This is an author produced version of Molecular Mechanism by which Prominent Human Gut Bacteroidetes Utilize Mixed-Linkage Beta-Glucans, Major Health-Promoting Cereal Polysaccharides.

White Rose Research Online URL for this paper:
http://eprints.whiterose.ac.uk/127256/

Article:
Tamura, Kazune, Hemsworth, Glyn R. orcid.org/0000-0002-8226-1380, Déjean, Guillaume et al. (7 more authors) (2017) Molecular Mechanism by which Prominent Human Gut Bacteroidetes Utilize Mixed-Linkage Beta-Glucans, Major Health-Promoting Cereal Polysaccharides. Cell Reports. pp. 417-430. ISSN 2211-1247

https://doi.org/10.1016/j.celrep.2017.09.049
**Molecular Mechanism by which Prominent Human Gut Bacteroidetes Utilize Mixed-Linkage Beta-Glucans, Major Health-Promoting Cereal Polysaccharides**

**Graphical Abstract**

**Highlights**
- The molecular mechanism of MLG utilization by *Bacteroides ovatus* is presented
- BoGH16<sub>MLG</sub> possesses structural features suited for efficient MLG degradation
- MLG utilization loci (MLGULs) serve as genetic markers for MLG catabolism
- Bacteroidetes MLGULs are ubiquitous in the gut microbiota of human populations

**Authors**
Kazune Tamura, Glyn R. Hemsworth, Guillaume Déjean, ..., Gideon J. Davies, Eric C. Martens, Harry Brumer

**Correspondence**
brumer@msl.ubc.ca

**In Brief**
Mixed-linkage β(1,3)/β(1,4)-glucan (MLG) is an important complex dietary polysaccharide (dietary fiber), the degradation of which in the human gut depends on the resident microbiota. Tamura et al. outline the molecular mechanism of MLG utilization by *Bacteroides ovatus* and reveal that the majority of surveyed humans possess MLG-utilizing Bacteroidetes.

**Data and Software Availability**
5NBO
5NBP

Tamura et al., 2017, Cell Reports 21, 417–430

October 10, 2017 © 2017 The Authors.
https://doi.org/10.1016/j.celrep.2017.09.049
Molecular Mechanism by which Prominent Human Gut Bacteroidetes Utilize Mixed-Linkage Beta-Glucans, Major Health-Promoting Cereal Polysaccharides

Kazune Tamura,1,4 Glyn R. Hemsworth,2 Guillaume Déjean,1 Theresa E. Rogers,4 Nicholas A. Pudlo,4 Karthik Urs,4 Namrata Jain,1,2 Gideon J. Davies,3 Eric C. Martens,4 and Harry Brumer1,7

1Michael Smith Laboratories, University of British Columbia, 2185 East Mall, Vancouver, BC V6T 1Z4, Canada
2Department of Biochemistry and Molecular Biology, University of British Columbia, 2350 Health Sciences Mall, Vancouver, BC V6T 1Z3, Canada
3Department of Chemistry, University of British Columbia, 2036 Main Mall, Vancouver, BC V6T 1Z1, Canada
4Department of Microbiology and Immunology, University of Michigan Medical School, 1150 West Medical Center Drive, Ann Arbor, MI 48109, USA
5Department of Chemistry, University of British Columbia, 3200 University Boulevard, Vancouver, BC V6T 1Z4, Canada
7Lead Contact

*Correspondence: brumer@msl.ubc.ca
https://doi.org/10.1016/j.celrep.2017.09.049

SUMMARY

Microbial utilization of complex polysaccharides is a major driving force in shaping the composition of the human gut microbiota. There is a growing appreciation that finely tuned polysaccharide utilization loci enable ubiquitous gut Bacteroidetes to thrive on the plethora of complex polysaccharides that constitute “dietary fiber.” Mixed-linkage β(1,3)/β(1,4)-glucans (MLGs) are a key family of plant cell wall polysaccharides with recognized health benefits but whose mechanism of utilization has remained unclear. Here, we provide molecular insight into the function of an archetypal MLG utilization locus (MLGUL) through a combination of biochemistry, enzymology, structural biology, and microbiology. Comparative genomics coupled with growth studies demonstrated further that syntenic MLGULs serve as genetic markers for MLG catabolism across commensal gut bacteria. In turn, we surveyed human gut metagenomes to reveal that MLGULs are ubiquitous in human populations globally, which underscores the importance of gut microbial metabolism of MLG as a common cereal polysaccharide.

INTRODUCTION

The composition and homeostasis of the human gut microbiota have a profound and intimate connection to various aspects of our physiology, health, and well-being (Littman and Pamer, 2011). Indeed, a multitude of diseases, such as type 2 diabetes, inflammatory bowel diseases (IBDs), and cancer, have been linked to alterations in the population and proportion of microbes in this highly complex and dynamic ecosystem that exists in our large intestine (Biedermann and Rogler, 2015; Fujimura et al., 2010; Kau et al., 2011; Schwabe and Jobin, 2013). The molecular mechanisms by which the microbiota exerts influence on human health are largely unresolved and undoubtedly complex yet may hold the key to personalized medicine through therapeutics that target the gut microbial ecosystem (Blanton et al., 2016; Haak et al., 2017; Kootte et al., 2012; Subramanian et al., 2015).

A major factor in shaping the composition and physiology of the gut microbiota is the influx of complex glycans—popularly known as “dietary fiber”—that evade degradation by the limited set of human-genome-encoded glycoside hydrolases (Hamaker and Tuncil, 2014; El Kaoutari et al., 2013; Koropatkin et al., 2012). Indeed, regular ingestion of plant polysaccharides is integral to maintaining a healthy balance of microbes in our lower gastrointestinal tract (De Filippo et al., 2010; Sonnenburg and Sonnenburg, 2014). Members of the Bacteroidetes, a dominant phylum in the human gut, possess an arsenal of polysaccharide utilization loci (PUL) to target a wide range of complex glycans (El Kaoutari et al., 2013). Analogous to the archetypal Bacteroides thetaiotaomicron starch utilization system (Sus), a hallmark of all Bacteroidetes PULs is the organization of genes clustered around tandem susC/susD homologs (encoding a TonB dependent transporter [TBDT] and a cell-surface glycan-binding protein [SGBP], respectively; Martens et al., 2009). Additional co-localized and co-regulated SGBP(s), a cohort of enzymes, and a transcriptional regulator typically make up a machinery that acts in concert to sense, break down, and import complex glycans (Grondin et al., 2017; Hemsworth et al., 2016). Many such PULs, each targeting specific glycan structures, have been identified by genomics and transcriptomics (see, e.g., the seminal study by Martens et al., 2011), but detailed functional characterization lags severely behind (reviewed in Grondin et al., 2017; Martens et al., 2014). Developing a precise understanding of the molecular details of complex glycan utilization by individual members of the microbiota is essential to designing targeted therapies based on prebiotics, probiotics, and symbiotics (Ciorka, 2012; Slavin, 2013), as well as novel therapeutic interventions.

