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Running title: In-depth characterization of marine alginate lyases **Key Words**: alginate lyase, family PL7, crystal structure, enzymology

Capsule

Background: Alginolytic systems from marine bacteria are crucial for algal biomass conversion, yet their molecular mechanisms remain poorly understood.

Results: Structural and biochemical characterization of two paralogous marine alginate lyases highlights details on complementary roles and differences with terrestrial enzymes.

Conclusion: Bacterial alginolytic enzymes are specifically adapted to the unique characteristics of the natural substrate.

Significance: Marine microbes evolved complex degradation systems targeting habitat-specific polysaccharides.

Abstract

Cell walls of brown algae are complex supramolecular assemblies containing various original, sulfated and carboxylated polysaccharides. Among these, the major marine polysaccharide component, alginate, represents an important biomass that is successfully turned over by heterotrophic marine bacteria. In the marine flavobacterium Zobellia galactanivorans the catabolism and uptake of alginate is encoded by operon structures that resemble the polysaccharide typical **Bacteroidetes** utilization locus (PUL). The genome of Z. galactanivorans contains seven putative alginate lyase genes, five of which are localized within two clusters comprising additional carbohydraterelated genes. This study reports the detailed biochemical and structural characterization of two of these. We demonstrate here that AlyA1_{PL7} is an endolytic guluronate lyase, while AlyA5 unsaturated cleaves units. α-Lguluronate (G) or β -D-manuronate (M) residues at the non-reducing end of oligo-alginates in an exolytic fashion. Despite a common jelly-roll fold, these striking differences of mode of action are explained by a distinct active site topology: an open cleft in AlyA1_{PL7}, whereas AlyA5 displays a pocket topology due to the presence of additional loops partially obstructing the catalytic groove. Finally, in contrast to PL7 alginate lyases from terrestrial enzymes bacteria. both proceed according calcium-dependent a mechanism suggesting an exquisite adaptation to their natural substrate in the context of brown algal cell walls.

Introduction

Brown algae dominate the primary production in temperate and polar rocky shores and represent a huge marine biomass. Indeed, coastal regions are considered as carbon sinks, retaining about 200.10^{12} g C per year (1). Their cell walls (CW) comprise a minor fraction of crystalline cellulose and a majority of anionic polysaccharides, alginates and sulfated fucoidans (2). Phlorotannins, which consist of halogenated and / or sulfated phenolic compounds (3,4) and 5% of proteins (5) complete this complex supramolecular assemblage. Among these compounds, alginate can account for up to 40% of the dry weight of the algal biomass (6). This linear polysaccharide is composed of β -D-mannuronate (M) and its C5 epimer α -L-guluronate (G), that are arranged in three types of repeating structures: poly-G stretches, poly-M stretches, and heteropolymeric random sequences (poly-MG) (7). The gelling properties strongly depend on the poly-G content of the algal polysaccharide, since these G-blocks form highly viscous solutions and gels through interconnected metal ion chelation (mainly Ca^{2+}) (8). Within the algal cell wall, the functional properties of alginate are modulated through changes in G- and M- content. These variations can be species specific or depend on season and environmental conditions; however, to date only very few studies exploring such relationships are available (9-12). Phylogenomic analysis of the carbohydrate metabolism of the model brown alga *Ectocarpus siliculosus* has revealed that brown algae horizontally acquired the alginate biosynthetic pathway from an ancestral Actinobacterium (13). Indeed, alginates are also produced as exopolysaccharides by some bacteria, such those belonging to the genera as Azotobacter and Pseudomonas (14.15). The main differences at the molecular level between algal and bacterial alginates are the presence of O-acetyl groups at C2 and/or C3 in the bacterial alginates (7,16) and their higher proportion of M units. From an ancestral bacterial exopolymer, alginate further evolved in brown algae into a pivotal CW polysaccharide, which is constrained by the necessity to interact with other CW components (2,13). These evolutionary constraints likely explain the loss of acetyl groups in algal alginates together with the increased importance of the G units and of their interactions with calcium ions to control gel properties. This hypothesis is consistent with the expansion of the mannuronan C5-epimerase gene families observed in the brown algae digitata (17-19) and Laminaria Ε. siliculosus (13).

Alginate constitutes an abundant nutrient resource for heterotrophic marine bacteria and thus plays in coastal ecosystems an ecological role similar to that of cellulosic and hemicellulosic biomass in terrestrial environments. However the comprehension of the molecular bases for the assimilation of algal alginates by marine bacteria remains at best fragmentary. We have recently reported the first operons specific for alginate assimilation in a marine bacterium, Zobellia galactanivorans (20). This flavobacterium, which is well known to degrade sulfated galactans from red seaweeds (21-23), is also able to use alginates from brown algae as sole carbon source (24). The alginolytic system of Z. galactanivorans encompasses seven

alginate lyases (AlyA1-AlyA7) belonging to four distinct families of polysaccharide lyases (PL, (25)), families PL 6, 7, 14 and 17. While *alyA1* and *alyA7* are isolated genes, alyA4, alyA5 and alyA6 are transcribed on the same mRNA. alyA2 and alvA3 are localized in a large operon carbohydrate-related comprising other genes, notably a TonB-dependent receptor and its associated SusD-like protein likely involved in oligo-alginate uptake (20). Such gene organization is typically found in Bacteroidetes and is referred to as a polysaccharide utilization locus (PUL) (26,27). With the exception of *alyA7*, all these genes are up-regulated in the presence of alginate (20). The complexity of this degradation system questions the exact role of the different alginate lyases. Notably, is there a functional redundancy providing a robustness to this alginolytic system? Do these enzymes display distinct substrate specificities? Do they proceed by different modes of action which could have synergistic effects? То discriminate between these hypotheses, which are not necessarily mutually exclusive, the indepth biochemical study of these enzymes is a prerequisite.

