Carbohydrate-active enzymes (CAZymes) are an important feature of bacteria in productive marine systems such as continental shelves, where phytoplankton and macroalgae produce diverse polysaccharides. We herein describe Maribacter dokdonensis 62–1, a novel strain of this flavobacterial species, isolated from alginate-supplemented seawater collected at the Patagonian continental shelf. M. dokdonensis 62–1 harbors a diverse array of CAZymes in multiple polysaccharide utilization loci (PUL). Two PUL encoding polysaccharide lyases from families 6, 7, 12, and 17 allow substantial growth with alginate as sole carbon source, with simultaneous utilization of mannuronate and guluronate as demonstrated by HPLC. Furthermore, strain 62-1 harbors a mixed-feature PUL encoding both ulvan- and fucoidan-targeting CAZymes. Core-genome phylogeny and pangenome analysis revealed variable occurrence of these PUL in related Maribacter and Zobellia strains, indicating specialization to certain "polysaccharide niches.” Furthermore, lineage- and strain-specific genomic signatures for exopolysaccharide synthesis possibly mediate distinct strategies for surface attachment and host interaction. The wide detection of CAZyme homologs in algae-derived metagenomes suggests global occurrence in algal holobionts, supported by sharing multiple adaptive features with the hydrolytic model flavobacterium Zobellia galactanivorans. Comparison with Alteromonas sp. 76-1 isolated from the same seawater sample revealed that these co-occurring strains target similar polysaccharides but with different genomic repertoires, coincident with differing growth behavior on alginate that might mediate ecological specialization. Altogether, our study contributes to the perception of Maribacter as versatile flavobacterial polysaccharide degrader, with implications for biogeochemical cycles, niche specialization and bacteria-algae interactions in the oceans.

Keywords: alginate, fucoidan, ulvan, PUL, macroalgae, EPS, pangenome, Zobellia galactanivorans
INTRODUCTION

Continental shelves are productive marine systems, where photosynthesis by pelagic phytoplankton and benthic macroalgae yields considerable amounts of organic matter. Polysaccharides constitute a major fraction of the algae-derived organic matter, with important roles in nutrient cycles and microbial metabolism (Hehemann et al., 2014; Arnosti et al., 2021). Consequently, diverse bacteria are specialized for the degradation of algal polysaccharides, colonization of algal surfaces and other types of biological interactions (van der Loos et al., 2019; Ferrer-González et al., 2020; Wolter et al., 2021).

Cultured bacterial strains are a valuable resource for studying the ecological and biogeochemical implications of microbial polysaccharide degradation, complementing molecular and metagenomic approaches on community level (Arnosti et al., 2011; Wietz et al., 2015; Matos et al., 2016; Reintjes et al., 2017; Grieb et al., 2020). Culture-based studies revealed the diversity and functionality of carbohydrate-active enzymes (CAZymes), which encompass polysaccharide lyases (PL), glycoside hydrolases (GH), carbohydrate-binding modules (CBM), carbohydrate esterases (CE), glycosyl transferases (GT), and auxiliary carbohydrate-active oxidoreductases (Lombard et al., 2014). CAZyme-encoding genes are frequently transferred between microbes, providing effective mechanisms of adaptation and niche specialization (Hehemann et al., 2016).

Flavobacteria, including the families Flavobacteriaceae and Cryomorphaceae, are major contributors to marine polysaccharide degradation. Comparable to the human gut, marine flavobacteria can degrade various polysaccharides through dedicated genetic machineries (Teeling et al., 2012; Fernández-Gómez et al., 2013). Flavobacterial CAZymes are typically clustered with susCD genes in polysaccharide utilization loci (PUL) for orchestrated uptake and degradation (Grondin et al., 2017). For instance, the marine flavobacterium Zobellia galactanivorans has complex biochemical and regulatory mechanisms for degrading laminarin, alginate, agar, and carrageenan (Hehemann et al., 2012; Thomas et al., 2012, 2017; Labourel et al., 2014; Ficko-Blean et al., 2017; Zhu et al., 2017). Comparable abilities have been described in the flavobacterial genera Formosa agariphila and Gramella forsetii through diverse PUL (Mann et al., 2013; Kabisch et al., 2014; Reisky et al., 2019). Also Maribacter, the “sister genus” of Zobellia, exhibits hydrolytic activity (Bakunina et al., 2012; Zhan et al., 2017). Accordingly, both Maribacter and Zobellia are abundant on macroalgal surfaces (Martin et al., 2015), and related PUL have been detected during phytoplankton blooms in the North Sea (Kappelmann et al., 2019). Furthermore, several Maribacter and Zobellia strains stimulate algal development by producing morphogenesis factors (Matsuo et al., 2005; Weiss et al., 2017). Hence, both genera are important from ecological and biotechnological perspectives.