Recently, comprehensive functional analysis has revealed the detailed molecular mechanisms by which individual PULs enable...
human gut Bacteroidetes to utilize predominant plant polysaccharides, including the matrix glycans, xyloglucan (Hemsworth et al., 2016; Larsbrink et al., 2014; Tauzin et al., 2016), xylan (Rogowski et al., 2015), β-mannan (Bøgenholm et al., 2017), and rhamnogalacturonan II (Ndeh et al., 2017). Mixed-linkage β(1,3)/β(1,4)-glycans (MLGs) (Figure 1A) from cereal grains constitute an additional key group of dietary glycans, whose utilization by gut microbes was previously unresolved at the molecular level. MLGs are abundant in the aleurone layer of common cereals, including oats (3%–8% dry weight) and barley (2%–20% dry weight; El Khoury et al., 2012). Beyond their obvious potential to contribute to energy intake (Cummings and Macfarlane, 1997; McNeil, 1984), MLGs have been linked to a range of health benefits, e.g., promoting healthy cholesterol and blood glucose levels, ameliorating insulin resistance, and mitigating metabolic syndrome (El Khoury et al., 2012). In particular, the cholesterol-lowering effect of oat MLG has long been recognized by the USA Food and Drug Administration (FDA) as well as the UK Joint Health Claims Initiative (JHCI) and been confirmed by subsequent studies (Othman et al., 2011).

The mechanisms behind these systemic benefits of MLG are, however, incompletely understood, in part due to a lack of understanding of MLG metabolism by individual members of the human gut microbiota. Thus, we report here the molecular characterization of a mixed-linkage glucan utilization locus (MLGUL) in the common symbiont B. ovatus. Identifying syntenic MLGUL...
in other Bacteroidetes revealed that, as the archetype, this MLGUL serves as a molecular marker for MLG utilization across the Bacteroidetes phylum, thereby enabling future functional prediction across species.

RESULTS

Identification of a Multi-gene Locus Responsible for MLG Utilization by B. ovatus

A putative MLGUL was previously identified in B. ovatus (Figure 1B) based on the presence of a tandem susC/susD homolog signature (Martens et al., 2009) and high-level expression of select genes in the presence of bMLG (Martens et al., 2011). Individual genes in the locus, BACOVA_02741-02745, were all substantially upregulated (125- to 298-fold) during growth on bMLG versus glucose as sole carbon sources (Table S1). BACOVA_02742 and BACOVA_02743 encode the signature TBDT/SGBP pair with 28% and 19% protein sequence identity to SusC and SusD, respectively. The putative MLGUL was additionally predicted to encode a second, non-homologous SGBP (BACOVA_02744), a hybrid two-component sensor/transcriptional regulator (HTCS) (BACOVA_02740), and up to three glycoside hydrolases (GHs).

BoGH16mLG (BACOVA_02741) is a member of a glycoside hydrolase family 16 (GH16) in the carbohydrate active enzymes (CAZy) classification (Cantarel et al., 2009). GH16 notably includes canonical bacterial MLG endo-glucanases (endo-MLGase) (Planas, 2000), along with a diversity of endo-galactanases and endo-galactanases (Eklof and Hehemann, 2016). BoGH3mLG (BACOVA_02745) is classified into glycoside hydrolase family 3 (GH3), whose members include exo-β-glucosidases (Fincher et al., 2017). Notably, we have determined that BACOVA_02738, which is predicted to encode a second GH3 exo-β-glucosidase, is unlikely to be part of the MLGUL for three reasons: (1) BACOVA_02738 was not significantly upregulated on MLG (1.6-fold versus glucose control; Table S1); (2) a corresponding gene is not found among syntenic loci (Figure 1B); and (3) the encoded enzyme was catalytically feeble compared to BoGH3mLG on β-glucosides relevant to MLG saccharification (vide infra).

To determine the correlation between the presence of the predicted MLGUL and growth of B. ovatus on MLG, we constructed an isogenic mutant of B. ovatus Δtdk (Larsbrink et al., 2014), in which a contiguous region of DNA-encoding genes BACOVA_02738–02745 was deleted (B. ovatus ΔMLGUL). Vis-à-vis the parent strain, the B. ovatus ΔMLGUL was able to grow equally well on glucose as the sole carbon source; however, the ability to grow on bMLG was completely abolished (Figure S1). Thus, the putative MLGUL is necessary to confer B. ovatus the ability to utilize MLG.

Enzymology and Structural Biology of BoGH16mLG, the Vanguard MLGase

Cellular Localization

The GH family membership of BoGH16mLG suggested a potential role as the vanguard enzyme catalyzing polysaccharide backbone cleavage at the cell surface as the essential first step in MLG utilization. Indeed, the presence of a predicted type II signal sequence (determined with LipoP 1.0; Juncker et al., 2003) suggested that the protein is membrane anchored via lipidation of the N-terminal cysteine residue (Paetzel et al., 2002). To validate this prediction, B. ovatus Δtdk was grown on minimal medium with either glucose or bMLG as a sole carbon source prior to immunolocalization of BoGH16mLG. As shown in Figure 2A, BoGH16mLG was clearly visualized on the outer surface of cells in which the presence of the polysaccharide induced MLGUL expression but was absent from cells grown on glucose (Figures S2C and S2F). Further analysis of cellular fractions by western blotting revealed the presence of BoGH16mLG in the membrane fraction, corroborating its attachment to the outer membrane (Figure 2C). Interestingly, BoGH16mLG was also detected in the lysate supernatant (soluble periplasmic or cytoplasmic proteins) and in the culture supernatant (secreted

Figure 2. Enzyme Localization Analysis

(A and B) Phase contrast microscopy and corresponding fluorescence microscopy images of B. ovatus Δtdk cells grown in minimal medium with bMLG as the sole carbon source probed with custom polyclonal antibodies against recBoGH16mLG (A) and recBoGH3mLG (B).

(C) Western blots of protein collected from the culture supernatant, cell lysate supernatant, and cell lysate membrane fraction of B. ovatus Δtdk cells grown in minimal medium with glucose or bMLG as a sole carbon source. See also Figure S2.
protein; Figure 2C). Whereas the former may represent anchored protein released into the soluble fraction during cell lysis, detection in the culture supernatant could result from packaging and release in outer membrane vesicles, which has previously been observed for other Bacteroidetes glycoside hydrolases (Elene
awy et al., 2014).