AlyA1 is a secreted modular protein with a N-terminal Carbohydrate Binding Module of the family 32 (CBM32) appended to a C-terminal PL7 module, while AlyA5 is predicted to be an outer membrane lipoprotein consisting of a lone PL7 catalytic domain. Moreover, these PL7 modules are extremely divergent with only 16% sequence identity and several large insertions are present in AlyA5 in comparison to AlyA1 (20). We describe here the detailed study of structurefunction relationships of the catalytic modules of two paralogous alginate lyases AlyA1 and AlyA5 from Z. galactanivorans, by a combination of crystallographic and biochemical approaches, including reaction product analysis by NMR spectroscopy.

Material and methods

Substrate materials

Sodium alginate samples with three different M/G ratios (0.5, 0.9 and 2.0) were provided by Danisco. Fractions of oligoguluronates, oligomannuronates and mixed MG oligosaccharides were prepared according to Haug *et al.* (9). This method has been reported to yield fragments with DP ranging from 4 to approx. 30 (28). Their respective M/G composition was controlled by NMR (data not shown). *Reduction of oligoguluronates*

Oligosaccharides were reduced following a protocol adapted from Abdel-Akher et al. (29). Briefly, 3 mg of sodium borohydride were added to 500 µl of oligoguluronates (1 mg.ml⁻¹) and the mixture was incubated overnight at room temperature. The reaction was acidified by adding drops of acetic acid until no more H₂ production was observed. The sample was then dried under vacuum, resuspended in 1 drop of 17.5 M acetic acid and 1 ml of methanol. Solvent was evaporated under a nitrogen flux. The amount of reducing ends was estimated by the ferricyanide assay (30) using a calibration curve with 20 -400 µg.ml⁻¹ glucose. Before the treatment, the ferricyanide assay measured 29.4 μ g.ml⁻¹ eq. glucose in the oligoguluronate solution. No reducing ends could be detected by the same assay after the reduction.

Phylogenetic analyses

Members of the PL7 family were selected CAZY database in the [http://www.cazy.org/(31)] and their sequences aligned using MAFFT with the L-INS-i algorithm and the scoring matrix Blosum62 (32). The alignment was manually edited in MEGA 4.1 (33). 180 sites were used to derive a phylogenetic tree using a maximum likelihood method conducted with the PhyML program (34) implemented online on Phylogeny.fr (35). Bootstrap values were calculated from 100 resamplings of the dataset.

Cloning of two genes coding for PL7 alginate lyases

Primers were designed to amplify the coding region corresponding to the mature protein for AlyA5 and for the catalytic module of AlyA1 only, hereafter called AlyA 1_{PL7} . The genes were amplified by PCR from Z. galactanivorans genomic DNA using the oligonucleotide primers as follows forward gggggGGATCCtgtaaagacaaacctaaggccact and reverse acg cccccGAATTCtcactgggcttctggggcgcttc and AlyA5, forward for ggggggAGATCTtccggtggttcgtccacccttc and reverse cccccGAATTCttaattgtgggttacgcttaggtttttg for the catalytic domain of AlyA1_{PL7}. Using the restriction sites EcoRI and BamHI for AlyA5, and EcoRI and BglII for AlyA1_{PL7}, the PCR products were then ligated into the expression vector pFO4, resulting in a recombinant protein with an N-terminal hexa-histidine tag. The plasmids were transformed into E.coli BL21(DE3) expression strains. All cloning procedures were performed as described in Groisillier et al., (36).

