

ORIGINAL ARTICLE

Functional characterization of polysaccharide utilization loci in the marine *Bacteroidetes* ‘*Gramella forsetii*’ KT0803

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Members of the phylum *Bacteroidetes* are abundant in many marine ecosystems and are known to have a pivotal role in the mineralization of complex organic substrates such as polysaccharides and proteins. We studied the decomposition of the algal glycans laminarin and alginate by ‘*Gramella forsetii*’ KT0803, a bacteroidetal isolate from North Sea surface waters. A combined application of isotope labeling, subcellular protein fractionation and quantitative proteomics revealed two large polysaccharide utilization loci (PULs) that were specifically induced, one by alginate and the other by laminarin. These regulons comprised genes of surface-exposed proteins such as oligomer transporters, substrate-binding proteins, carbohydrate-active enzymes and hypothetical proteins. Besides, several glycan-specific TonB-dependent receptors and SusD-like substrate-binding proteins were expressed also in the absence of polysaccharide substrates, suggesting an anticipatory sensing function. Genes for the utilization of the beta-1,3-glucan laminarin were found to be co-regulated with genes for glucose and alpha-1,4-glucan utilization, which was not the case for the non-glucan alginate. Strong synteny of the PULs of ‘*G. forsetii*’ with similar loci in other *Bacteroidetes* indicate that the specific response mechanisms of ‘*G. forsetii*’ to changes in polysaccharide availability likely apply to other *Bacteroidetes*. Our results can thus contribute to an improved understanding of the ecological niches of marine *Bacteroidetes* and their roles in the polysaccharide decomposition part of carbon cycling in marine ecosystems.

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Introduction

Bacteroidetes represent important contributors to the mineralization of complex organic substrates such as proteins and polysaccharides in the marine realm (Kirchman 2002; Bauer *et al.*, 2006; González *et al.*, 2008; Teeling *et al.*, 2012; Fernández-Gómez *et al.*, 2013). An important source of such organic matter is the marine phytoplankton that has been estimated to be responsible for about 50% of the global net primary production (Field *et al.*, 1998). Polysaccharides constitute a substantial fraction of this primary

production, and algae can be composed of up to 50% or more of polysaccharides (Kraan, 2012). Marine *Bacteroidetes* are assumed to have a key role in the utilization of these algal polysaccharides (Kirchman, 2002; Pinhassi *et al.*, 2004; Bauer *et al.*, 2006; González *et al.*, 2008; Teeling *et al.*, 2012; Fernández-Gómez *et al.*, 2013), but only for few cultivable members this has been investigated in functional studies so far (Thomas *et al.*, 2012). Recently, it could be shown that during a North Sea diatom bloom specialized populations of *Bacteroidetes* coated in the successive decomposition of algal-derived organic matter (Teeling *et al.*, 2012). This supports the assumption that the wealth of naturally occurring algal polysaccharide structures and compositions provides distinct ecological niches for heterotrophic marine bacteria (Grossart *et al.*, 2005; Hehemann *et al.*, 2010; Gomez-Pereira *et al.*, 2010; Gomez-Pereira *et al.*, 2012). However, so far

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respective niche specializations of different members of the *Bacteroidetes* are only poorly resolved. In order to gain deeper insights into the relationship between abundant algal polysaccharides and the gene repertoires of adapted *Bacteroidetes*, we studied the isolate '*Gramella forsetii*' KT0803 (Eilers *et al.*, 2001). This strain belongs to the class *Flavobacteria* of the phylum *Bacteroidetes* and has been isolated from eutrophic coastal surface waters near the island Helgoland in the German Bight (54°11'03"N, 7°54'00"E). A previous whole-genome analysis (Bauer *et al.*, 2006) of '*G. forsetii*' has revealed various genetic modules encoding carbohydrate-active enzymes (CAZymes) and transporters, frequently arranged in so-called polysaccharide utilization loci (PULs; Sonnenburg *et al.*, 2010). Such PULs have also been reported for commensal *Bacteroidetes* from the human intestine, and are specifically induced by oligosaccharides (Martens *et al.*, 2011), including such of algal origin (Hehemann *et al.*, 2012). In this proteogenomic study, '*G. forsetii*' was studied during growth on two typical algal glycans, laminarin and alginate, in comparison to a glucose-supplemented control. Beta-1,3-glucans of the laminarin type act as a storage polysaccharide in brown algae, *Chryso-phyceae*, diatoms and haptophytes such as *Phaeo-cystis* spp. where they can make up about 30–50% of the cell dry weight (Beattie *et al.*, 1961; Janse *et al.*, 1996; Davis *et al.*, 2003). Alginate on the other hand is a cell wall polysaccharide as well as an intracellular component in brown algae that can account for up to 40% of algal dry weight (Haug *et al.*, 1966; Davis *et al.*, 2003). It is an unbranched copolymer composed of 1,4-linked beta-D-mannuronate (M) and alpha-L-guluronate (G) that are arranged in alternating M-blocks, G-blocks and mixed M/G-blocks. Relevant proteins for the decomposition of these polysaccharides were determined by stable isotope labeling, followed by subcellular fractionation and high-accuracy mass spectrometry. This combinatory approach allowed for an enhanced precision in protein quantification and revealed two major PULs of '*G. forsetii*' that were specifically controlled by either alginate or laminarin. Data of this study furthermore indicate a cross-regulation of genes acting on similar polysaccharides such as laminarin and 1,4-glucans and revealed that some proteins so far designated as 'hypothetical' might contribute to polysaccharide decomposition. Particular progress was achieved with respect to possible substrate specificities within the highly variable group of TonB-dependent receptors (TBDRs), which are known to be of crucial importance for uptake of glycans by marine Gram-negative bacteria (Tang *et al.*, 2012).

Materials and methods

Culture conditions and metabolic labeling

'*G. forsetii*' KT0803 was grown at 25 °C and 170 r.p.m. in an artificial seawater medium, as

described by Schut *et al.* (1993), in batches of 100 ml in 500 ml shaking flasks until the mid-exponential phase ($OD_{600} \approx 0.5$). The medium was supplemented with either ^{14}N naturally labeled or ^{15}N -labeled ammonium chloride ('isotopic tracer', 99%; Euriso-Top, Saint-Aubin Cedex, France). As algae-specific carbon sources, laminarin (Sigma-Aldrich, Munich, Germany; source: brown alga *Laminaria digitata*; composition: about 25 glucose units) and sodium alginate (Sigma-Aldrich; source: brown algae; composition: 61% mannuronic acid, 39% guluronic acid with 60–400 uronic acid units) were added to a final concentration of 0.2%. The control cultures were supplemented with glucose (Carl Roth GmbH & Co. KG, Karlsruhe, Germany) to a final concentration of 0.2%. All of these experiments were carried out in triplicates (Supplementary Figure S1A).