Here, we describe CAZyme content and hydrolytic capacities of Maribacter dokdonensis strain 62–1, isolated from an alginate-supplemented microcosm at the Patagonian continental shelf (Wietz et al., 2015). This highly productive marine region harbors frequent phytoplankton blooms and abundant coastal macroalgae, indicating regular availability of polysaccharides (Acha et al., 2004; García et al., 2008). Our CAZyme characterization in the pangenomic context illustrates the role of CAZymes, PUL and exopolysaccharide-related genes in niche specialization among Maribacter and Zobellia. The finding of diverse traits for interactions with algae, together with the detection of CAZyme homologs in macroalgae-derived metagenomes, highlight the predisposition of Maribacter spp. to algae-related niches and substrates. Notably, Maribacter dokdonensis 62-1 has been isolated from the same sample as Alteromonas sp. 76–1 with shown capacities for alginate and ulvan degradation (Koch et al., 2019b), illustrating that distantly related hydrolytic strains co-occur in the same habitat. Comparison of their CAZyme machineries illuminated whether these strains might employ different ecophysiological strategies or compete for resources. These eco-evolutionary perspectives into CAZyme diversity and corresponding niche specialization contribute to the understanding of ecophysiological adaptations behind community-level polysaccharide degradation (Teeling et al., 2012, 2016). Considering the abundance and biogeochemical relevance of algal polysaccharides, our study adds further evidence to the eco-evolutionary role of CAZymes in marine flavobacteria.

MATERIALS AND METHODS

Isolation and Cultivation

Strain 62–1 was isolated in April 2012 from a microcosm with surface seawater collected at the Patagonian continental shelf (47.944722 S, 61.923056 W) amended with 0.001% sodium alginate (Wietz et al., 2015). Purity was confirmed by PCR amplification of the 16S rRNA gene after several rounds of subculturing. Alginate utilization was analyzed in seawater minimal medium (SWM) (Zech et al., 2009) supplemented with 0.2% sodium alginate (cat. no. A2158; Sigma-Aldrich, St. Louis, MO) as sole carbon source in comparison to SWM + 0.4% glucose. Precultures were grown from single colonies for 24 h, washed three times with sterile SWM, and adjusted to an optical density of 0.1 measured at 600 nm (OD600). Main cultures were inoculated with 1% (v/v) of washed preculture in triplicate, followed by cultivation at 20°C and 100 rpm with regular photometric measurements (diluted if OD600 > 0.4).

Substrate Quantification

At each OD measurement, subsamples of 5 mL were filtered through 0.22 μm polycarbonate filters into combusted glass vials and stored at −20°C. Alginate concentrations were quantified by High Performance Liquid Chromatography (HPLC) of its monomers mannurionate and guluronate after chemical hydrolysis (20 h, 100°C, 0.1 M HCl) in combusted and sealed glass ampoules. Samples were neutralized with 6 N NaOH, desalted using DionexOnGuard II Ag/H cartridges (Thermo Fisher Scientific, Waltham, MA), and eluted with 100 mM sodium acetate tri-hydrate in 100 mM NaOH. Concentrations were determined in three dilutions per sample (0.01, 0.002, 0.001%) using a Carbowax PA 1 column (Thermo Fisher Scientific) and pulsed amperometric detection according to
Mopper et al. (1992). A calibration curve was generated using hydrolyzed 1% alginate solution ($R^2 = 0.97$). Glucose concentrations were measured using samples diluted to 0.001% with MilliQ followed by HPLC with NaOH (18 mM) as eluent and a Carboxapac PA 1 column (Thermo Fisher Scientific). A calibration curve was generated using 24 concentrations from 0.025 to 10 μM glucose ($R^2 = 0.99$).