Protein expression and purification procedures

For both enzymes the same procedure was applied. Two milliliters pre-cultures of the recombinant E. coli BL21(DE3) cells were grown overnight at 37°C in LB medium containing ampicillin (100 µg.ml⁻ ¹). 200 ml of autoinducible ZYP medium with ampicillin (37) were inoculated with 200 µl of pre-culture and incubated for three days at 20°C and 200 rpm. Cells were centrifuged for 20 min at 5000 rpm, 4°C and pellets were stored at -20°C. Cells were re-suspended in 20 ml of Buffer A (25 mM Tris-HCl pH 7.5, 200 mM NaCl, 5 mM imidazole) containing a cocktail of antiproteases (Complete EDTA-free, Roche) and DNase. Cells were lysed with a French press. Samples were then centrifuged for 2 h at 20000 g. The supernatant was loaded on a Hyper Cell PAL column charged with 0.1 M NiSO₄ and equilibrated with Buffer A. Proteins were eluted with a linear gradient between Buffer A and Buffer B (25 mM Tris-HCl pH 7.5, 200 mM NaCl, 1 M imidazole) in 60 ml at a flow rate of 1 ml.min⁻¹ and collected in fractions of 1 ml. Fractions showing the presence of the recombinant proteins by sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) were pooled. Aliquots of 5 ml were further purified on a size exclusion chromatography column (Superdex 75 16/60 pg, GE Healthcare) using Buffer C (25 mM Tris-HCl pH 7.5, 200 mM NaCl) at a flow rate of 1 ml.min⁻¹. Fractions showing the presence of the pure proteins were pooled for AlyA1_{PL7} (data not shown), while two samples, separating the fractions of monomeric and dimeric forms of AlyA5 (data not shown), where prepared. Dynamic Light Scattering (DLS) measurements were performed on 40 µl of concentrated protein solutions (> 1mg.ml^{-1}) in a quartz cuvette on a ZetaSizer Nano-S instrument (Malverne Instruments).

Enzymatic assays

Protein concentration was determined by measuring the absorbance at 280 nm. Theoretical molar extinction coefficients ε_{280} were computed from the protein sequences on the ProtParam server (http://www.expasy.ch/tools/protparam.ht ml). Alginate lyase activity was assayed by measuring the increase in absorbance at 235 nm (A_{235}) of the reaction products (unsaturated uronates) for 5 min in a 1 cm quartz cuvette containing 0.5 ml of reaction mixture in a thermostated spectrophotometer. One unit of activity was defined as an increase of one unit in A₂₃₅ per minute.

The effect of temperature, NaCl and chelating agents on AlyA1_{PL7} activity were tested in 100 mM Tris-HCl buffer pH 7.5 using sodium alginate 0.05% as substrate and 5 μ l of purified enzyme at 1.2 μ M. The chelating agents EDTA or EGTA were either incubated at 1 mM final

concentration with AlyA1_{PL7} (overnight, 4°C) or added directly in the reaction mixture. The effect of pH on AlyA1_{PL7} activity was tested using various buffers at 100 mM: citrate (pH 3.0 to 6.0), 2morpholinoethanesulfonic acid-NaOH (MOPS, pH 6.0 to 7.9), HEPES (pH 7.0 to 8.0), Tris-HCl (pH 7.0 to 8.5) and glycine-NaOH (pH 8.5 to 10.0). Kinetic parameters were determined at 30°C in 100 mM Tris-HCl pH 7.5, 200 mM NaCl with 10 alginate concentrations ranging from 0.075 to 2.5 mg.ml⁻¹. Substrate concentrations were derived from the molecular mass of a M or G monosaccharide (194 g.mol^{-1}) . Reactions were performed in triplicates. A molar extinction coefficient ε =8500 L.mol⁻ ¹.cm⁻¹ (38) was used for the reaction products (unsaturated uronic acids) The kinetic parameters were estimated by a hyperbolic regression algorithm with Hyper32

(http://homepage.ntlworld.com/john.easterby/s oftware.html).

AlyA5 enzymatic assays on alginate polysaccharides were conducted as for AlyA1_{PL7}, using 5 μ l of the purified enzyme at 2.7 μ M. Reactions on saturated or unsaturated oligosaccharides were performed at 30°C in 100 mM Tris-HCl buffer pH 7.5, 4 mM CaCl₂, 50 μ g of substrate and 5 μ l of enzyme at 5.6 μ M. *Carbohydrate electrophoresis (C-PAGE)*

After enzymatic degradation, aliquots were boiled for 10 min before analysis. Oligosaccharides were analyzed by carbohydrate polyacrylamide gel electrophoresis (C-PAGE) (Zablackis and Perez 1990; Jackson 1996). Aliquots (5 µl) were mixed with 15 µl of loading buffer (10% sucrose and 0.01% phenol red). 5 µl were electrophoresed through a 6% (wt/vol) stacking and a 28% running, 0.75mm thick, polyacrylamide gel in 50 mM Tris-HCl, 1 mM EDTA buffer (pH 8.7). Gels were stained with alcian blue 0.5% (wt/vol) followed by silver nitrate 0.4% (wt/vol) (Cowman et al., 1984; Zablackis and Perez 1990). The coloration was developed with 7% sodium carbonate (wt/vol) containing formaldehyde and the reaction was stopped by adding concentrated acetic acid.

Monitoring of alginate lyases reaction by ¹H NMR

The action of alginate lyases was directly followed at 30°C in the NMR tube on crude alginates, G-, M- and alternating MG-blocks. The substrates were dissolved at a concentration of 5 mg.ml⁻¹ in a 100 mM deuterated phosphate buffer, pH 7.6, with 200 mM NaCl and 5 mM CaCl₂. The NMR experiments were performed on a Bruker DRX-400 MHz spectrometer using a 5.0 mm ${}^{1}\text{H}/{}^{13}\text{C}/{}^{15}\text{N}/{}^{31}\text{P}$ ONP probe or on 600 III а Bruker Avance MHz spectrometer using 5.0mm а $^{1}H/^{13}C/^{15}N/^{31}P$ inverse detection OXI probe, and a 2.5 mm ¹H/¹³C inverse detection probe, all equipped with zgradients and controlled by Topspin 1.3 and 2.1 software, respectively. The 1 H NMR experiments were recorded at 30 and 70 °C using a spectral width of 3000 Hz, an acquisition time of 5.62 s, a pulse width of 8.5 µs, a relaxation delay of 4 s, and 128 scans. The residual HOD signal was suppressed using the NOESY presaturation pulse sequence in which applied during saturation was the relaxation delay of 4 s and the mixing time of 50 ms.