**Substrate and Product Specificity**

To verify the leading catalytic role of BoGH16\_MLG in MLG utilization and determine the specificity of the enzyme for individual β-glucans, recombinant BoGH16\_MLG produced in *E. coli* (recBoGH16\_MLG; Figures S3A and S3B) was screened for hydrolytic activity against a library of polysaccharides. No activity was observed on tamarind xyloglucan, beechwood xylan, wheat arabinoxylan, carob galactomannan, konjac glucomannan, synthetic carboxymethylcellulose, synthetic hydroxyethylcellulose, *Xanthomonas campestris* xanthan gum, or *Ulva* sp. ulvan. In this initial screen, BoGH16\_MLG was minimally active on all-β(1,3)-glucans, including *Laminaria digitata* laminarin, yeast β-glucan, and *Alcaligenes faecalis* curdlan, but exhibited high specific activity on bMLG. The optimum pH of 6.5 (consistent with function in the distal human colon) and maximum temperature range of 45°C to 55°C was determined using bMLG as substrate (data not shown).

Subsequent Michaelis-Menten kinetic analysis at the pH optimum and 37°C demonstrated that BoGH16\_MLG is a highly predominant mixed-linkage β(1,3)/β(1,4)-glucanase (MLGase), with a 33-fold higher specificity constant, $k_{cat}/K_m$, for bMLG (Figure 1A) over laminarin, an all-β(1,3)-glucan with single β(1,6)-linked glucosyl branches (Figure 3A; Table S2; Martin et al., 2007). BoGH16\_MLG was even less efficient on the other two all-β(1,3)-glucans for which activity was initially observed: the $k_{cat}/K_m$ was 147-fold higher for bMLG than yeast β-glucan (similar in structure to laminarin but with longer β(1,6)-linked glucose branches; Lowman et al., 2011) and nearly four orders of magnitude higher than that of high curdlan, a 22-kDa, non-branched β(1,3)-glucan (Harada et al., 1968; Figure 3A; Table S2).

Detailed product analysis was employed to determine the mode of hydrolysis, *endo* versus *exo*, and linkage specificity of recBoGH16\_MLG to gain information on the nature of the MLG cleavage products at the *B. ovatus* cell surface. High-performance liquid chromatography (HPLC) analysis at selected time points in the hydrolysis showed the initial production of very large oligosaccharide fragments, which were progressively converted into shorter species and ultimately to two distinct oligosaccharides in the limit digest (Figure 3B). This product evolution indicates that BoGH16\_MLG operates through an *endo*-dissociative mode of action, in which the MLG polysaccharide is stochastically cleaved along the backbone.

Comparison with oligosaccharide standards (Figure 3B) and additional liquid chromatography-mass spectrometry (LC-MS) analysis (data not shown) revealed that the limit digest products were the mixed-linkage trisaccharide, G4G3G (Glcβ(1–4) Glcβ(1–3) Glc), and the mixed-linkage tetrasaccharide, G4G4G3G (Glcβ(1–4) Glcβ(1–4) Glcβ(1–3) Glc). Thus, BoGH16\_MLG specifically hydrolyzes β(1,4)-linkages of glycosyl residues that are immediately preceded by a β(1,3)-linked glucosyl residue (toward the non-reducing chain end). This specificity is typical of

![Figure 3. BoGH16\_MLG Kinetics and MLGUL GHs Product Analysis](image)
bacterial endo-MLGases within GH16 (Gaiser et al., 2006; McGregor et al., 2017; Planas, 2000).

To provide more refined insight into BoGH16<sub>MLG</sub> substrate specificity, Michaelis-Menten kinetics was determined for a series of chromogenic glycosides (Figures S4A and S4B; Table S3). receBoGH16<sub>MLG</sub> had no activity on the ortho-chloro-para-nitrophenyl (CNP) β-glycosides of glucose (G-CNP), cellobiose (G4G-CNP), cellotriose (G4G4G-CNP), or para-nitrophenyl (pNP) β-glucoside (G-pNP). Weak activity was observed on the pNP and CNP β-glycosides of laminaribiose (G3G), consistent with a requirement for a β(1,3) linkage spanning the −2 to −1 active-site subsites (GH subsite nomenclature according to Davies et al., 1997), as was indicated by the bMLG limit-digest analysis (vide supra). Likewise, G4G3G-CNP and G4G4G3G-CNP were specifically and efficiently hydrolyzed to release the aglycone, with no cleavage of the internal glycosidic bonds. The specificity constants (k<sub>cat</sub>/K<sub>m</sub> values) for CNP release from these mixed-linkage tri- and tetrasaccharides were 800- and 1,500-fold greater than that of G3G-CNP, respectively, which indicate that potential −3 and −4 subsites contribute 17 kJ/mol and 1.6 kJ/mol to transition state stabilization (∆∆G<sub>T</sub>). Indeed, a very significant contribution from the −3 subsite is a common feature of GH16 endo-MLGases (Gaiser et al., 2006; McGregor et al., 2016; Planas, 2000).

**BoGH16<sub>MLG</sub> Tertiary Structure**

Three-dimensional structures of receBoGH16<sub>MLG</sub> were solved by X-ray crystallography to reveal the molecular basis for MLG recognition and hydrolysis. The apo structure of receBoGH16<sub>MLG</sub> was determined to a resolution of 1.8 Å by molecular replacement using the structure of *Zobellia galactanivorans* laminarinase ZgLamC<sub>GH16-E142S</sub> (PDB code 4CRQ; Labourel et al., 2015) as a search model (see Table S4 for processing and refinement statistics). The crystal contained two polypeptide chains in the asymmetric unit corresponding to residues I55–L271 of wild-type BoGH16<sub>MLG</sub> for both chains (residue numbering is from transcriptional start site according to the genomic sequence). No electron density was observed for the N-terminal His<sub>6</sub>-tag and subsequent 15 amino acids in either chain of the recombinant protein, which suggests that residues C19–D34 of the wild-type enzyme constitute a flexible linker sequence to distance the catalytic module from the outer membrane surface (residues M1–S18 comprise the predicted signal peptide); the sidechain of C19 would constitute the site of terminal lipida
tion (Paetzl et al., 2002). The overall fold of BoGH16<sub>MLG</sub> consists of a β-jelly roll architecture typical of other GH16 members (Davies and Sinnott, 2008): two antiparallel β sheets stack on top of each other with the concave face forming the polysaccharide substrate binding cleft. The end-on arrangement of the two chains in the asymmetric unit hinted at the possibility of the formation of a dimer (Figure 4A). Size-exclusion chromatography, however, indicated that BoGH16<sub>MLG</sub> exists as a monomer in solution (data not shown), which, together with steric considerations of polysaccharide binding through the active-site cleft, indicates that end-on contacts observed between chains A and B are artifacts of crystal packing.