Proteome fractionation

Cellular protein extracts were separated into three different fractions: the intracellular soluble, the membrane-bound and the extracellular soluble proteome (Supplementary Figure S1B). For purification of extracellular proteins, cells were centrifuged (6600 g at 4 °C for 10 min) and the resulting supernatant was precipitated at 4 °C using 10% trichloroacetic acid, according to the protocol published by Antelmann *et al.* (2001). The precipitated proteins were pelleted via centrifugation (40 000 g at 4 °C for 60 min), washed with ethanol and solubilized in 8 M urea and 2 M thiourea. To extract the membrane-bound and intracellular protein fractions, the cells were disrupted in a French press (Simaminc SLM, Rochester, NY, USA; three cycles) followed by removal of cell debris by centrifugation (40 000 g at 4 °C for 20 min). Subsequent ultracentrifugation (100 000 g at 4 °C for 60 min) allowed for separation of intracellular soluble and membrane-bound proteins. Purification of the membrane fraction was performed, as described by Eymann *et al.* (2004), omitting the n-dodecyl- β -D-maltoside treatment. Accordingly, pelleted bacterial membranes were homogenized, washed for 30 min with a highly concentrated saline solution (20 mM Tris-HCl pH 7.5, 10 mM EDTA and 1 M NaCl) and ultracentrifuged (100 000 g at 4 °C for 60 min). Following this, the same steps were carried out using a carbonate buffer (100 mM Na_2CO_3 -HCl pH 11, 10 mM EDTA and 100 mM NaCl). Finally, purified membrane and intracellular proteins were solubilized in 50 mM triethylammonium bicarbonate buffer (Sigma-Aldrich) supplemented with the Complete Protease Inhibitor (Roche, Berlin, Germany). After determination of the protein concentration, triplicates of ^{15}N -labeled protein extracts from all three different subcellular fractions were combined and then mixed with equal amounts of individual ^{14}N -labeled protein samples in a 1:1 ratio for all substrates before proteome analysis.

1D-PAGE and protein measurements

From each ^{14}N -labeled triplicate combined with the ^{15}N -labeled pooled protein extracts, 15 μg of each fraction were separated by 1D-polyacrylamide gel electrophoresis (10% acrylamide, protein ladder by Thermo Fisher Scientific Inc., Waltham, MA, USA), stained with Coomassie G-250 'blue silver' (Candiano *et al.*, 2004), eluted from the gel and digested using trypsin. Peptides were subsequently desalted by ZipTip columns (Millipore, Billerica, MA, USA) according to the manufacturer's guidelines. For liquid chromatography-MS/MS measurements, peptides were subjected to a reversed-phase C18 column chromatography operated on a nanoACQUITY-UPLC (Waters Corporation, Milford, MA, USA) and separated as described by Otto *et al.* (2010). Mass spectrometry (MS) and MS/MS data were determined using an online-coupled LTQ-Orbitrap mass spectrometer (Thermo Fisher Scientific Inc.).

Data processing

The *.dta files were extracted from *.raw files with the BioworksBrowser 3.3.1 SP1 (Thermo Fisher Scientific Inc.) with no charge state deconvolution and deisotoping. The data were searched by SEQUEST version v28 (rev.12, Thermo Fisher Scientific Inc.) against a '*G. forsetii*' target-decoy protein sequence database. A set of common laboratory contaminants was included and all data were compiled using the BioworksBrowser. The following search parameters were used: enzyme type, trypsin (KR); peptide tolerance, 10 p.p.m.; tolerance for fragment ions, 1 a.m.u.; b- and y-ion series; variable modification, methionine (15.99 Da). A maximum of three modifications per peptide was allowed. In the second iteration, the mass shift of all amino acids completely labeled with ^{15}N -nitrogen was taken into account in the search parameters. Resulting *.dta and *.out files were assembled and filtered using DTASelect (version 2.0.25; parameters: -y 2 -c 2 -C 4 -here -decoy Reverse_ -p 2 -t 2 -u -MC 2 -i 0.3 -fp 0.005).

Relative protein quantification and analysis of variance

Relative quantification of the shotgun proteomics data was carried out according to the principle of pooled reference samples published by MacCoss *et al.* (2003) using the Census software (Park *et al.*, 2008) as described in Otto *et al.* (2010). Basically, according to the filtered database search peptide identification data provided by DtaSelect, both intensities for ^{14}N - (sample) and ^{15}N - (pooled reference) mass traces were extracted for each peptide identification in the respective mass spectrometric data and were compared within Census to yield a ratio between the sample mass trace (^{14}N) and the pooled reference mass trace (^{15}N). These peptide ratios were then combined to yield

sample/reference ratio values on the protein level. Finally, peptide ratios and combined protein ratios were exported (R^2 -values >0.7 and only unique peptides; proteins failing to be relatively quantified were checked manually in the graphical user interface for on/off proteins). Only proteins identified and quantified with at least two peptides were taken into account for further analyses. For correcting any potential systematic bias that might have been introduced by pooling of the ^{15}N -labeled samples and mixing with unlabeled protein samples, the quantitative data of a complete GeLC-MS run were median-centered. For all ratios, \log_2 -values and averages of the biological replicates were determined. TMEV (version 4.4; Saeed *et al.*, 2003) was applied to carry out the analysis of variance with the following settings: number of groups 5; P -values based on permutation (1000) resulting in P -values <0.01 as significance threshold. Quantification and analysis of variance data are provided in Supplementary Table S1.

Gene functions and protein localization

Pairwise identities of homologous genes were determined by means of the Geneious alignment software (Drummond *et al.*, 2010). Details on protein functions were obtained from precomputed data available on the UniProt Knowledgebase (Magrane, UniProt consortium 2011), the ENZYME database (Bairoch, 2000) and by using the JCoast software (Richter *et al.*, 2008). Information on the classification of CAZymes was derived from the CAZy database (Cantarel *et al.*, 2009). Subcellular localizations of all proteins were predicted according to Romine (2011); the data are provided in Supplementary Table S2.