Prior to an enzymatic reaction, a reference spectrum of the substrate was acquired. The sample was then removed from the probe, the enzyme was added at the proper concentration and the tube was put back into the probe. The reaction was monitored as a function of time. To monitor the progress of degradation of the sample, series of ¹H NMR spectra were recorded at 30 °C using the multizg or multi-zgvd programs from the Bruker library.

Analysis of end products by NMR and mass spectrometry

The enzymatic degradation was performed on 10 ml of 5 mg.ml⁻¹ alginate samples with addition of alginate lyases from *Zobellia galactanivorans* at 30 °C for 18 hours. The samples were then centrifuged, the supernatant lyophilized and dissolved in 0.1M ammonium acetate. The oligouronates were fractionated according to their size by preparative gel filtration chromatography on a ÄKTA system with a 16*60 Superdex 30 column (Amersham, **Biosciences**). Oligosaccharides were eluted with ammonium acetate 0.1 M at a flow rate of ml.min⁻¹ 0.8 at room temperature. Unsaturated oligoalginates were detected by A₂₃₅ and the relative amounts of the different end products were determined as the peak area. The fractions belonging to the same peak were pooled, lyophilized and purified further by repeating the SEC procedure on the Superdex 30 column. The pure fractions were lyophilized and analyzed by ESI-MS using a Bruker Esquire-LC mass spectrometer (Bruker Daltonics Inc., Billerica, MA, USA). The samples were dissolved in 1:1 MeOH-H₂O and were directly injected into the electrospray with a flow of 2 μ l.min⁻¹. The mass range scanned was from 100 to 1100 atomic mass units (amu). All acquisitions were recorded in both negative and positive mode and treated by Bruker Daltonics esquire LC 4.5, data analysis version 3.0. To determine the structure of the end products, the oligosaccharide signals were assigned from onedimensional proton and selective CSSF TOCSY with Keeler filter (selcssfdizs.2) from Bruker library NMR spectra and from two-dimensional ¹H-¹H TOCSY NMR experiments using a mixing time of 90 ms. The NMR spectra were referenced at 30 °C using acetone ($\delta_{\rm H}$ 2.225, $\delta_{\rm C}$ 31.05) as internal reference.

Protein crystallization and 3D crystal structure determination

The first crystallization-screening experiments for both the catalytic domain of AlyA1_{PL7} and the full length dimeric sample of AlyA5 were carried out at 292 K with the sitting-drop vapor diffusion method in 96 well Corning plates (no. 3551). The crystallization screening was performed with JCSG+ and PACT screens

from Qiagen, which makes a total of 192 conditions in two 96-well plates. A crystallization robot (Proteomics Solutions, Honeybee) was used for dispensing the drops that contained 300 nl protein solution that were mixed with 150 nl of reservoir solution After visual identification of initial crystallization conditions, these were further optimized in 24 well Linbro plates by the hanging-drop vapor diffusion method. For AlyA1_{PL7} optimal conditions were obtained for a solution of 11.3 $mg.ml^{-1}$ protein concentration supplemented with oligoguluronate (0.1 mg.ml⁻¹). Drops of 2 μ l volume of this protein solution were mixed with 1 µl of crystallization solution that contained 0.2 M KSCN and 28% PEG-MME 2000, and equilibrated against a reservoir containing 500 µl. For AlyA5 (dimer) the optimized conditions were obtained for a protein solution of 7.7 mg.ml⁻¹. Drops of 2 μ l volume of this protein solution were mixed with 1 µl of crystallization solution that contained 22% PEG 3350 and 0.2 M sodium/potassium tartrate, and equilibrated against a reservoir containing 200 µl. The native crystals of both alginate lyases were flash-frozen at 100 K in a crystallization solution supplemented with 10% glycerol. Data were collected to 1.43 Å resolution for AlyA1_{PL7} and 1.75 Å for AlyA5 on beamline PROXIMA-1 equipped with an ADSC Quantum Q315r detector (SOLEIL, St Aubin, France) and a wavelength of 0.98 Å. All data sets were treated, scaled and converted with the program suite XDS (39). The 3D structures were solved by the molecular replacement method with ALY-1 from Corynebacterium sp. (PDB 1UAI) for the crystals of AlyA1_{PL7} and A1-II' from Sphingomonas sp.A1 (PDB 2Z42) as initial model, in the case of AlyA5. The starting phases were optimized using the programs ARP/wARP (40) and REFMAC 5 (41) as part of the CCP4 suite (42). The refinement was carried out with REFMAC 5 and the final model building with Coot (43). Water molecules were added

automatically with REFMAC-ARP/wARP and visually verified. In all cases, the stereochemistry of the final structure was evaluated using PROCHECK (44). All further data collection and refinement statistics are summarized in Table 3.