**Genome Sequencing and Taxonomy**

Genomic DNA was extracted using the PeqGold DNA Isolation Kit (PEQLAB, Germany) according to the manufacturer's instructions. The genome was sequenced with Illumina technology using a GAIIx sequencing platform on paired-end libraries prepared with the Nextera XT DNA Kit (Illumina, San Diego, CA). A total of 83 contigs (0.5–330 kb, average 55 kb) were assembled using SPAdes v3.0 (Bankevich et al., 2012), followed by error correction using BayesHammer (Nikolenko et al., 2013) and gene prediction using the IMG pipeline (Markowitz et al., 2012). The draft genome has been converted to EMBL format and gene prediction using the IMG pipeline (Markowitz et al., 2012). The draft genome has been converted to EMBL format using EMBLmyGFF3 (Norling et al., 2018) and deposited at ENA under PRJEB40942. Phylogenetic analysis was carried out with 92 core genes identified using UBCG (Na et al., 2018), with Capnocytophaga ochracea DSM 7271 as outgroup. The resulting nucleotide alignment was visually confirmed for consistency and the best substitution model (GTR + G) computed using ModelTest-NG (Darriba et al., 2020). A maximum-likelihood phylogeny with 1000 bootstrap replicates was calculated using RaxML v8.2 (Stamatakis, 2014) on the CIPRES Science Gateway (Miller et al., 2010).

**Comparative Genomics**

Genomes of 62–1 and related strains (Supplementary Table 1) were compared using bioinformatic software. Average nucleotide identities were calculated using the Enveomics web application (Rodriguez-R and Konstantinidis, 2016). Core, accessory and unique genes were identified from protein-translated genes using OrthoFinder (Emms and Kelly, 2019) using a 30% identity cutoff. CAZymes were identified using dbCAN2 (Zhang et al., 2018), only considering hits with e-value $< 10^{-15}$ and > 65% query coverage. Gene annotations and PUL boundaries were manually curated based on the CAZy and UniprotKB-Swissprot databases (Lombard et al., 2014; Bateman et al., 2017). Sulfatases were identified using SulfAtlas v1.1 (Barbeyron et al., 2016a), only considering hits with e-value $< 10^{-1}$ and > 40% query coverage. Genes were assigned to KEGG classes and pathways using KAAS and KEGG Mapper (Moriya et al., 2007; Kanehisa and Sato, 2020). PUL homologies were analyzed by custom-BLAST in Geneious Pro v72 and PULDB (Terrapon et al., 2018). Genes for downstream processing of alginate monomers ($kdgA$, $kdgF$, $kdgK$, and $dehR$) were identified by searching homologs from Gramella forsetii (NCBI assembly GCA_000060345.1). Putative $kdul$ and $kduD$ genes for processing unsaturated uronates were identified by searching homologs from Gramella flava (NCBI assembly GCA_001951155.1). Signal peptides were predicted using SignalP v5.0 (Almagro Armenteros et al., 2019).

**RESULTS AND DISCUSSION**

Strain 62–1 was isolated from alginate-supplemented seawater collected at the Patagonian continental shelf (Wietz et al., 2015). Colonies on solid medium are round, smooth, and yellowish-colored. Genome sequencing resulted in a draft genome (83 contigs) with a cumulative length of 4.6 Mb, encoding 4,103 predicted proteins. Core genome-based phylogeny revealed clear assignment to Maribacter from the Flavobacteriaceae (Figure 1), with 99.8% 16S rRNA gene similarity and 97.8% average nucleotide identity to Maribacter dokdonensis DSW-8T (Supplementary Figure 1). Hence, strain 62–1 is a novel member of this flavobacterial species. M. dokdonensis DSW-8T originates from South Korean and hence subtropical waters, demonstrating occurrence of closely related strains on global scales.