Results and Discussion

Proteome analysis

In the present proteome analysis, 1691 proteins were identified, which corresponds to 47% of all protein-coding genes in '*G. forsetii*'. A considerable proportion (12%) of identified proteins could be assigned to the outer membrane, with one-third showing significant upregulation in alginate- or laminarin-grown cultures. Proteins enriched in the membrane fraction most frequently consisted of transporters (12%, mainly TBDRs), CAZymes (5%, primarily glycosyltransferases, GTs), signal-transducing components (3%), SusD-like proteins (2%) and proteins with unknown functions (35%).

A genome-wide *in silico* localization prediction indicated that the '*G. forsetii*' extracellular proteins are in most cases attached to its outer membrane (Supplementary Table S2). This was supported by broad overlaps of proteins detected in the extracellular and the membrane proteome (Supplementary Figures S2A and S3). The exposure

of proteins to the extracellular space is essential for bacteria in order to efficiently bind macromolecular substrates and decompose them into smaller components that are suitable for uptake into the cell. Conceivably, it would be an economical advantage for marine microorganisms to keep the majority of substrate-binding proteins and hydrolytic enzymes to themselves by attaching them to the cell envelope rather than secreting them into the surrounding water. Such a strategy would prevent a loss of these proteins by diffusion and increase the efficiency of breakdown/uptake coupling (see also Azam and Malfatti, 2007). Intriguingly, at the same time, such membrane-anchored enzyme systems would be a means of cell attachment to nutrient-rich particles like marine snow. Indeed, about 74% of all secreted proteins found in the extracellular fraction of 'G. forsetii' were anchored to the outer membrane and thus associated with the cell (Supplementary Figures S2 and S3, Supplementary Table S3).

The proteome analysis revealed that both laminarin and alginate notably affected the protein expression patterns of the membrane and extracellular fractions, whereas distinct quantitative changes of proteins from the intracellular fraction were observed in glucose-grown cultures (Supplementary Figure S2B). In total, a differential expression of 703 proteins (20% of all annotated genes) was observed. Seven outer membrane proteins showed a particular strong increase in abundance. These proteins are encoded in two large alginate- and laminarin-dependent PULs (Supplementary Figures S4A and B). The membrane fraction from glucose-grown cells was mainly dominated by inner membrane enzymes with functions in signal transduction, protein translocation, electron transport and ATP synthesis (Supplementary Figure S4C). Moreover, certain TBDRs were found to be highly abundant in all investigated cell samples.

Laminarin and alginate utilization loci

PUL-like systems are frequently found in members of the *Bacteroidetes* and typically comprise outer membrane Sus-like lipoproteins, such as SusC-like TonB-dependent transporters and SusD-like substrate-binding proteins, which function in concert with linked CAZymes, as described for members of the anaerobic genus *Bacteroides* of the human intestine (Martens *et al.*, 2009, 2011). The proteome analysis of this study demonstrated that in 'G. forsetii' two PULs are involved in alginate and laminarin degradation. These PULs comprise common components (TBDR, SusD, PKD domain proteins and CAZymes) and were found to be present in other marine *Flavobacteriaceae* as well (Figures 1 and 2).

The laminarin-dependent PUL of 'G. forsetii' encodes three 1,3-beta-glucanases, that is, two family-16 glycoside hydrolases (GH16 laminarinases) and one GH3 (Supplementary Table S4).

On the basis of their localization at the cell surface and according to information in the CAZy database (Cantarel *et al.*, 2009) both GH16 proteins (GFO_3466 and 3468) may constitute endo-acting enzymes that initiate polymer breakdown by randomly cleaving laminarin into smaller oligosaccharides. The latter can be taken up by a TonB-dependent transporter into the periplasmic space, where subsequent hydrolysis of terminal glucose residues by the presumably exo-acting GH3 (GFO_3467) may take place (Figure 3). Both, the hypothetical outer membrane protein (GFO_3465, Supplementary Table S7) encoded in this PUL, and one of the two GH16 laminarinases (GFO_3466) feature a PKD domain, known to participate in near-surface protein-protein interactions (Jing *et al.*, 2002). This suggests the existence of a physical link between the proteins, possibly connecting the substrate-binding SusD and the glycoside hydrolases.

The much larger alginate-dependent PUL encompasses four operons (Figure 2, Supplementary Tables S5). Upstream of the typical activity unit (operon 3 with TBDR, SusD, PKD and CAZymes) three polysaccharide lyase (PL) genes are located (operon 1 and 2 with PL6, PL7 and PL17). Downstream, an inner membrane transporter and genes for cytoplasmic processing of carbohydrates are encoded (operon 4 with genes for MFS permease, Sdr, KdgK and KdgA). On the basis of our expression analysis and consistent with transcriptome studies on the *Flavobacterium* species *Zobellia galactanivorans* (Thomas *et al.*, 2012), 'G. forsetii' gradually catabolizes alginate into 2-keto-3-deoxy-6-phosphogluconate (KDPG), which is likely directly funneled into the Entner-Doudoroff pathway (Preiss and Ashwell, 1962; Figure 3). A highly coordinated action of all alginate lyases (in operon 1, 2 and 3) and the cytoplasmic proteins 2-dehydro-3-deoxy-D-gluconate 6-dehydrogenase (Sdr), 2-dehydro-3-deoxygluconokinase (KdgK) and KDPG aldolase (KdgA; operon 4) is strongly supported by the proteome analysis, which revealed a consistent regulation pattern of these proteins within the alginate PUL (Supplementary Table S5).

The PKD-domain-containing proteins present in both, the alginate (GFO_1155) and the laminarin cluster (GFO_3465), most likely perform different functions as is indicated by their low similarity (51.4%) and striking differences in molecular weight (34.0 kDa for the alginate- and 96.6 kDa for the laminarin-induced proteins). PKD-domain-containing proteins and alginate lyases or laminarinases, respectively, were only upregulated in the two PULs described.

Carbohydrate transporters and associated SusD-related proteins

Efficient nutrient uptake is a critical step in the competition of marine bacterioplankton.