The atomic coordinates and structure factors (codes 3zpy, and 4be3) have been deposited in the Protein Data Bank, Research Collaboratory for Structural Bioinformatics, Rutgers University, New Brunswick, NJ (http://www.rcsb.org/).

Results

Phylogenetic analysis

The PL7 family has been recently subdivided in subfamilies based on sequence similarities (25). To identify the subfamily for the three PL7 enzymes AlyA1, AlyA2 and AlyA5 from Z. galactanivorans, 35 additional selected sequences from the same family were used to derive a phylogenetic tree (Figure 1). This approach clearly distinguished the five existing subfamilies within the PL7 family (SF1 - SF5). It reveals that the three PL7 enzymes encoded in the Z. galactanivorans genome belong to different subfamilies. AlyA1 is distantly related to enzymes classified in SF3 and thus is a new member of this subfamily. AlyA5 can also be confidently assigned to SF5. AlyA2 was found to belong to a wellsupported group composed of 4 other enzymes from marine Flavobacteriaceae. A condition imposed by Lombard et al. (25) when creating subfamilies, was that the group contained at least five members. Thus, we propose that AlyA2 and the four other enzymes in the same group belong to a new subfamily of PL7. This new subfamily, which we hereby define as SF6, appears to be conserved only in marine representatives of the Flavobacteriaceae, possibly representing a niche-specific evolution of PL7 enzymes. As subfamilies correlate in general with substrate specificities, they can help predict the preferred substrate of new enzymes (25). To date, two alginate lyases have been

characterized in SF3 and seven in SF5 (CAZY database, December 2012). They all are classified as $poly(\alpha-L-guluronate)$ lyases (EC 4.2.2.11). Thus, we hypothesize that AlyA1 and AlyA5 should preferentially cleave G motifs (Figure 2). No prediction can be made on AlyA2 specificity, since none of the proteins belonging to this new subfamily SF6 has yet been characterized.

Overexpression and purification of AlyA1_{PL7} and AlyA5

To characterize the enzymatic activities and structural properties of the new PL7 enzymes from Z. galactanivorans, the coding sequences of the mature proteins (full-length for AlyA2 and AlyA5 and only the catalytic module for AlyA1) were cloned into pFO4 expression vector. Expression tests in E. coli BL21(DE3) grown in ZYP medium (37) were successful for the recombinant proteins AlyA1_{PL7} and AlyA5, but failed for AlyA2 (data not shown). We thus focused on the study of AlyA1_{PL7} and AlyA5. High amounts of soluble recombinant proteins were obtained and purified to homogeneity using a nickel affinity chromatography followed by gel filtration. The two steps of purification yielded 30 mg of recombinant AlyA1_{PL7} from 200 ml of ZYP medium. The apparent molecular mass was between 26.5 kDa and 29 kDa (estimated from the calibration of the size-exclusion column and the SDS-PAGE, respectively). This is concordant with the theoretical molecular mass of 27.5 kDa calculated for the protein sequence of the catalytic domain using ProtParam implemented in the ExPASy Proteomics Server (45). The recombinant AlyA5 eluted from the final size exclusion chromatography as two distinct peaks (data not shown), corresponding to apparent molecular masses of 69.5 kDa and 38.3 kDa. SDS-PAGE analyses showed that these two populations contained only one protein form under denaturing conditions, with an apparent molecular mass of 42 kDa. Compared with the theoretical

molecular mass of 38.3 kDa, the proteins eluted in the first and second peaks correspond roughly to a dimer and a monomer of AlyA5, respectively. This was confirmed by the difference in radius of gyration that was measured by dynamic light scattering as 2.5 nm and 3.2 nm for the mono-disperse samples of purified monomeric and dimeric form of AlyA5 respectively. The purification of the recombinant AlyA5 yielded 2.2 mg of monomeric form and 10 mg of dimeric form from culture of 200 ml. The dimeric form was stable in solution, as it eluted as a single peak after a second size exclusion chromatography, corresponding to а molecular mass of 69.5 kDa (data not shown).

Characterizationoftherecombinant AlyA1

Biochemical properties of AlyA1_{PL7}: The action of the recombinant catalytic domain of AlyA1_{PL7} on alginate induced a strong increase in the absorbance at 235 nm, confirming that it is an alginate lyase acting via the β -elimination mechanism, producing the 4,5-unsaturated 4-deoxy-Lerythro-hex-4-enepyranosyluronate (Δ) on the non-reducing end (Figure 2). The optimal temperature and salinity were 30°C and 200 mM NaCl, respectively (Figure 3A and B). AlyA1_{PL7} showed the highest activity at pH 7.0 in Tris-HCl buffer (Figure 3C). This is similar to the optimal pH determined for other alginate lyases from marine bacteria (46). No activity was detected outside the range pH 4.0-8.5. The choice of the buffering molecule appeared to be critical, as exemplified by the 40% decrease in activity in MOPS buffer at pH 7.0 compared to Tris-HCl. Addition of the chelating agents, EDTA or EGTA, in the reaction mixture reduced the enzyme activity by 90% and 95%, respectively (data not shown). This inhibition was only incomplete when the chelating agents were incubated with the enzyme overnight but not added to the reaction mixture (43% and