The sidechains of the conserved GH16 catalytic residues (Planas, 2000), comprising Glu-143 (nucleophile), Asp-145 (electrostatic “helper”), and Glu-148 (acid/base), are presented on the same face of one β strand (i8), pointing into the active-site cleft. Notably, these residues are contained in an EXDXXE consensus sequence that is typical of bacterial GH16 laminarinases (i8,13-specific endo-glucanases). The insertion of an extra amino acid (underlined), typically methionine, results in a so-called “β-bulge” secondary structural motif that is not found among canonical bacterial MLGases, which instead possess a regular β strand (Barbeyron et al., 1998; Michel et al., 2001).

Commensurate with this observation, the closest eight structural homologs identified using the Dali server (Holm and Rosenström, 2010) feature a β-bulge active-site motif (Table S5). Specifically, the top match (Z score = 29.3) was the structure of laminarinase “ZgLamC<sub>GH16-E142S</sub>” from *Zobellia galactanivorans* (PDB code 4CTE; Labourel et al., 2015), which has 38% amino acid identity and superimposed with BoGH16<sub>MLG</sub> with a root-mean-square deviation (RMSD) value of 2.0 Å over 211 out of 231 Cα pairs. In comparison, the closest GH16 homolog with a regular active-site β strand was the lichenase (MLGase) from *Paeonia
culus macerans* (PDB code 1MAC; Hahn et al., 1995), which has a comparable Z score of 25.1 and an RMSD value of also 2.0 Å over 212 Cα pairs, despite having only 22% amino acid identity with BoGH16<sub>MLG</sub>.

Soaking crystals of the wild-type enzyme with G4G4G3G yielded a product complex with 1.8 Å resolution (Table S4). The complete tetrasaccharide was modeled in electron density spanning substrates −1 to −4 in the active-site cleft of chain A, whereas the electron density for the fourth glucosyl residue in substrate −4 was not resolved in chain B. This is most likely due to disorder of this residue because the corresponding −4 Glc in chain A is fully solvent exposed, makes no contact with the enzyme, and has significantly higher B factors (Figure 4B). These structural observations are consistent with kinetic data for chromogenic MLG oligosaccharides (Table 1), which likewise suggest the existence of three primary negative subsites, −1 to −3, and a weakly interacting −4 subsite.

In both chains A and B, the three glucosyl residues spanning substrates −1 to −3 are well defined and virtually identical. The reducing-end glucosyl residue in the −1 subsite is in the β-conformation, with the C1 hydroxyl hydrogen bonded to Tyr-181, which is observed in a dual conformation in both chains of the G4G4G3G complex (Figure 4C). Interestingly, this dual conformation is not observed in the apo form of the enzyme; Tyr-181 is “swung in” to the active site in chain B, whereas it is “swung out” in chain A, the sidechain from chain A stacking on top of the chain B sidechain (Figure 4D). The conformation of this sidechain will be key to determining the nature of the positive substrate-binding subsites; indeed, comparison with other GH16 endo-glucanases clearly suggests the presence of two positive subsites (Gaiser et al., 2006; Planas, 2000). Whether the dynamics observed for Tyr-181 are an artifact of crystallization or perhaps play a role in substrate binding and product release is unclear in the absence of an enzyme-substrate complex spanning the positive subsites.

With regard to specific interactions in the negative subsites, subsite −1 is further characterized by hydrogen bonds between Glu-143 and the C2 hydroxyl, Trp-125 and the C6 hydroxyl, as well as Glu-148 and the ring oxygen and the C1 hydroxyl. This
glucose is also positioned by a stacking interaction with Trp-125 and Trp-129 (Figure 4E), both of which are conserved across all GH16 laminarinases. At subsite –2, highly conserved Arg-97 forms a hydrogen bond with the C6 hydroxyl, and Asn-60 hydrogen bonds to the C2 hydroxyl as well as to the glucosidic bond oxygen between the –1 and –2 sugars. Another conserved residue, Trp-138, serves as a platform that stacks with the subsite –2 glucose. In subsite –3, the main interaction is stacking against Trp-58, which also forms a hydrogen bond to the glucosidic bond oxygen between the –3 and –4 sugars (Figure 4E). Together, these interactions in subsite –3 are responsible for 17 kJ/mol of transition-state stabilization (vide supra).

**Downstream Saccharification of Mixed-Linkage Oligosaccharides Produced by BoGH16**

To elucidate the mechanism for the downstream conversion of the oligosaccharide products of BoGH16, to glucose for primary metabolism, we examined the biochemistry of the two predicted exo-β-glucosidases, BoGH3MLG and BACOVA_02738(GH3), associated with the MLGUL.

**Cellular Localization of BoGH3**

BoGH3MLG and BACOVA_02738(GH3) were unambiguously predicted by SignalP 4.0 (Petersen et al., 2011) to contain a secretion signal peptide, whereas LipoP 1.0 (Juncker et al., 2003) additionally indicated a type II lipoprotein signal sequence (Paetzel et al., 2002) in BoGH3MLG only. The same *B. ovatus* Δtk cultures used for BoGH16MLG localization, containing glucose or bMLG as the sole carbon source, were probed using polyclonal antibodies independently raised against recombinant BoGH3MLG and the BACOVA_02738(GH3) gene product. Neither protein was detected on the cell surface by fluorescence microscopy, especially in the presence of bMLG, which induces BoGH16MLG production (Figures 2B and S2A). BoGH3MLG induction by bMLG was confirmed by a western blot of cellular fractions, which also confirmed that this enzyme is membrane anchored (Figure 2C).

In contrast, the BACOVA_02738(GH3) gene product was detected to a higher degree in *B. ovatus* cells grown in minimal medium with glucose as a sole carbon source compared to cells induced with bMLG (Figure S2B). The lack of upregulation by bMLG is consistent with transcriptional analysis, which showed a limited change in expression in bMLG versus glucose (1.6-fold), which was two orders of magnitude lower than all MLGUL members (Table S1). The higher detection in uninduced cells is explained by the high basal expression of BACOVA_02738(GH3) (more than an order of magnitude greater than all MLGUL members; Table S1). The lack of detection in

---

**Figure 4. BoGH16MLG Structural Biology**

(A) The overall structure of the BoGH16MLG G4G4G3G asymmetric unit containing two polypeptide chains shown from orthogonal views with the bound oligosaccharides in yellow and the transparent surface representation in white. Chain A cartoon is shown in cyan, and chain B cartoon is shown in slate blue throughout the figure.

(B) Mixed-linkage tetrasaccharide ligand modeled into chain A of BoGH16MLG with the opaque surface representation in gray and the oligosaccharide colored according to B factors. The glucose in subsite –4 is outside of the active site cleft and has significantly higher B factor than the glucose units in subsites –1 to –3.