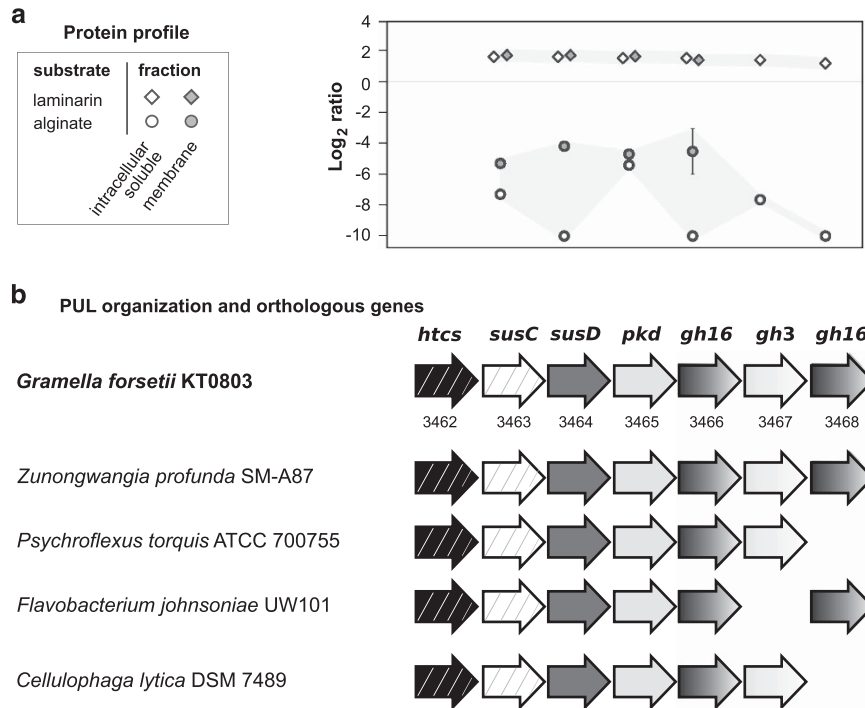


Figure 1 Expression profile (a) and gene organization of the laminarin utilization operon. (a) Protein ratios of the intracellular soluble and membrane protein fractions of cultures grown on laminarin compared with alginate are shown. Except for a signal transduction system (GFO_3462), all genes of the laminarin operon were overexpressed in the presence of laminarin. Consistency in protein levels is highlighted by the gray background that combines all data points of the corresponding substrate and indicates a uniform cluster regulation. Error bars refer to the s.d. of all biological replicates ($n = 3$), median-centered protein ratios are given in Supplementary Table S4. (b) The gene organization of the laminarin-specific PUL in 'G. forsetii' is conserved among other marine *Flavobacteriaceae* as exemplified. Identity/similarity values are listed in Supplementary Table S9. *htcs*, hybrid two-component system; *susC*, SusC-like TBDR; *susD*, SusD-family protein; *pkd*, PKD-domain-containing protein; *gh*, glycoside hydrolase.

In heterotrophic Gram-negative bacteria, TBDRs/transporters have a pivotal role in the uptake of complex organic matter such as from algal polysaccharides. This is especially reflected by a high number of TBDR-encoding genes in the genomes of marine *Flavobacteria*, compared with monomer specialists within the class *Alphaproteobacteria* (Fernández-Gómez *et al.*, 2013). The importance of TBDR in marine bacterioplankton is corroborated by *in situ* proteome studies of samples from the South Atlantic (in particular at coastal upwelling zones; Morris *et al.*, 2010), the Antarctic Southern Ocean (Williams *et al.*, 2012) and the coastal North Sea (Teeling *et al.*, 2012) where TBDRs dominated the set of expressed transporters. These transporters consist of an outer membrane TBDR that is linked to the cytoplasmic membrane via a TonB–ExbBD complex. The latter allows usage of the proton motive force across the cytoplasmic membrane for transport through the outer membrane by the selective opening of an outer membrane pore (Krewulak and Vogel 2011). Owing to their substrate-binding function and high sequence diversity, the TBDR component is suggested to mediate substrate specificity (Schauer *et al.*, 2008; Tang *et al.*, 2012).

Our proteome study identified 33 of the 40 'G. forsetii' TBDRs (83%) in the membrane fraction,

21 (53%) of which were differentially regulated under the three investigated substrate conditions (Figure 4). These findings suggest that some TBDRs are constitutively expressed and act as substrate sensors, whereas other TBDRs are dynamically upregulated only when these substrates become available. Laminarin exerted the most significant influence on TBDR expression (28% of TBDR genes responded to laminarin, 13% to alginate; Supplementary Figure S5). Among the laminarin-induced TBDRs we found (I) one encoded in the laminarin-specific PUL (GFO_3463), (II) one encoded next to a gene for a highly induced hypothetical inner membrane protein (Supplementary Table S7, GFO_2191) and (III) three TBDRs that were also identified in glucose-grown cultures (GFO_0361, 2138 and 1,4- α -glucanase associated; GFO_1029 next to two large outer membrane lipoproteins with adhesion domains), indicating regulatory overlaps between laminarin and its monomeric component glucose.

Half of all TBDR genes (20 of 40) were located next to genes encoding outer membrane SusD-like proteins. These proteins are unique to *Bacteroidetes* (Thomas *et al.*, 2011), and have been suggested to be involved in oligosaccharide binding as well as to interact with TBDRs (Shipmann *et al.*, 2000; Cho and Salyers, 2001). In addition to the 15 known