58%, respectively). This suggests that the interaction between the enzyme and divalent cations takes place during substrate fixation, and not before. To determine the mode of action of AlyA1_{PL7}, the products resulting from the degradation of alginate were sampled periodically and analyzed by C-PAGE. After two minutes, the polysaccharide fraction was already largely degraded (Figure 3E). Following the reaction from 2 min to 14 h, the size of the detected oligosaccharides decreased with time, rapidly reaching the smallest degree of polymerization (DP) of two. The appearance of intermediate larger oligosaccharides, DP4 to DP20, at early stages demonstrates that AlyA1_{PL7} was acting with an endo-lytic mode on alginate.

Substrate specificity of AlyA1_{PL7}: The enzyme activity was measured on three polymeric alginate substrates, chosen for their difference in the M/G ratio. The kinetic parameters were clearly correlated with the guluronate content of the substrate. The K_m determined on G-rich alginate samples was 4-fold lower than on M-rich alginate samples, while the inverse relation is true for k_{cat}/K_m (Table 1). This that AlyA1_{PL7} preferentially suggests cleaves G-stretches. The substrate specificity was then determined by carrying out the enzymatic reaction directly in the NMR tube. The ¹H NMR chemical shifts were assigned according to previously published data on alginate polyand oligosaccharides (47-51). The structure of the reducing end sugar can be easily identified by the characteristic chemical shift of its anomeric proton signal. Thus a lyase activity on a G-M or G-G diads will lead to the appearance of a doublet at 4.70- $^{3}J_{HH}$ with 4.75 ppm = 8.6 Hz corresponding to the β -anomeric proton of a G residue. A lyase activity on M-M or M-G diads will give a signal at 4.75-4.80 ppm with a ${}^{3}J_{HH} < 2$ Hz corresponding to the β -anomer of an oligomer formed by the cleavage of such a diad. Figure 4 shows that when the crude alginate was submitted to the action of $AlyA1_{PL7}$, the characteristic doublet of G-reducing ends, H1 at 4.74 ppm with ${}^{3}J_{HH} = 8.6$ Hz, appeared in the NMR spectrum, demonstrating that the enzyme requires a G-unit adjacent to the cleavage site on the non reducing end. The chemical shifts of the protons of the unsaturated non-reducing end (further denoted as 'delta' and symbolized by Δ) are dependant on the nature of the nearest sugar residue. This neighbour sugar residue can be identified from the shift of the H-4 signal of Δ . A H4(Δ) doublet at δ 5.75 ppm indicates that the neighbour is a G residue while the chemical shift of the H-4 signal appears at 5.67 ppm if the neighbour to the unsaturated non-reducing end is a M residue. As shown in Figure 4, the intensity of the H4 Δ G signal is much higher than that of H4 Δ M, indicating that $\Delta G(X)_n G$ oligosaccharides with X being a G or M sugar are predominantly formed by AlyA1_{PL7}.

The uronic acid residue initially involved at the other end of the linkage cannot be identified from hydrolysis of crude alginate since β-elimination on β-Dα-Lmannuronosyluronate or on gulupyranosyluronate produces in both 4-deoxy-L-erythro-hex-4cases enepyranosyluronate. Thus, to further investigate the substrate specificity of the enzyme, that is to determine if $AlyA1_{PL7}$ performs β-elimination indifferently on G-M and G-G diads, the degradation of poly-G, poly-M, and poly-MG purified blocks also monitored by ¹H was NMR spectroscopy. As shown in Figure 5A, the spectrum of the poly-MG blocks remains unchanged, indicating that the enzyme is not active on this substrate. No cleavage is either observed for the MM-diads (Figure 5B). The minor products seen in the NMR spectrum arise from the cleavage of some GG diads present in the polymer, as demonstrated by the chemical shift of the H4 Δ signal. Finally, a clear cleavage of GG-blocks is observed (Figure 5C), confirming that the enzyme cleaves only

between two guluronate-units of the polysaccharide chain.

Characterization of end products: The oligosaccharides produced by degradation of the crude alginate with AlyA1_{PL7} were separated by size exclusion chromatography. The separation profile of the oligosaccharides showed three major peaks (data not shown) and the molar fractions of the different products were estimated from the SEC chromatogram (Table 2). The ESI-MS analysis showed *m*/*z* of 351.0, 526.8, and 703.0 [M-H] indicating di-, tri- and tetrasaccharide fragments, respectively. The lyase produced mainly trisaccharide and tetrasaccharide oligomers with a total content of 41% respective 36%. Around 19% of the isolated oligomers were disaccharide and only a small amount of pentamers and hexamers were observed.