(C) Tyr-181 rotamers observed in the complex structure with the 2Fo-Fc map of the tyrosines shown contoured at 0.5σ in gray.

(D) Tyr-181 residues observed in the apo structure with the 2Fo-Fc map of the tyrosines shown contoured at 0.5σ in gray.

(E) Wall-eyed stereo view of the active site of chain A of the BoGH16MLG G4G4G3G complex. Hydrogen bonding interactions are shown as dashed black lines, sugars are shown in yellow with its 2Fo-Fc map contoured at 1σ in orange, and the conserved GH16 active site residues are shown in purple. Hydrophobic stacking interactions in addition to hydrogen bonds position the mixed-linkage oligosaccharide in the negative subsite of BoGH16MLG.

See also Figures S5–S7 and Tables S4 and S5.
Table 1. Kinetic Parameters for the Hydrolysis of Various Substrates by BoGH3$_{MLG}$ and BACOVA_02738(GH3)

<table>
<thead>
<tr>
<th>Enzyme</th>
<th>Substrate</th>
<th>$k_{cat}$ (s$^{-1}$)</th>
<th>$K_m$ (mM)</th>
<th>$k_{cat}/K_m$ (s$^{-1}$ mM$^{-1}$)</th>
<th>Assay</th>
</tr>
</thead>
<tbody>
<tr>
<td>BoGH3$_{MLG}$</td>
<td>β-Glc-pNP</td>
<td>59.5 ± 1.46</td>
<td>2.95 ± 0.14</td>
<td>20.2</td>
<td>pNP</td>
</tr>
<tr>
<td></td>
<td>gentiobiose (G6G)</td>
<td>ND</td>
<td>ND</td>
<td>0.0571</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>cellobiose</td>
<td>5.52 ± 0.19</td>
<td>7.47 ± 0.48</td>
<td>0.739</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>cellotriose</td>
<td>22.1 ± 0.3</td>
<td>0.859 ± 0.033</td>
<td>25.7</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>cellotetraose</td>
<td>17.3 ± 0.5</td>
<td>0.687 ± 0.044</td>
<td>25.2</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>cellopentaose</td>
<td>19.4 ± 0.8</td>
<td>0.777 ± 0.060</td>
<td>25.0</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>cellohexaose</td>
<td>17.4 ± 0.4</td>
<td>0.747 ± 0.041</td>
<td>23.3</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>laminaribiose</td>
<td>28.0 ± 1.1</td>
<td>1.90 ± 0.12</td>
<td>14.7</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>laminaritriose</td>
<td>34.2 ± 1.0</td>
<td>0.911 ± 0.052</td>
<td>37.5</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>laminaritetraose</td>
<td>31.3 ± 2.3</td>
<td>0.896 ± 0.135</td>
<td>34.9</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>laminaripentaose</td>
<td>39.5 ± 3.4</td>
<td>1.27 ± 0.20</td>
<td>31.1</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>MLGO$_3$ A (G3G4G)</td>
<td>61.6 ± 1.6</td>
<td>0.997 ± 0.040</td>
<td>61.8</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>MLGO$_3$ B (G4G3G)</td>
<td>24.7 ± 1.3</td>
<td>0.521 ± 0.064</td>
<td>47.4</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>MLGO$_4$ A (G3G4G3G)</td>
<td>55.7 ± 2.7</td>
<td>1.33 ± 0.12</td>
<td>41.9</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>MLGO$_4$ B (G4G3G3G)</td>
<td>30.8 ± 2.0</td>
<td>0.736 ± 0.106</td>
<td>41.8</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td></td>
<td>MLGO$_4$ C (G4G3G4G)</td>
<td>15.7 ± 0.3</td>
<td>0.601 ± 0.031</td>
<td>26.1</td>
<td>HK/G6PDH</td>
</tr>
<tr>
<td>BACOVA_02738 (GH3)</td>
<td>β-Glc-pNP</td>
<td>0.212 ± 0.004</td>
<td>2.53 ± 0.13</td>
<td>0.0838</td>
<td>pNP</td>
</tr>
</tbody>
</table>

ND, not determined (in cases where Michaelis-Menten curve fitting was not feasible, individual $k_{cat}$ and $K_m$ values are not reported and $k_{cat}/K_m$ value was determined from linear curve fit to initial-rate data in the [$S$] $<$ $K_{m(apparent)}$ range). Data are represented as the fitted parameters ± SD. See also Figures S4C and S4D.

*Biologically relevant substrates that BoGH3$_{MLG}$ encounters in the periplasmic space.

Minimal medium containing bMLG is due to high amounts of induced MLGUL proteins diminishing the presence of the BACOVA_02738(GH3) gene product when normalized to total protein (Figure S2B).

**Substrate Product Specificity of BoGH3$_{MLG}$ and BACOVA_02738(GH3)**

Initial activity screening on chromogenic pNP glycosides (see Experimental Procedures) revealed that both recBoGH3$_{MLG}$ and recBACOVA_02738(GH3) are specific exo-β-glucosidases (activity on other pNP glycosides was undetectable at micromolar enzyme concentrations). However, recBACOVA_02738 (G3H3) is strikingly feeble compared to recBoGH3$_{MLG}$ on G-β,pNP ($k_{cat}/K_m$ values of 0.084 mM$^{-1}$ s$^{-1}$ versus 20 mM$^{-1}$ s$^{-1}$; Figures S4C and S4D; Table 1). Further, measuring Michaelis-Menten kinetic parameters on cello- and laminari-oligosaccharides was not feasible due to overall poor activity and low protein production yields (data not shown). These kinetic results corroborate the above comparative genetic and transcriptional analyses, collectively suggesting BACOVA_02738(GH3) is not integral to the MLGUL. Hence, this enzyme was not characterized further.