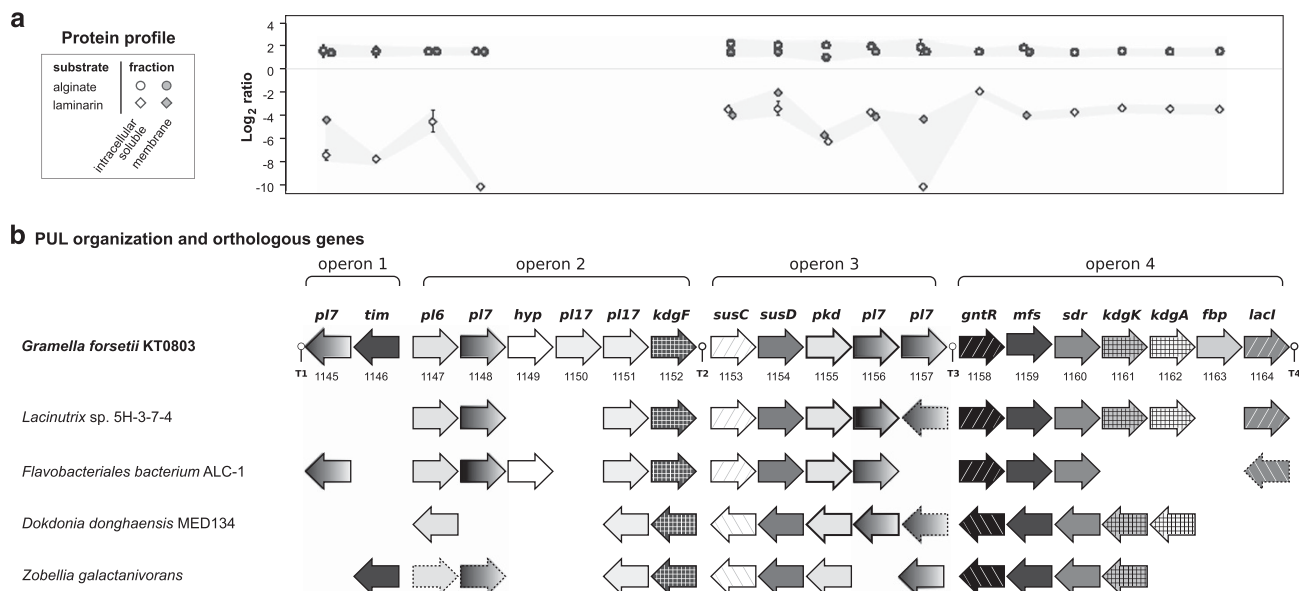


Figure 2 Differential expression (a) and genomic structure (b) of the alginate utilization cluster. (a) Shown are protein ratios of the intracellular soluble and membrane protein fractions of cultures grown on alginate compared with laminarin. The gray background marks the clustering of protein patterns in corresponding cultures. According to the CAZy database, GFO_1150 and 1151 are fragments of a single PL17 enzyme, which probably hindered its protein identification. Error bars indicate the s.d. of the biological replicates ($n = 3$), protein ratio values can be found in Supplementary Table S9. (b) The four putative operons of the alginate-specific PUL are flanked by thermostable stem-loop structures of possible transcriptional terminators (T, terminator; changes in Gibbs free energy: $\Delta G(T1) - 6.9 \text{ kcal mol}^{-1}$, $\Delta G(T2) - 20.0 \text{ kcal mol}^{-1}$, $\Delta G(T3) - 21.3 \text{ kcal mol}^{-1}$, $\Delta G(T4) - 8.1 \text{ kcal mol}^{-1}$). Homologous gene equivalents are present in related marine *Flavobacteriaceae* (identity/similarity values are listed in Supplementary Table S5). Dotted lines mark genes that belong to a separate operon, but were considered in this overview, because of their orthology to corresponding genes in 'G. forsetii'. *susC*, SusC-like TBDR; *susD*, SusD-family protein; *pkd*, PKD-domain-containing protein; *pl*, polysaccharide lyase; *tim*, TIM-barrel-domain-containing protein; *kdgF*, pectin degradation protein; *gntR*, GntR family transcriptional regulator; *mfs*, MFS permease; *sdr*, short-chain dehydrogenase/reductase family protein with a putative function as 2-dehydro-3-deoxy-gluconate-6-dehydrogenase; *kdgK*, 2-dehydro-3-deoxy-gluconokinase; *kdgA*, KDPG/KHG aldolase; *fbp*, fructose-1,6-bisphosphatase; *lacI*, LacI family transcriptional regulator; *hyp*, hypothetical protein.

SusD family genes (PF07980; Figure 4b upper section; Bauer *et al.*, 2006), this study revealed five additional genes with novel SusD-like domains (PF12771 and PF12741; Figure 4b, lower section) in the genome of 'G. forsetii'. Fourteen (70%) of these in total 20 *susD*-related genes were expressed and exhibited similar expression patterns as their colocalized TBDRs, but in comparison with the TBDRs a lower fraction of the *susD*-like genes were regulated (only 25% of *susD* genes were regulated, mainly laminarin-dependent with glucose overlaps). Alginate- or laminarin-induced upregulation of TBDR genes occurred irrespective of whether these were colocalized with *susD* genes (50%, Figure 4b) or not (45%, Figure 4a).

Tandems of *susC* and *susD* genes in 'G. forsetii' are frequently associated with CAZyme genes (45%; Bauer *et al.*, 2006). However, only few SusD-like proteins from GH-associated PULs were expressed under the tested conditions (44%, in contrast to 82% in non-GH-containing operons). Such GH-linked *susD*-like genes belonged to (I) the alginate and laminarin PUL (GFO_1154 and 3464) and were also associated with genes for (II) an alpha-amylase (GH13; SusD-like GFO_2139) and (III) a fructan-specific hydrolase (GH32; SusD-like GFO_0011). The GH32-associated *susD*-like gene belongs to an

operon (GFO_0008-11, Supplementary Figure S6, Supplementary Table S9) that comprises a glucose-specific TBDR gene (GFO_0010). Surprisingly, the GH13-associated *susD*-like gene is linked to an aforementioned laminarin-induced *susC*-like gene (GFO_2138), which is part of a 1,4-alpha-glucan utilizing PUL (GFO_2129-41; Figure 5, Supplementary Table S9).

Among the group of inner membrane transporters, differential expression was detected for three genes of the major facilitator superfamily (MFS) transporters (5 of 18 genes identified on protein level). On the basis of their annotation, the regulated secondary carriers are specific for hexuronate/hexarate (GFO_1159, upregulated in alginate), alpha-gluco-sides and galactose/glucose (GFO_2136 and 3326, both exclusively detected in glucose cultures). In contrast, high numbers of ATP-binding cassette (ABC) transporters (20 of 28 genes) were found in the proteome of all examined substrates, but exhibited relatively constant, and therefore substrate-independent, expression levels. Like other marine *Bacteroidetes*, ABC transporters of 'G. forsetii' almost entirely lack periplasmic solute-binding proteins that usually mediate substrate specificity (Bauer *et al.*, 2006). This stands in stark contrast to the expression of such