The uniform sized fractions were not separated further and the structure and amount of the oligosaccharides present in each fraction were determined by NMR. The NMR spectrum of the disaccharide fraction showed that only ΔG was produced. The NMR spectra of the fraction corresponding to trisaccharides showed the existence of two products, ΔGG and ΔMG , present in amounts of 88 and 12% respectively. This was evidenced by the observation of two H4 Δ signals, one at 5.75 ppm corresponding to Δ at the non reducing end with a G neighbour and one at 5.67 ppm for Δ with an M neighbour. The H2 of G at the reducing end also has a characteristic chemical shift, depending on the neighbouring sugar units, being of 3.47 ppm with G as neighbour and 3.54 ppm with M. The amount of ΔGG (88%) and ΔMG (12%) was determined by integration of the H4 Δ G and H4 Δ M signals.

The NMR spectra of the fraction containing the tetrasaccharides also show the concomitant presence of several patterns. Three H4(Δ) signals were assigned to Δ GGX, Δ GMX and Δ MXX, according to the value of their chemical shift at 5.74 ppm, 5.75 ppm and 5.67 ppm respectively. Two well separated signals were observed for the H2 G-reducing end with different neighbours, one with a characteristic chemical shift at 3.44 ppm for the reducing end with the G neighbour and 3.51 ppm for the reducing end with the M neighbour. Since only G-reducing end sugars are present, the products are ΔGGG , Δ GMG and Δ MXG. Also two Δ H2M signals were observed in the selective TOCSY experiments (Figure 6). Thus four different tetrasaccharides are present Δ GGG, Δ GMG, Δ MGG and Δ MMG. Integration of the H4 Δ G and H4 Δ M signals indicated that ΔGGG and ΔGMG together represent 84% of the total content of tetrasaccharides while ΔMGG and AMMG account for 16% of the molecules.

Minimal recognition pattern of AlyA1_{PL7}: The NMR analyses of the end products reveal a high content of di-, triand tetrasaccharides and only very low contents of penta- and hexasaccharides. This would be in agreement with a oligosaccharide minimal of size recognized, bound cleaved and by AlyA1_{PL7} being a pentasaccharide. Further experiments showed that at least the hexasaccharide G6 is indeed cleaved by $AlyA1_{PL7}$ (data not shown), while the accumulation of tetrasaccharides in the end-products indicate that these are not further cleaved. Purified pentasaccharides were not available but their degradation pattern can be derived from the NMR analysis of the products obtained after enzymatic degradation. Accordingly, the structures of the pentasaccharides that can be cleaved by AlyA1_{PL7} are: GGGGG, GGMGG, GGGMG and GGMMG (Table 4).

Crystal structure of AlyA1_{PL7}: The 3D crystal structure of AlyA1_{PL7} was determined at 1.43 Å resolution by molecular replacement (Table 3). The asymmetric unit of the triclinic unit cell contained two molecules giving rise to a

global solvent content of 41%. For both molecules 248 out of 252 residues could be modelled into the electron density, the missing residues being located at the Nterminal of the polypeptide chain. In addition, the crystal structure of AlyA1_{PL7} contains two calcium ions and 749 water molecules. The overall structure of AlyA1_{PL7} is that of a PL7 β -sandwich jelly roll fold, formed by two anti-parallel β sheets stacked against each other (Figure 7A). The outer, convex sheet is composed of 5 β -strands while the inner, concave sheet is composed of 7 β -strands forming a groove that harbours the catalytic active site. Consistent with its membership in PL7 SF3, AlyA1_{PL7} displayed lowest rmsd of 1.5 upon structural superimposition to Aly-1 from Corynebacterium sp. (1UAI). The structural similarity to A1-II' from Sphingomonas sp.A1 is reflected by an rmsd of 1.76. As is typical for enzymes displaying this fold, the major structural differences are located in the flexible loops that delimit and surround the catalytic groove (Figure 8A). In particular, and in contrast to A1-II' from Sphingomonas sp.A1 (PDB 2ZAA), two of these loops display two short helical structures (residues Gly265 to Asn269 and Gly359 to Gln365, see Figure 7A, B; Figure 8; Figure 9). A third loop localized between the strands $\beta 8$ and $\beta 9$ (Gly324-Asp335) is absent in the structure of Aly-1 (1UAI) and displays a largely different conformation than the equivalent loop in A1-II', in this way participating to the open accessibility of the active site groove in AlyA1_{PL7} (Figure 7A, 8A). This is in contrast to A1-II' which displays a tunnel shaped active site (52). Possibly, the three loops of trapped an $AlyA1_{PL7}$ are in open conformation in this structure and move to form a tunnel upon substrate binding. The electron density maps defining these loops in our structure, however, are perfectly defined and the potential flexibility of these loops thus remains an open question. At the bottom of the groove (Figure 10A), three strictly conserved residues, Gln321, His323 and Tyr420 that were reported to be involved in the lytic activity in A1-II' from *Sphingomonas sp*.A1 (52), form the active site of $AlyA1_{PL7}$.

