μM). Aliquots were removed at regular intervals for up to 1 h and, after boiling for 10 min to inactivate the enzyme, the amount of the product was quantified by comparison to known standards. The activity of wild type and mutant Roch12X3 (10 μM substrate) was determined in 50 mM sodium phosphate buffer, pH 7.0 at 37°C against 2-chloro-4-nitrophenyl α-1-fose (Carbohydryl) was determined by continuously monitoring the release of 2-chloro-4-nitrophenol product at 400 nm.

The molar concentration of polysaccharide available to exo acting enzymes was determined by incubating the glycans at 1 mg ml−1 with a large excess of enzyme for 1 h to ensure the reaction goes to completion. The molar concentration of product at this point was then determined using the appropriate monosaccharide detection kit (detailed above). This equates to the molar concentration of total available substrate, thus enabling mg ml−1 of substrate to be converted to molarity.

Where kcat/Km alone is reported this was determined by monitoring the initial rate at a range of substrate concentrations significantly below the Km such that a linear plot of V0 vs [S] was obtained. Thus, the initial rate of substrate hydrolysis gives a direct readout of kcat/Km, using the equation kcat/Km = V0/[S]E0. Curve fitting to determine kcat/Km was performed using an analytical CARBOPACTM PA-100 anion-exchange column (Dionex) equipped with a guard column. The system had a loop size of 100 μl and was run at a flow rate of 1 ml min−1 at a pressure of 2 bar. Reactions were detected by pulsed amperometric detection (PAD) with electrode settings of E1 = 0.05, E2 = +0.6 and E3 = −0.6. The elution conditions were 0–10 min, 100 mM NaOH; 10–25 min, 100 mM NaOH with a 0–75 mM sodium acetate gradient; and 25–35 min, 100 mM sodium acetate containing 500 mM sodium acetate. For analysis of the very large oligosaccharides produced by the GH89 xylanase a 0-100% sodium borohydride gradient was run over 25–35 min, 100 mM NaOH with a 0–75 mM sodium acetate gradient; and 25–35 min, 100 mM NaOH containing 500 mM sodium acetate. The glucurono-xylooligosaccharides, which were re-glycosylated by the remaining Proteinase K, were assayed against BGX, WAX and CX at 37°C and 100°C, respectively. The xyloglucan heptasaccharide and linear xylooligosaccharides were used as standards.

Analysis of the GH89 (BACOVA_03433) products reduced with sodium borohydride. To analyse the specificity of BACOVA_03433, the recombinant enzyme (5 μM final) was incubated with 20 mg of CX resuspended in 0.5 ml 50 mM sodium phosphate buffer, pH 7.0 at 37°C for 24 h. The same amount of CX was incubated separately without the enzyme as a control. Both samples were then incubated overnight with NaBH4 (10 mg per sample) in the presence of NH4OH, pH 10.8 at 25°C. After incubation, samples were freeze-dried, resuspended in 500 μl of water and loaded separately onto Bio-gel P2 size-exclusion chromatography (Bio-Rad laboratories Inc.). Water was applied to the column (0.5 ml min−1) using a Pharmacia CKB peristaltic pump to elute carbohydrate products. A 4 μl aliquot of each of the 120 fractions (2 ml) was spotted on silica gel TLC plates, which were developed in butanol:acetic acid 2:1:1 and carbohydrate products detected by spraying with 0.5% (w/v) orcinol in 10% sulphuric acid and heating to 70°C for 10 min. The spots were visualised by cooling in a speed-vac. Each sample (enzyme and control) was then re-suspended in 20 μl of water and hydrolysed with 0.3 M HCl for 3 h at 100°C. The products of acid
hydrosylation were neutralized with 1 M NaOH and analysed by HPAEC using the vapour diffusion sitting drop method. Crystals of BACOVA_03438 in complex with L-galactose were determined to be P21 21 21 and a solvent content of 53%). The structure BACOVA_03438 was solved by molecular replacement in MOLREP using pdb file with accession code ZDY as the search model. The model underwent cycles of model building in OOT and refinement in REFMAC. Five percent of the observations were used to monitor the progress of refinement. The model was submitted to the PDB_REDO server to improve final model quality. The model was validated using Molprobity. The crystal and data collection statistics are reported in Supplementary Table 12.

Genetic manipulation of Bacteroides ovatus. Knockout strains lacking either PUL-XyII, PUL-Xyl or both PULs were created using counterselectable allelic exchange as described previously. The inactive genotype GH98 (BACOVA_03438) in P. tetracystica lacking both catalytic residues was made using the Quickchange site directed mutagenesis kit (Agilent) following the manufacturer’s protocol (Supplementary Table 10). The mutated gene was then cloned into pEX and integrated into the genome of B. ovatus using the gene counterselectable allelic exchange methodology as for the PUL knockouts. Signature-tagged strains of wild type and the AGH98 mutant, for determining the ratios of each during co-culture by qPCR, were made by integration of unique sequences into one of two att sites in the Bacteroides ovatus genome through homologous recombination of pNB2 conjugated into the Bacteroides from S17-1. pI pI E. coli as described previously. Following positive selection on BH-agar containing 250 mg l⁻¹ of tetracycline (1 µg ml⁻¹), genomic DNA was extracted (QIAamp DNA Mini Kit; Qiagen) and PCR with primers flanking each possible att insertion site was used to select for Bacteroides strains that had successfully tagged the integration site. Primers used for all genetic manipulations are shown in Supplementary Table 13.

Sharing assays. Bifidobacterium adolescentis and Bacteroides ovatus cells were grown anaerobically to mid/late-log phase in CNM (Sigma-Aldrich) and TYG rich media, respectively. Cells were washed in PBS and used, in approximately equal proportions, to inoculate Minimal Medium (based on medium devised by Van der Ploeg et al. 2015) containing 0.5% (w/v) BGX, WAX or CX as the sole carbon source. Cells were grown anaerobically to mid/late-log phase in CNM (Sigma-Aldrich) and TYG rich medium (CNM). Aliquots were taken at different phases of growth, serially diluted and plated onto CNM and CNM with 200 µg ml⁻¹ tetracycline (1 µg ml⁻¹). Crystals of BACOVA_03438 in complex with L-galactose were determined to be P21 21 21 and a solvent content of 53%). The structure BACOVA_03438 was solved by molecular replacement in MOLREP using pdb file with accession code ZDY as the search model. The model underwent cycles of model building in OOT and refinement in REFMAC. Five percent of the observations were used to monitor the progress of refinement. The model was submitted to the PDB_REDO server to improve final model quality. The model was validated using Molprobity. The crystal and data collection statistics are reported in Supplementary Table 12.

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Author contributions
D.N.B. and H.J.G. conceived the study. A.R., J.A.B, N.T., E.C.I., A.B., A.M.D., H.Z., T.E.R., E.C.M., P.D., B.H., H.J.G. and D.N.B. designed the experiments. A.R., J.A.B, N.T., E.C.L, A.M.D., H.Z., T.E.R., E.C.M, P.D., B.H., H.J.G. and D.N.B. wrote the manuscript.

Additional information
Accession codes: Coordinates and structure factors for RACOVA_03438 have been deposited in the RCSB Protein Data Bank under accession code 4UPC.

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Corrigendum: Glycan complexity dictates microbial resource allocation in the large intestine

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