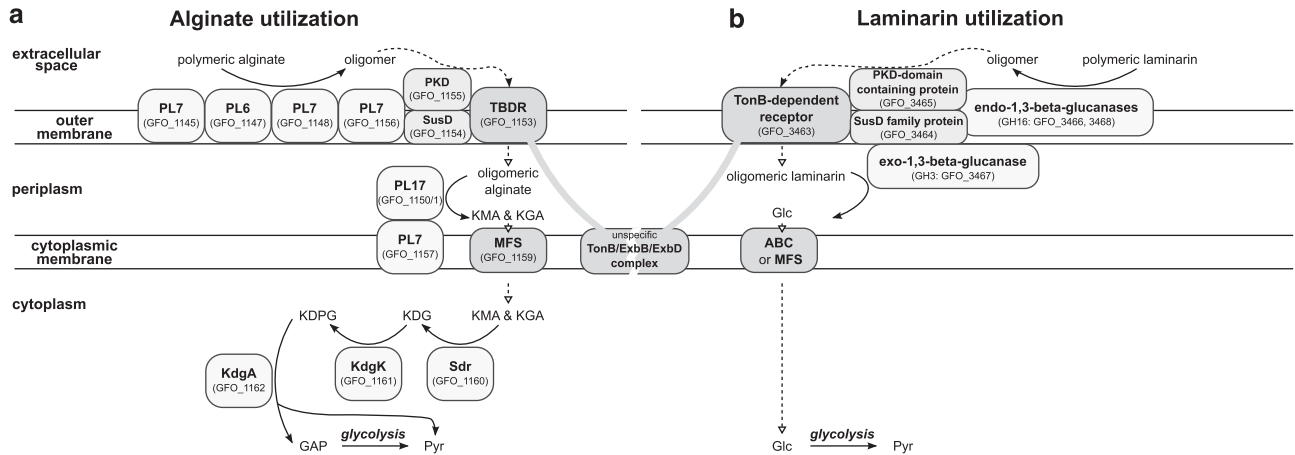


Figure 3 Tentative model of the alginate (a) and laminarin (b) utilization pathways in '*G. forsetii*'. Protein localizations were predicted *in silico* according to (Romine, 2011); Supplementary Table S2). (a) Four outer membrane alginate lyases of polysaccharide families 6 and 7 (PL6 and PL7, endo-acting according to ExPASy) probably catalyze the depolymerization of alginate into hetero- or homo-oligomeric alpha-L-guluronic acid (GA) and beta-D-mannuronic acid (MA). The PKD-domain-containing protein (PKD) and a SusD-related oligosaccharide-binding protein (SusD) may function as bridges delivering the oligomeric substrate to the TBDR. Transfer through the outer membrane is mediated by a linkage of the TBDR to the energizing TonB–ExbBD complex. According to their suggested localization, periplasmic alginate lyases (putative oligoalginate lyase PL17 and guluronate/ mannuronate lyase PL7) might exhibit oligoalginate specificity. This allows for breakdown into 4-deoxy-5-keto-mannuronic or guluronic acid (KMA and KGA). The monomers can be transported into the cytoplasm by means of an MFS permease and step by step converted into 2-keto-3-deoxygluconate (KDG; by a putative 2-dehydro-3-deoxy-gluconate-6-dehydrogenase, Sdr), keto-deoxy-phospho-gluconate (KDPG; by a 2-dehydro-3-deoxy-gluconokinase, KdgK) and pyruvate (Pyr) as well as glyceraldehyde-3-phosphate (GAP; by KDPG aldolase, KdgA). (b) The polysaccharide laminarin is metabolized into monomeric glucose by two outer membrane endo-1,3-beta-glucanases (GH16) and a periplasmic exo-1,3-beta-glucanase (GH3). Nutrient-binding and outer membrane transport are carried out by the same protein families as present in the alginate utilization pathway (PKD, SusD and TBDR). Uptake of glucose into the cytoplasm is conducted by a constitutively expressed MFS permease (MFS) or ABC-type transporter (ABC), followed by glycolysis resulting in the generation of pyruvate.

periplasmic-binding proteins in *Alphaproteobacteria*, which was recently found in a metaproteomic study of Antarctic surface waters (Williams *et al.*, 2012), and might indicate a housekeeping function of ABC transporters in '*G. forsetii*'.

CAZymes

GTs participate in the biosynthesis of glycans and glycoconjugates that mediate various functions such as surface adhesion, biofilm formation, signaling or energy and carbon storage (Taniguchi *et al.*, 2002). Genome analyses of several marine *Bacteroidetes* (Bauer *et al.*, 2006; Fernández-Gómez *et al.*, 2013) have shown high abundances of GT genes. Of the 60 genes predicted for '*G. forsetii*' (Bauer *et al.*, 2006), 39 were proteomically identified in this study, with a dominance of the families GT1 and GT2. Probably owing to their basic cellular function, expression of GTs was similar under all substrate conditions.

Many members of glycoside hydrolase families exhibited non-differential expression or were not identified (Supplementary Table S8). Interestingly, however, two collocated genes, one encoding a GH13 (amylase/cyclomaltodextrinase, GFO_2133) and one encoding a GH65 (maltose phosphorylase, GFO_2134) were significantly upregulated in glucose- and laminarin-grown cultures. Both the genes affiliated to a large alpha-1,4-D-glucan PUL (Supplementary Figure 5, Supplementary Table S9) together with adjacent glycolytic genes. The maltose

phosphorylase likely obtains its substrate maltose from the adjacent amylase/cyclomaltodextrinase that, owing to its dual function, may decompose both maltodextrins (1,4-linked D-glucose oligomers) and its precursors glycogen, starch and amylose (long-chained glucose polymers). Absence of two further cluster-encoded amylases (GFO_2132 and 2141) on the protein level suggests focusing on dextrins as parent substrates. Two additional components of this cluster, an alpha-glucoside-specific MFS permease and a beta-phosphoglucomutase, were identified in a single replicate of glucose-grown cells that provide the cytoplasmic glycolysis machinery with glucose-6-phosphate. '*G. forsetii*' notably regulated cell surface and intracellular utilization genes of the alpha-1,4-D-glucan PUL in a differential manner (Figure 5). The outer membrane proteins TBDR and SusD were selectively activated in the presence of polymeric laminarin, whereas intracellular tri-/di-/monosaccharide processing enzymes and a transporter were strongly induced by monomeric glucose.

Sensing of substrates

The majority of hybrid sensor/response regulator signal transduction systems was identified in the membrane or intracellular fraction (9 of 12), some of which responded specifically to laminarin (GFO_1684, 1697 and 2345), but this response was insignificant in comparison with glucose-grown