**CAZymes in the Phylogenomic Context**

Maribacter dokdonensis strain 62–1 encodes 90 putative CAZymes predicted by dbCAN2, corresponding to 2% of all protein-encoding genes (Table 1 and Supplementary Table 1). As described in detail below, CAZymes commonly clustered with susCD genes, the hallmark of PUL in Bacteroidetes (Grondin et al., 2017). The presence of 12 polysaccharide lyases from PL families 6, 7, 12, and 17 illustrates specialization toward alginate, confirmed by physiological experiments (Figure 2). Most PL12 are classified as heparinases, but co-localization with known alginate lyase families indicates alginoletic activity. Strain 62–1 furthermore encodes PL33 and PL40 lyases that potentially target ulvan (Table 1). CAZyme numbers and diversity match the hydrolytic potential of related Maribacter and Zobellia strains included for comparison (Supplementary Table 1), corroborating the adaptation of these taxa to algal substrates and surfaces (Bakunina et al., 2012; Martin et al., 2015; Kwak et al., 2017; Zhan et al., 2017; Chernysheva et al., 2019).

We contextualized CAZyme patterns with phylogenetic relationships (Figure 1 and Supplementary Table 2) and a general overview of the Maribacter pangeneome (Supplementary Table 3). Core genome-based phylogeny resolved three lineages, each with distinct signatures of CAZymes and exopolysaccharide-related genes (Figure 1). Maribacter lineages 1 and 2 encode more PLs than lineage 3 (Wilcoxon rank-sum test, $p = 0.01$), including an entire additional PUL for alginate degradation. Moreover, one PL7 and one PL12 are unique to lineage 1 (see detailed paragraph below). In contrast, a sizeable PUL encoding a PL10 pectate lyase with CE8 methyltransferase domain is restricted to lineage 3 (Figure 1). This PL10-CE8

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1https://www.geneious.com

2https://rstudio.com
Alginic Acid Degradation Relates to Two Alginolytic PUL

The array of predicted alginic acid lyase genes allowed considerable growth with alginic acid as sole nutrient source (Figure 2). Growth was only slightly lower than with the monosaccharide glucose, indicating excellent adaptation for polysaccharide degradation. HPLC demonstrated that concentrations of both monomeric building blocks of alginic acid, mannuronic (M) and guluronic (G), decreased at the same rate (Figure 2).

Predicted alginic acid lyase genes are encoded in two major PUL, plus additional, single genes dispersed throughout the genome (Figure 3 and Supplementary Table 2). AlgPUL1 harbors four adjacent PL6-12-6-17 lyase genes, co-localized with a susCD pair and all genes for downstream processing of alginic monomers (Figure 3A). Hence, AlgPUL1 presumably encodes the complete metabolic cascade from external polymer breakdown (PL6, PL12), oligosaccharide hydrolysis (PL17) to polysaccharide (PL1).
monomer processing (kdgA, kdgF, kdgK, and dehR). The co-localization of PL families with structural and catalytic diversity (Xu et al., 2017) presumably facilitates access to various alginate architectures, e.g., relating to polymer length or the ratio between M and G in poly-G, poly-M or mixed stretches. Upstream of AlgPUL1 is a cluster encoding one GH3 and two GH144 genes, together with susCD and sulfatase genes (Figure 3). The two regions are separated by a type II-C CRISPR-Cas locus, signifying a mobile genomic region that might facilitate the exchange of adjacent CAZymes. Both GH144 genes share ~50% amino acid identity to an endo-glucanase of Chitinophaga pinensis (Abe et al., 2017), indicating glucosidase or endoglucanase activity. Their occurrence in all Maribacter and Zobellia (orthologous groups 0002512 and 0002513; Supplementary Table 3), many marine flavobacteria as determined by BLASTp (data not shown) as well as human gut microbes (McNulty et al., 2013) indicates ecological relevance across diverse habitats. The smaller AlgPUL2 encodes two PL7 lyases (Figure 3B) and might be auxiliary to AlgPUL1, potentially being activated by particular M/G architectures or environmental signals (Lim et al., 2011). The prediction of different signal peptide variants in the two PL7 (Supplementary Table 2) indicated that one PL7 is freely secreted whereas the other is anchored to the cell membrane. This complementary extracellular localization might boost alginolytic activity. Even without transcriptomic data, the combined physiological, chemical, and genomic evidence illustrates a functional and effective alginolytic pathway.