To investigate oligosaccharide substrate preference of the BoGH3$_{MLG}$, we conducted initial-rate kinetics analyses on a series of gluco-oligosaccharides of distinct linkage composition and degrees of polymerization (d.p.). The non-reducing-end glucose was hydrolyzed from all-β(1,4)-linked cello-oligosaccharides (d.p. 2–6), all-β(1,3)-linked laminari-oligosaccharides (d.p. 2–5), and mixed-linkage β(1,3)/β(1,4)-gluco-oligosaccharides (d.p. 3 to 4; 5 examples) with comparable efficiencies, according to classic Michaelis-Menten saturation kinetics (Figures S4E and S4F; Table 1). In this series, only cellobiose (G4G) was poorly hydrolyzed by BoGH3$_{MLG}$ vis-à-vis the β(1,3)-linked congener laminaribiose (G3G) and all other gluco-oligosaccharides (e.g., G4G has a $k_{cat}/K_m$ value 20-fold lower than G3G; Table 1). The β(1,6)-linked diglucoside gentiobiose (G6G) was also a very poor substrate, with a $k_{cat}/K_m$ value 260-fold lower than that of G3G. Gluco-oligosaccharides with a β(1,3)-linked glucosyl unit at the non-reducing end all have slightly higher $k_{cat}$ Values compared to those with a β(1,4) linkage in this position, which typically contributes to higher $k_{cat}/K_m$ values for the former when substrates of equal d.p. are compared. However, the magnitude of these differences, which are often less than 2-fold, indicate that BoGH3$_{MLG}$ is essentially agnostic to β(1,3) versus β(1,4) linkages. These results also suggest that, in addition to a single negative subsite (−1) commensurate with its exo activity, BoGH3$_{MLG}$ has only two positive subsites that contribute to catalysis: in each gluco-oligosaccharide series, tetrasaccharides and larger are hydrolyzed with identical $k_{cat}/K_m$ values to the corresponding trisaccharides.

Product analysis following extended incubation of BoGH3$_{MLG}$ with G4G4G3G and G4G3G demonstrated that BoGH3$_{MLG}$ completely degrades the BoGH16$_{MLG}$ limit digest products to glucose. HPLC also revealed that laminaribiose (G3G) is the only new intermediate formed during the course of hydrolysis (Figure 3B). This demonstrates that BoGH3$_{MLG}$ sequentially hydrolyzes one glucose unit at a time from the non-reducing end of MLG oligosaccharides, viz.: G4G4G3G → G + G4G3G (also present in the starting mixture) → G + G3G → G + G. Notably, the individual $k_{cat}$ and $K_m$ values for each step are nearly identical (Table 1).
BoGH3\(_{\text{MLG}}\) and BACOVA_02738(GH3) Primary Structures

Despite extensive efforts, we were unable to crystallize the key \(\beta\)-glucosidase BoGH3\(_{\text{MLG}}\) for experimental tertiary structure determination. However, BoGH3\(_{\text{MLG}}\) has 63% sequence identity to a \(B.\ ovatus\) \(\beta\)-glucosidase (BoGH3B; PDB code 5JP0; Hemsworth et al., 2018) from the xyloglucan utilization locus (Figure S5A) and, as such, was amenable to tertiary structure homology modeling. Phyre2 (Kelley et al., 2015) utilized PDB code 5JP0 as the sole template, and 728 out of 742 residues (98% of the sequence, excluding the signal peptide) were modeled with 100% confidence. The model suggests that BoGH3\(_{\text{MLG}}\) possesses a homologous three-domain architecture with the active site being formed at the interface of the \((\alpha/\beta)_8\) triosephosphate isomerase (TIM) barrel and \(\alpha/\beta\) sandwich domains (Figure S5B). The predicted catalytic nucleophile and acid/base residues are Asp-309 and Glu-453, respectively, based on primary and tertiary alignment (Figures S5A and S5C). Two tryptophan residues were modeled on opposite sides of the entrance to the active site pocket (Figure S5D), forming a narrow “coin slot,” which may confer a preference toward \(\beta\)(1,3)- and \(\beta\)(1,4)-linked glucans and poor activity against \(\beta\)(1,6)-linked gentiobiose (Table 1). In contrast, enzymes that lack a homologous Trp-453 have a more open entrance to the active site and show broad activity against \(\beta\)(1,2)- and \(\beta\)(1,6)-linked glucans in addition to \(\beta\)(1,3)- and \(\beta\)(1,4)-linked glucans (Karkehadi et al., 2014; Pozzo et al., 2010).

In comparison, BACOVA_02738(GH3) possess catalytic residues homologous to BoGH3\(_{\text{MLG}}\) and similar GH3 members, despite having only 31% sequence identity to BoGH3\(_{\text{MLG}}\) (Figure S5A). The most similar characterized GH3 member to BACOVA_02738(GH3) among ~300 members identified in the CAZy Database is a Chrysosporium lucknowense \(\beta\)-glucosidase with 39% sequence identity (Dotenko et al., 2012).

Syntenic MLGULs Are Molecular Markers of MLG Utilization across the Bacteroidetes

Refined functional characterization of the catalytic specificity of GH components significantly increases confidence in the use of individual PULs as genetic markers of complex carbohydrate metabolism among Bacteroidetes (Cuskin et al., 2015; Larsbrink et al., 2014; Rogowski et al., 2015; Sonnenberg et al., 2010). The MLGUL characterized here represents the only PUL in \(B.\ ovatus\) that confers growth on MLG from cereals. To understand the wider distribution of MLG metabolic capacity among symbiotic Bacteroidetes in the human gut, we correlated the presence of a syntenic MLGUL with growth on bMLG for 354 individual Bacteroidetes strains representing 29 different species.

A total of 121 strains across just 7 of the species were able to grow on bMLG (Figure 5). In particular, 33 of 33 \(B.\ ovatus\) strains (including the type strain ATCC 8483) grew well on bMLG, as did 44 of 45 strains of the closely related \(B.\ xylanivorans\). The majority of \(B.\ uniformis\) strains tested (33 out of 35) were also competent bMLG utilizers. The limited penetration of the MLGUL across the genus clearly demonstrates nutrient-niche specialization among individual Bacteroidetes species.

Comparative analysis of available genomic sequences revealed that strains able to grow on bMLG as the sole carbon source harbor a syntenic MLGUL (Figure 1B). Previous transcriptional analysis demonstrated that the syntenic MLGUL found in \(B.\ celluloslyticus\) is also activated during growth on bMLG (McNulty et al., 2013). Concordance between the presence of a syntenic MLGUL and the ability to utilize MLG is further highlighted by the lack of a MLGUL in the \(B.\ uniformis\) ATCC 8492, one of only two strains of \(B.\ uniformis\) that could not grow on bMLG. Based on this analysis, we can also predict MLG utilization ability in two sequenced species of \(Prevotella,\ Pr.\ copri\) DSM 18205 and \(Pr.\ multituberculosis\) DSM16608, important members of the Bacteroidetes from the human gut and oral cavity, respectively (Figure 1B).