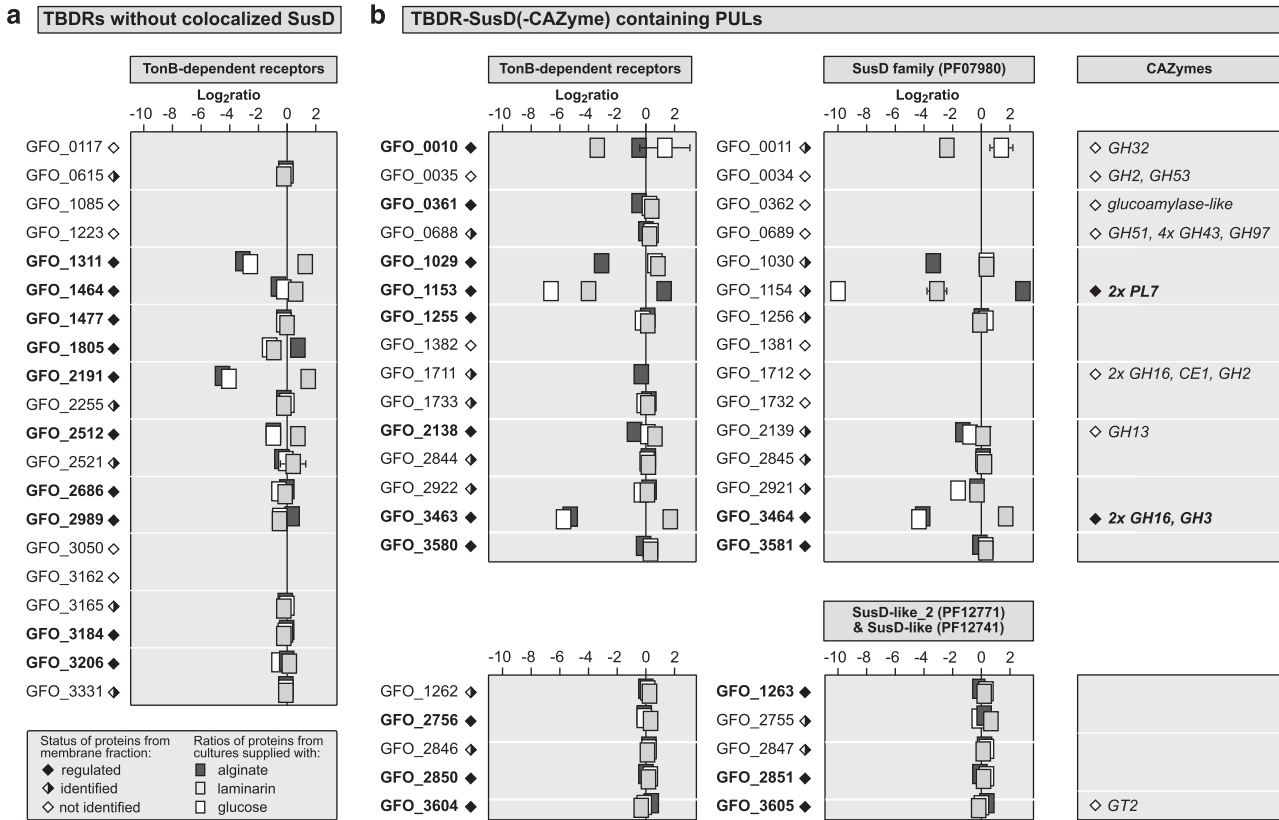


Figure 4 Regulation patterns of (a) TBDRs and (b) TBDR-SusD PULs frequently clustered with CAZymes. Alginate- and laminarin-dependent relative quantitative changes in protein amounts of SusD-independent and SusD-associated TBDRs as enriched in the membrane fraction. Protein profiles illustrate similarities in regulation patterns of TBDRs and corresponding SusD proteins. Colocalization of CAZymes was almost exclusively determined for operons containing SusD-related proteins with domain PF07980. Except for those associated with the major alginate or laminarin-specific PUL (in bold), all CAZyme genes in these operons remained unexpressed (in italics). Locus tags of 'G. forsetii' (GFO) proteins are indicated.

cells (Supplementary Table S9). Also, high numbers of sensors and regulators of conventional non-hybrid two-component systems were determined in the proteome (28 of 62 predicted signal transduction elements, 45%). However, except for a few glucose-induced response regulators (GFO_1017, 1420 and 1778), most of these proteins showed no differential expression. These findings, together with the omnipresence of various two-component signaling proteins under all substrate conditions, suggest a constitutive function, in which rather specificities than high quantities of signal transduction systems mediate the distinct upregulation of polysaccharide-associated genes. We observed a limited expression of ECF-sigma/anti-sigma factors (6 of 21 genes; 29%), which indicates that sensing of alginate, laminarin and glucose was not predominantly carried out via TonB-dependent transducers (TBDRs that are linked to a cytoplasmic sigma factor by inner membrane-localized anti-sigma factors; Koebnik, 2005). Instead, sensing was likely initiated by TBDR-enabled uptake of oligosaccharides into the periplasm in conjunction with (hybrid) two-component signal transduction to the cytoplasm. Only one sigma/anti-sigma system (GFO_1018-19) and an ECF-type sigma factor (GFO_3248) had an

elevated expression in laminarin-supplied cultures, however, these genes were not found adjacent to a TBDR gene.

Glycan-specificity and suggested functional assignments of proteins with unknown functions

Of all identified proteins, 27% were annotated as hypothetical ones. Of these hypothetical proteins, 70 displayed laminarin- or alginate-dependent expression (Supplementary Tables S6 and S7) and were predominantly detected in the membrane-bound fraction. Presence of the two polysaccharides resulted in increased expression levels of proteins that were likely involved in outer membrane assembly and maintenance of the cell envelope's integrity (alginate-dependent GFO_0835 and 1564; laminarin-dependent GFO_0244). Moreover, laminarin induced a marked upregulation of an imelysin-like protein with putative laminarin-binding function (GFO_2514) and its neighboring hypothetical protein (GFO_2515; Supplementary Figure S7). A variety of proteins with unknown functions also seemed to participate in TonB-dependent transport in laminarin-grown cultures (GFO_0462, 2190 and 2728). In addition a TonB-like hypothetical protein

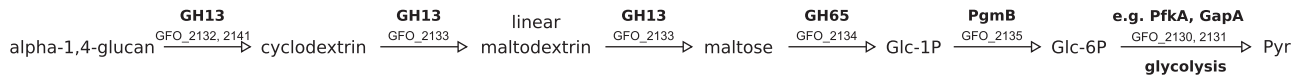
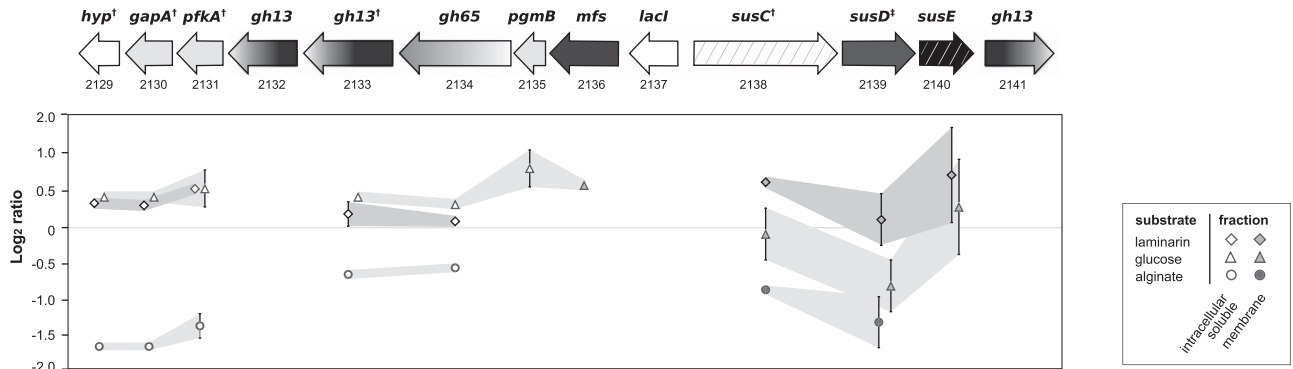
a Alpha-1,4-glucan utilization pathway**b Organization of gene cluster and protein profile**