Comparative genomics revealed homologs of AlgPUL1 in most related Maribacter and Zobellia strains, whereas AlgPUL2 is restricted to Maribacter lineages 1 and 2. The observed structural
dissimilarity of alginateolytic PUL among *Maribacter* and *Zobellia* overall corresponds to core-genome relationships, supporting the notion of lineage-specific genomic signatures (Supplementary Figure 2). Notably, a PL12 only occurs in the AlgPUL1 variant of lineage 1 (Figure 1). This PL was presumably acquired from distant *Bacteroidetes* taxa, considering 65% amino acid identity to FNH22_29785 from *Fulvivirga* sp. M361 (Cytophagales). In contrast, lineages 2 and 3 encode a PL7 in this position, or solely harbor the PL6 and PL17 (Supplementary Figure 2). An almost identical AlgPUL2 occurs in *Fulvivirga eckloniae* (locus tags C1H87_08155-08120), with 65% amino acid identity between the respective PL7 lyases. The original isolation of *F. eckloniae* from the alginate-rich macroalga *Ecklonia* (Lee et al., 2017) highlights this PUL as adaptation to algae-related niches. In *Zobellia*, AlgPUL1, and AlgPUL2 occur in mixed combinations and in distant genomic regions (Thomas et al., 2012), suggesting internal recombination events (Supplementary Figure 2).

*Maribacter dokdonensis* 62–1 encodes a unique PL12 (locus tag 00457) within an exopolysaccharide-related cluster, including several unique GTs (Figure 3C). This cluster might represent a link between the degradation and biosynthesis of polysaccharides, considering that other bacteria regulate EPS production and release via PLs or GHs (Bakkevig et al., 2005; Köseoglu et al., 2015). The PL12 has 39% amino acid identity to ATE92_1054 of the North Sea isolate *Ulvisibacter* sp. MAR-2010-11 (Kappelmann et al., 2019), indicating ecological relevance of horizontally transferred PL12 homologs in distant habitats. The PL12 is among the two lyases without predicted signal peptide (Supplementary Table 2) and hence likely retained within the cell, supporting an intracellular role in EPS metabolism. However, only transcriptomic data can confirm whether the PL12 indeed targets alginate and might be co-regulated with EPS-related genes.

### A PUL Related to Ulvan and Fucoidan

Strain 62–1 harbors additional PUL for the degradation of other algal polysaccharides. The co-localization of two PL40 lyases, one PL33 from subfamily 2 as well as several sulfatases indicates activity towards a sulfated polysaccharide, presumably ulvan ([Figure 3D](#)). Although PL33 are classified as chondroitin or gellan lyases, our findings suggest an extended substrate range, potentially specific to subfamily 2. The two PL40 variants (locus tags 01347 and 01356) only have 30% identity and hence do not originate from duplication. Notably, PL40_01347 lacks a signal peptide compared to PL40_01356 and the PL33 (Supplementary Table 2), suggesting different secretory behavior and complementary functionality. The PL40_01347 homolog is conserved in most *Maribacter* and *Zobellia* genomes (Supplementary Figure 2), whereas PL40_01356 and the PL33 are missing in *Zobellia*. The latter PLs share 70 and 77% amino acid identity with a CAZyme pair in *Formosa agariphila* (locus tags BN863_10330 and BN863_10340, respectively), originally annotated as heparinase and PL8 but confirmed as PL33 and PL40 using the latest CAZy database (Mann et al., 2013; Lombard et al., 2014). Their absence in *Zobellia* but presence in distantly related *Arenibacter* spp. ([Figure 1](#)) indicates separate acquisition after speciation, potentially via transfer events between *Formosa* and *Maribacter*.

The combination of PL33 and PL40 differs from ulvanolytic PUL in other bacteria, which largely comprise ulvan lyases from PL families 24, 25, and 28 (Foran et al., 2017; Reisky et al., 2018; Koch et al., 2019b). Furthermore, the PUL in strain 62–1 also encodes α-fucosidases, including GH29, GH95, GH97, GH117, and GH141 comparable to a CAZyme plasmid in the verrucomicrobium *Lentimonas* (Sichert et al., 2020). Furthermore, a co-localized GH28 might remove galacturonic acid from fucoidan (Sichert et al., 2020), further metabolized by adjacent kduA and kduD genes into the Entner-Doudoroff pathway (Salinas and French, 2017). These observations indicate an alternative hydrolytic activity toward fucoidan, supported by fucose importer, fuconolactonase and fucanate dehydrogenase genes ([locus tags 01371, 01373, 01376, respectively](#)) encoded downstream of the PUL.