DISCUSSION

A Molecular Model for MLG Utilization by \(B.\ ovatus\)

Our current suite of data suggests a model by which the MLGUL gene products work in concert to enable the utilization of MLG (Figure 6), analogous to that of other PUL-encoded systems (Grondin et al., 2017). Thus, BoGH16\(_{\text{MLG}}\) is anchored to the outer membrane, where it plays a leading role in fragmenting large MLG polysaccharide chains (typical d.p. 700–5,000, depending upon the plant species of origin; Lazaridou et al., 2004; Zheng et al., 2011) into oligosaccharides that can be imported into the periplasm via the TBDT. Notably, the specific limit digest products of BoGH16\(_{\text{MLG}}\) endo-hydrolysis identified here, viz. the trisaccharide G4G3G and the tetrasaccharide G4G4G3G (Figure 3B), have been shown previously to bind the periplasmic sensor domain of the HTCS encoded by BACOVA_02740 (Ko 300 \(\mu\)M and 400 \(\mu\)M, respectively), whereas the intact MLG polysaccharide does not (Martens et al., 2011). Monomeric glucosic, all-\(\beta\)(1,4)-linked cello-oligosaccharides, and all-\(\beta\)(1,3)-linked laminari-oligosaccharides are also not bound by the HTCS (Martens et al., 2011), indicating that the unique linkages present in MLG are central to inducing the MLGUL system. Thus, there is an essential yet distant interplay between the outer-membrane-localized MLGase and the HTCS in specific nutrient sensing.

It is therefore likely that the BoGH16\(_{\text{MLG}}\) limit digest products, or minimal repeats of these structures ([(G4G3G)\(_{4,6}\)(G4G3G)\(_{4,6}\)]), comprise the main products transported through the TBDT in vivo. Recent studies on inulin (\(\beta\)(2,1)-fructan) utilization suggest that some TBDTs are able to import longer polysaccharide chains (e.g., d.p. > 20; Rakoff-Nahoum et al., 2016). Regardless of length, the efficient exo-hydrolytic activity of BoGH3\(_{\text{MLG}}\) in the periplasm is sufficient to completely saccharify all imported oligosaccharides to glucose (Figure 3B), to feed primary metabolism in the cytosol. In this process, the trisaccharide substrate of the HTCS, G4G3G, will always be generated, regardless of the imported saccharide chain length, ensuring continual production of the MLGUL upregulation signal until substrate is exhausted. Interestingly, BoGH3\(_{\text{MLG}}\) will never encounter cellobiose (G4G), toward which it has relatively weak activity (Figure S4F; Table 1), in this process; the final step of saccharification of MLGOs is the hydrolysis of the competent substrate laminaribiose (G3G).

Figure 1
Previous phylogenetic analyses of GH16 have suggested that the evolution of the active-site β-bulge motif EXDXXE, which is widespread among clan GH-B (comprising GH16 and GH7), to a regular β-strand motif EXDXE is a defining feature that delineates endo-β(1,3)-glucanases (laminarinases; EC 3.2.1.39 and EC 3.2.1.6) from mixed-linkage endo-β(1,3)/β(1,4)-glucanases (licheninases; EC 3.2.1.73), respectively (Barbeyron et al., 1998; Michel et al., 2001). In this context, the observation that BoGH16 MLG is highly specific for MLG, despite having a β-bulge motif, was surprising.

Using the CAZy Database as a starting point (http://www.cazy.org/GH16_characterized.html), together with mining of the primary literature, we generated a contemporary maximum-likelihood phylogeny of all biochemically characterized GH16 members (Figure S6). This analysis indicates that, although canonical, normal β-strand MLGases do form a monophyletic group as previously observed, MLGase activity is in fact widespread among the historical “laminarinae” group, in which BoGH16MLG is itself positioned. Despite currently limited and disparate kinetic data for individual enzymes, it also appears that it is not possible to define further substrate-specific clades within this group based on molecular phylogeny alone, in light of weak bootstrap support. This precludes defining any single evolutionary event giving rise to unique trajectories for the further diversification of extant laminarinases and MLGases in this clade. Instead, it appears that diverse, subtle mutations have allowed the independent evolution of predominant laminarinase or MLGase activity numerous times across a flat evolutionary landscape. As such, we suggest that this GH16 subgroup should be more generally referred to as the “laminarinase/MLGase group” going forward.

Detailed tertiary structural comparison of 10 β-bulge-containing members of this laminarinase/MLGase group revealed, however, that predominant laminarinases harbor a significantly more protruding loop (which is often, but not always, longer) connecting strands β2 and β3 than predominant MLGases (Figures S7A and S7B). Structural superposition with the BoGH16MLG-G4G4G3G complex indicates that this loop in predominant laminarinases would clash with MLG in the negative subsites, instead favoring binding of an all-β(1,3)-glucan that curves away from this loop. Such curvature is exemplified by the ZgLamC_{G16-1425}thio-β(1,3)-trisaccharide structure (Figure S7A; PDB code 4CTE; Labourel et al., 2015). Indeed, Ilari et al. (2009) observed that shortening the homologous loop in...
LamA from the archeon *Pyrococcus furiosus* (Figure S7A; PDB code 2VY0) by 4 amino acids increased the activity toward MLG by 10-fold. Likewise, BglF from *Nocardiopsis* sp. F96 (Figure S7B; PDB code 2HYK) and LamR from *Rhodothermus marinus* (Figure S7B; PDB code 3ILN), which have a 3.3- and 8.5-fold greater specificity constant and specific activity, respectively, toward MLG than laminarin, also have a smaller loop, similar to BoGH16MLG in this position. The canonical, regular β strand MLGase from *Paenibacillus macerans* (Figure S7C; PDB code 1MAC) and *Bacillus licheniformis* (Figure S7C; PDB code 1GBG) similarly have a small loop at this position.

Taken together, these analyses reveal a complex evolutionary landscape that computational phylogenetic analysis fails to resolve. Despite using a manually curated, structure-based sequence alignment as input, the maximum-likelihood numerical approach did not delineate the members of the laminarinase/MLGase group on the basis of the distinct active-site loop differences observed in tertiary structures (Figure S7). Instead, the phylogeny was likely obfuscated by diverse, random variations in amino acid composition across the entire β sandwich domain, which clearly limits large-scale, unsupervised phylogenetic analysis of these GH16 members. Moreover, analysis of both MLG and laminarin specificity (as a minimum) for individual members of this group, in light of their tertiary structures, is essential to avoid potential mis-annotation of these enzymes.