Figure 5 Proposed pathway for the decomposition of alpha-1,4-glucans in '*G. forsetii*' (a), organization of the cluster and its protein profile (b). (a) An alpha-1,4-linked glucose-polymer (glycogen, starch and amylose) is converted into oligomeric dextrin (by GH13 alpha-amylase, GFO_2132, GFO_2141 or GFO_2133), dimeric maltose (by dextrinase function of GFO_2133), glucose 1-phosphate (Glc-1P; by GH65 maltose phosphorylase, GFO_2134) and finally glucose-6-phosphate (Glc-6P; by PgmB, GFO_2135). The latter metabolite can be directly used by glycolysis and metabolized into pyruvate (Pyr). (b) Intracellular components contained in the operon located upstream were upregulated in both glucose- and laminarin-supplied cultures (†, highest protein quantity changes in glucose-grown cultures), whereas membrane proteins of the downstream located operon exhibited upregulation particularly in laminarin-grown cultures (‡ $P < 0.01$). Except for the two alpha-amylases (GFO_2132 and GFO_2141) and the LacI family regulator, all genes were found to be expressed in this proteome study. According to the predicted subcellular localization, log₂ ratio values are shown for the respective subcellular fraction of each identified protein (error bars: s.d.; protein ratios: see Supplementary Table S9). Locus tags are given as numbers below the genes. *hyp*, hypothetical protein; *gapA*, glyceraldehyde-3-phosphate dehydrogenase A; *pfkA*, 6-phosphofructokinase; *gh*, glycoside hydrolase family; *pgmB*, beta-phosphoglucosyltransferase; *mfs*, MFS permease—possibly alpha-glucoside transporter; *lacI*, LacI family transcriptional regulator; *susC*, SusC-like TBDR; *susD*, SusD-family protein; *susE*, putative SusE-like protein.

(GFO_2900) showed increased expression in glucan-supplied cells. Its functional assignment is supported by a similar membrane expression pattern detected for the associated TBDR component ExbD (GFO_2902). Furthermore, a laminarin-dependent putative SprE-like gliding protein (GFO_3274) and an alginate-dependent hydrolase (GFO_0138) were upregulated in polysaccharide-supplied cultures.

Ecological implications

'*G. forsetii*' KT0803 is able to specifically respond to the presence of the algal polysaccharides laminarin and alginate. Its glycan utilization strategy comprises a distinct regulation of regulons with mostly cell surface proteins that allow for binding, uptake and degradation of complex sugars. The availability of either laminarin or alginate induced two PULs that share common characteristics (TBDR, SusD, PKD and CAZymes). Upregulation of the PKD-domain-containing proteins and laminarinases or alginate lyases, respectively, was restricted to these major PULs and thus highlights the importance of such clusters for polysaccharide decomposition. Moreover, the presence of homologs of PUL-encoded genes in related marine *Flavobacteriaceae* indicates a conserved glycan utilization process. To accommodate the varied composition and complexity of polysaccharides, PULs seem to be

organized as matching modules, in which genes can be rearranged, modified, added or removed in order to allow an evolutionary adaptation to the breakdown of specific substrates. Accordingly, utilization of the low-complexity substrate laminarin involves a smaller PUL compared with that of the more complex polysaccharide alginate.

'*G. forsetii*' lives in coastal surface waters and occupies an ecological niche that is defined by polysaccharide resources from algae. A respective adaptation is evident from the high number of *susC/susD*-like gene pairs and CAZymes in '*G. forsetii*' that notably exceed those found in the oligotrophic ocean-adapted *Flavobacteria*, *Polaribacter* spp. and *Dokdonia* spp. (Fernández-Gómez *et al.*, 2013). One-third of all TBDRs in '*G. forsetii*' were involved in the laminarin-specific response. In addition, partial co-regulation of TBDRs by both laminarin and its monomer glucose indicates that not only the glycosidic linkage type but also sugar composition controls PUL expression in *Bacteroidetes*. Likewise, crosstalk between the PULs specific for the beta-1,3-glucan laminarin and the alpha-1,4-glucans in '*G. forsetii*' demonstrates how the catabolism of two polysaccharides that are composed of the same monomer, but differ in linkage type can be efficiently co-regulated. A selected set of TBDRs and SusD-like proteins was found to be constitutively expressed under all tested substrate conditions. This suggests that TBDRs do not only act together with SusD-like proteins in binding

and uptake of specific substrates, but also have a general function in the early recognition of nutrient availability, probably via periplasmic signal transduction systems. Likewise, Thomas *et al.* (2012) suggested the presence of 'sentry' proteins waiting to recognize complex sugars and to trigger the alginate degradation pathway in the marine flavobacterium *Zobellia galactanivorans*. Moreover, a transcriptome study of the gut bacterium *Bacteroides plebeius* by Hehemann *et al.* (2012) showed a basal expression of selected CAZymes without an algal polysaccharide, but in the presence of its respective monomer. Such 'sensing' mechanisms may ensure a rapid response when algal polymers become available.

On overall, this proteogenomic study of the marine *Bacteroidetes* 'G. forsetii' allowed a detailed functional characterization of a series of enzymes, transporters and sensing proteins that concertedly participate in the consumption of typical algal polysaccharides. It can serve as a useful starting point in assessing substrate utilization capacities and metabolic versatilities of other *Bacteroidetes* representatives, such as members of the genera *Polaribacter* and *Formosa*, which have been identified as key organisms in the bacterioplankton during an algae bloom (Teeling *et al.*, 2012), or related bacteria in the human intestine (Martens *et al.*, 2011). Such functional analyses of model bacteria have the potential to establish protein biomarkers with well-characterized specificities that will be valuable for the interpretation of future metaproteomic *in situ* studies and thereby will ultimately deepen our understanding of the concerted polysaccharide breakdown by complex natural microbial communities.

Conflict of Interest

The authors declare no conflict of interest.

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