### Ecological Implications

To establish a broader ecological context, we searched homologs of the PL33–PL40 pair (specific to *Maribacter*) and the PL12 within the EPS gene cluster (unique to strain 62–1) in 102 microbial metagenomes from marine plants. The co-detection of PL33 and PL40 homologs on diverse algae and seagrasses from global locations ([**Figure 4A**](#) and [Supplementary Table 4](#)) illustrates that relatives of strain 62–1 occur in such niches worldwide. The presence of PL33–PL40 homologs on brown, green and red macroalgae indicates that predicted hydrolytic capacities are functional *in situ* and support establishment in algal holobionts. The brown macroalgae *Macroystis* and *Ecklonia* are rich in alginate whereas ulvan can constitute ~40% of the green algae *Ulva* (Kidgell et al., 2019), highlighting the importance of related CAZymes for associated bacteria and why *Maribacter* spp. are common algal epibionts (Martin et al., 2015).

Wide detection of the PL12 unique to strain 62–1 ([Supplementary Table 4](#)) supports the presumed role of the lyase and adjacent EPS genes in surface attachment. This gene arrangement might permit the formation of specific biofilm structures, helping to reduce competition with co-existing bacteria. EPS on algal surfaces could also provide a protective matrix, minimizing diffusion of secreted CAZymes and retaining hydrolysis products for maximal uptake (Vetter et al., 1998). In turn, EPS could also be advantageous in pelagic waters from which 62–1 has been isolated, where aggregation on self-produced EPS could constitute a protective refugium and facilitate survival when algal substrates are unavailable (Decho and Gutierrez, 2017). In general, the diversity of EPS genes on fine phylogenetic levels might mediate distinct interactions with algal hosts, considering that EPS can be strain-specific determinants of host interaction (Lee et al., 2016; Deo et al., 2019).

In addition to metagenomic analyses, we searched strain 62–1 for genes encoding characterized algae-adaptive traits in *Zobellia galactanivorans*, a closely related hydrolytic flavobacterium (Barbeyron et al., 2016b). This approach identified homologous genes for degradation of mannitol and digeneaside in strain 62–1 ([**Figure 4B**](#) and [Supplementary Table 4](#)). Furthermore, we found
GH43-GH10 and GH13-GH65 gene pairs targeting the algal carbohydrates xylan and maltose (Figure 4C). We also detected most homologs for anhydrogalactose utilization and hence the second step in carrageenan metabolism, but no GH127 or GH129 anhydrogalactosidase genes for initial hydrolysis (Ficko-Blean et al., 2017). Hence, Maribacter might utilize hydrolysis products from primary degraders, employing a secondary “harvester” strategy for carrageenan on red macroalgae (Hehemann et al., 2016). However, agarase or porphyrinase homologs were not detected (Figure 4B), indicating an overall narrower niche range than Zobellia.

In addition to carbohydrate utilization, we found other features typical for interactions with macroalgae. For instance, genes for the detoxification of iodine and nitrous oxide (Figure 4B and Supplementary Table 4) likely counteract algal defense mechanisms (Ogawa et al., 1995; Verhaeghe et al., 2008). Detoxification of superoxide and hydrogen peroxide is associated with two dismutases and 21 thioredoxin-like oxidoreductases (Supplementary Table 4), an almost twofold higher count despite an ∼1 Mb smaller genome than Zobellia (Barbeyron et al., 2016b). The predicted ability to detoxify methanol is likely advantageous during demethylation of pectinous algal substrates (Koch et al., 2019a). Moreover, strain 62–1 encodes traits that might be advantageous to algal hosts. For instance, a putative homoserine lactone lyase (Figure 4B and Supplementary Table 4) might interfere with communication of bacterial competitors and prevent their biofilm formation. This process, often termed quorum quenching, might antagonize resource competitors and modulate the composition of the holobiont (Wahl et al., 2012). Presence of a PKS might allow biosynthesis of thallusin, an essential algal morphogen produced by both Zobellia and Maribacter (Matsuo et al., 2003; Weiss et al., 2017). Another conserved feature is the complete pathway for biosynthesis of biotin (vitamin B7; locus tags 03353–03359 in strain 62–1), a common characteristic of macroalgal epibionts (Karimi et al., 2020). Strain 62–1 misses a single gene in biosynthetic pathways.
Comparison With Alteromonas sp. 76–1 From the Same Habitat