**Mining Metagenomic Data Reveals the Ubiquity of MLG Utilization in the Human Gut and Beyond**

Using syntenic MLGULs as genetic markers, we surveyed the publicly available metagenome data of 426 adults to understand the capacity of human populations to derive nutrition
from cereal MLGs. We were able to distinguish the species of origin based on nucleotide sequence except for MLGULs from *B. ovatus* and *B. xylanisolvens*, which were strikingly similar at 97% nucleotide identity. The *B. ovatus*/*B. xylanisolvens* and *B. uniformis* MLGULs are the most prevalent; both are observed in about 70% of the total human cohort (Figure 7). The *Pr. copri* MLGUL is more often the sole MLGUL of an individual than that of *B. cellulosilyticus* when only one is present (Figure 7, cyan lines), despite the latter being more frequent in total. Overall, 92.5% of the subjects harbor at least one of the five different MLGULs identified in this study, irrespective of nationality or whether they are diseased. MLGULs are ubiquitously detectable despite variability in sampling depth across different metagenomics sequencing projects (Figure 7). The prevalence of MLGULs across different nationalities is consistent with MLG from cereal grains being a ubiquitous component of the human diet. Indeed, the importance of MLG utilization is underscored by the upregulation of the MLGUL in the ceca of mice fed a complex plant cell wall diet (Martens et al., 2011). Similar widespread global distribution in human populations has been observed for xylloglucan utilization loci (Larsbrink et al., 2014). These observations are sharply contrasted by the PUL that mediates utilization of the red algal polysaccharide porphyran, which is essentially confined to subjects from Japan, where seaweed is a common part of the diet (Hehemann et al., 2010; Larsbrink et al., 2014). Interestingly, we were unable to detect MLGULs in four unweaned infants sampled in the Japanese metagenome project (data not included in our analysis of adult metagenomes). This is consistent with a dearth of Bacteroidetes in infants who receive the bulk of their nutrition from milk and are not yet consuming plant polysaccharides (Kurokawa et al., 2007).

Moving beyond the human microbiota, we can likewise predict MLG utilization ability in *Dysgonomonas gadei* and *Pr. oryzae* (formerly Xylanibacter oryzae) based on the presence of a syntenic MLGUL. These species are commonly found in the termite hindgut and decomposing rice straw, respectively. This provides direct evidence that an analogous MLG utilization system is employed by Bacteroidetes operating in environments beyond the human gut.

**Conclusions**

Complex carbohydrates that promote the growth of beneficial microbes in our distal large intestine are a cornerstone of a healthy diet. MLGs in particular have long been known to impart healthful effects (Othman et al., 2011), yet its mechanism of utilization for fermentation by gut microbes was unknown. Our work here sheds light on the fine-tuned mechanism that *B. ovatus* and other Bacteroidetes has evolved to efficiently utilize MLGs in the highly competitive environment of the human gut microbiota. The finding that a majority of humans possess microbes that can utilize this ubiquitous cereal polysaccharide highlights the relevance of potential therapeutic interventions based on MLG utilization to the general population. The present study also sets the stage for future work to understand the quantitative contributions of individual members of the microbiota and their cognate enzymes to MLG utilization in the human gut (Patrascu et al., 2017; Zhong et al., 2015).

**EXPERIMENTAL PROCEDURES**

**Microbiology**

*B. ovatus* gene deletions were constructed using allelic exchange as previously described (Koropatkin et al., 2008). Anaerobic growth profiles were
measured as previously described (Martens et al., 2011). Details of localization analysis by immunofluorescence microscopy and immunoblotting are provided in the Supplemental Experimental Procedures.

**Cloning, Expression, and Purification of Recombinant Enzymes**

The genes encoding BoGH16<sub>macL</sub>, BoGH3<sub>macL</sub>, and BACOVA<sub>02738</sub>(GH3) were cloned into expression vectors for recombinant production in E. coli. Details of cloning strategies, production, and purification are provided in the Supplemental Experimental Procedures.

**Enzyme Kinetics and Product Analysis**

Thorough kinetic analysis was conducted on a panel of polysaccharides, oligosaccharides, and chromogenic substrates. Products of enzymatic reactions were analyzed by HPAC-PAD and HILIC-MS. Details of enzymatic assays, analytical methods, as well as sources of commercial substrates are provided in the Supplemental Experimental Procedures.

**X-Ray Crystallography**

Crystals of BoGH16<sub>macL</sub> were screened and optimized by sitting drop vapor diffusion method. The structures of the apo- and G4G4G3G<sub>macL</sub>-BoGH16<sub>macL</sub> were solved by molecular replacement. Details of crystallization, data collection, and structure solution are provided in the Supplemental Experimental Procedures.

**Bioinformatics**

Phylogenetic analysis of select GH16 and GH3 sequences was conducted based on structure-guided alignment. Metagenomic survey was carried out by nucleotide BLAST of MLGUL sequences against various metagenome sequence data. Details are provided in the Supplemental Experimental Procedures.

**Statistical Analysis**

All kinetic assays were done in triplicate. Michaelis-Menten parameters are reported as fitted values ± SD throughout the article. All growth experiment results are averages of two biological replicates.

**DATA AND SOFTWARE AVAILABILITY**

The accession numbers for the atomic coordinates and structure factors of apo- and G4G4G3G-complexed BoGH16<sub>macL</sub> reported in this paper are PDB: 5NBO and PDB: 5NBP, respectively.

**SUPPLEMENTAL INFORMATION**

Supplemental Information includes Supplemental Experimental Procedures, seven figures, and five tables and can be found with this article online at https://doi.org/10.1016/j.celrep.2017.09.049.

**AUTHOR CONTRIBUTIONS**

K.T. cloned, expressed, and purified recombinant enzymes; conducted and analyzed kinetics for hydrolysis of polysaccharides, oligosaccharides, and chromogenic substrates; determined hydrolysis products; conducted phylogenetic and structural analyses; carried out metagenomics survey; and wrote the article. G.R.H. determined the crystal structures of apo- and G4G4G3G-complexed BoGH16<sub>macL</sub>. G.D. conducted enzyme localization analyses. T.E.R., N.A.P., and K.U. conducted reverse genetics and growth analyses. N.J. synthesized G3G-CNPh. G.J.D., E.C.M., and H.B. designed and directed research and co-wrote the article with input from all authors.

**ACKNOWLEDGMENTS**

Work in Vancouver was supported by operating grants from the Canadian Institutes for Health Research (MOP-137134 and MOP-142472) and infrastructure support from the Canadian Foundation for Innovation (Project #30663) and the British Columbia Knowledge Development Fund. Work at the University of Michigan was supported by the U.S. NIH (GM124136). G.J.D. is supported by the Royal Society “Ken Murray” Research Professorship. We thank Diamond Light Source (Harwell, UK) for access to beamlines I02 and I03 (proposal mx9948) that contributed to the results presented here. We thank Constance M. Bahr (Koropatkin group; U. Michigan) for invaluable assistance with microscopy. We thank Adriana Cabrera (Brumer group) for preparing laminarin by sodium borohydride reduction of laminarin. We thank Nicholas McGregor (Brumer group) for assistance with LC-MS. We thank Alexander H. Viborg (http://research.ahv.dk/) for access to CAzy database tools. We thank Hila Behar (Brumer group) for assistance with BLAST analysis of human metagenome sequences.

Received: June 2, 2017

Revised: August 30, 2017

Accepted: September 14, 2017

Published: October 10, 2017; corrected online November 7, 2017

**REFERENCES**