The gammaproteobacterial strain Alteromonas sp. 76–1, isolated from the same seawater sample as Maribacter dokdonensis 62–1, has a comparable predisposition toward polysaccharide degradation (Koch et al., 2019b). The presence of alginolytic and ulvanolytic systems in both 62–1 and 76–1 supports the notion that alginate and ulvan commonly occur at the Patagonian continental shelf. Although the two strains might hence compete for these resources, their corresponding CAZyme repertoires substantially differ. While sharing similar numbers of PL6 and PL7 alginate lyase genes (Figure 5A), PL12 are restricted to Maribacter and PL18 to Alteromonas, indicating different alginolytic strategies. Furthermore, concerning ulvanolytic activity, Alteromonas encodes PL24 and PL25 lyases on a CAZyme plasmid opposed to PL33 and PL40 on the Maribacter chromosome (Figure 5A), potentially influencing the regulation and transfer of these genes.

We hypothesize that differences in CAZymes and PUL for the same substrates mediate ecological specialization. For instance, different PL combinations might specialize for different subsets of these polysaccharides, e.g., relating to polymer length, the ratio of individual carbohydrate monomers, sulfatation or the degree of side-chain decorations. For instance, the Alteromonas-specific PL18 might boost alginolytic activity by acting as bifunctional lyase on both poly-G and poly-M stretches (Li et al., 2011). Furthermore, strain-specific GH genes might allow the separation into distinct “polysaccharide niches” and minimize competition. For instance, a GH109 with presumed N-acetylgalactosaminidase activity (Lombard et al., 2014) only occurs in Maribacter (Figure 5A). Finally, ecological specialization might include “temporal niche differentiation” considering variable doubling times when degrading alginate (Figure 5B). The delayed exponential phase in Maribacter suggests K-strategy, compared to r-strategy in the faster-growing Alteromonas (Pedler et al., 2014).

CONCLUSION

The diversity of CAZymes and PUL targeting alginate, ulvan and other algal carbohydrates illustrates a marked predisposition of Maribacter dokdonensis 62–1 for polysaccharide degradation, exemplified by distinct growth with alginate as sole carbon source. The variety of EPS-related genes might facilitate switching between free-living and surface-associated lifestyles, potentially connected to the specific colonization of algal hosts. The overall conservation of adaptive features in Maribacter and the “sister genus” Zobellia highlights their importance as macroalgal epibionts, supported by detection of PL homologs in metagenomes from brown, green and red macroalgae on global scales. The presence of both adverse (utilization of algal substrates, evading algal defense) and advantageous traits (vitamin production, biofilm control) indicates versatile lifestyles, which potentially change depending on physicochemical conditions and algal health (Kumar et al., 2016). Different hydrolytic machineries and doubling times of strain 62–1 compared to the co-occurring Alteromonas sp. 76–1 might avoid competition and result in colonization of discrete niches. These insights contribute to the understanding...
of flavobacterial CAZymes and hydrolytic activity in context of biogeochemical cycles, niche specialization and ecological interactions in the oceans.

DATA AVAILABILITY STATEMENT

The datasets presented in this study can be found in online repositories. The names of the repository/repositories and accession number(s) can be found in the article/Supplementary Material.

AUTHOR CONTRIBUTIONS

LW conducted genomic analyses and contributed to writing. MM performed growth experiments and HPLC. JK contributed to genomic analyses. RD performed genome sequencing. MS co-performed growth experiments and HPLC. LW conducted genomic analyses and contributed to writing. MM contributed to the final version of the manuscript.

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SUPPLEMENTARY MATERIAL

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