Characterization of the recombinant AlyA5

Biochemical properties of AlyA5: When the dimeric form of AlyA5 was incubated with alginate, recordings of A₂₃₅ showed a kinetic in two phases (Figure 3F). In the first phase, the absorption increased, showing that AlyA5 cleaved alginate by a β-elimination mechanism and confirming its annotation as an alginate lyase. This was followed by a decrease in A_{235} . suggesting the transitory nature of the UVabsorbing material. Such a profile is typical for exolytic lyases producing unsaturated monosaccharides (53). Indeed, products subsequently these reaction convert non-enzymatically to the most stable 5-keto-structure, i.e., 4-deoxy-Lerythro-5-hexoseulose uronic acids (DEH), which does not absorb at 235 nm due to the transfer of the unsaturation to the ketone group (38, 54, 55). The addition of at least 2 mM Ca^{2+} in the reaction mixture was essential for the activity (data not shown). AlyA5 showed the highest activity at pH 7.0 in MOPS buffer (Figure 3D) and displayed a narrow range of tolerance. The choice of the buffering molecule was critical, as exemplified by the 50% decrease in activity in Tris-HCl buffer at pH 7.0 compared to MOPS. To analyze from which end of the substrate the exolytic enzyme attacks, AlyA5 activity was monitored on polyG blocks that had their reducing ends reduced using sodium the digestion. borohydride prior to Compared to the reaction with classical polyG substrate, AlyA5 retained 75% of its activity on such reduced substrates, indicating that the enzyme cleaves of the unit at the non-reducing end. The 25% decrease in activity can be attributed to loss of material during the successive evaporation/resolubilization steps which are part of the reduction protocol.

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Substrate specificity of AlyA5: To assess the substrate specificity of AlyA5, the activity was tested on MG-, M- and Gblocks. The reactions were followed by the absorbance at 235 nm and by ¹H NMR (Figure 3F; Figure 11). The three substrates were degraded, indicating that the enzyme has a broad substrate tolerance and can cleave M-M, M-G and G-G linkages at the non-reducing end. The activity was however depending on the block structure. Specific activities were calculated in the first phase of the reaction, corresponding to the increase in A_{235} (Figure 3F). AlyA5 was much more active on G blocks (449.3 U.mg⁻¹) than on MG blocks (13.5 U.mg⁻¹) or M blocks (2.0 U.mg⁻¹). On the NMR spectra (Figure 11), in addition to the characteristic H4 Δ signals indicating formation of unsaturated oligosaccharides, other signals were observed in the region 1.6-2.6 ppm and 3-4 ppm (vide infra).

Characterization of end products: The degradation of poly-G was followed by NMR to characterize the end products resulting from AlyA5 activity. Figure 12 shows that during the course of the reaction. the concentration of the disaccharide ΔG increased while the concentration of the longer unsaturated oligosaccharides decreases. А small amount of the unstable Δ -(4,5)-unsaturated pyranose monomer was also identified in spectra. the NMR As the reaction progressed, signals corresponding to protons from CH₂ groups appeared in the NMR spectra (Figure 12). These proton signals have chemical shifts typical for CH₂ groups close to a keto group or hemiketal (56). Thus the NMR data confirms that one of the reaction products is DEH resulting from the spontaneous conversion of the unsaturated monosaccharide Δ . The concentration of the unsaturated disaccharide always stays low during the course of the reaction, compared to the signal intensities from

DEH. This indicates that ΔG is a minor product of the degradation reaction.

Minimal recognition: Incubation of the unsaturated trisaccharide Δ GG with AlyA5 resulted in the formation of Δ and ΔG detected by ¹H NMR (Figure 13). The amount of ΔG was much higher than that of Δ confirming that the unsaturated monosaccharide is not stable in aqueous Concordantly, solution. signals corresponding to deoxy protons appeared in the 1.5-2.6 ppm region of the ¹H-NMR spectra. As seen in Supplemental Figure S7, some Δ MG was present as traces in the solution containing ΔGG . ΔMG was degraded during the course of the reaction into Δ and ΔG indicating that AlyA5 is also able to cleave the Δ -M glycosidic linkage. The fact that ΔM was not observed strengthens the conclusion that AlyA5 attacks its substrate from the nonreducing end. The enzyme was also active on ΔMM producing $\Delta + \Delta M$ (data not shown). The activity on ΔGM was not studied but the combined data above suggest that this compound should also be a substrate for AlyA5. The saturated oligosaccharides MMM and GGG were also found to be substrates of AlyA5, producing ΔM and ΔG , respectively. Disaccharides were not substrate for the enzyme. Thus, the minimal recognition oligosaccharides are saturated and unsaturated trisaccharides, and the enzyme is able to cleave of a G, M or Δ unit from the non reducing end.

Crystal structure of AlyA5: AlyA5 represents the first 3D crystal structure of an exolytic alginate lyase belonging to the family PL7 SF5. The structure was determined at 1.75 Å resolution and the asymmetric unit contains two independent molecules. For both molecules, the atomic positions for 312 amino acids out of 337 residues could be constructed into the electron density, with 22 residues missing at the N-terminal and 3 at the C-terminal ends of the polypeptide chains. The crystal structure trapped tartrate and glycerol molecules, present in the buffer solutions, for both independent AlyA5 molecules. In addition the asymmetric unit contained 623 water molecules, equally distributed around the two alginate lyase molecules. The overall fold is very similar to that of AlyA1_{PL7} (rmsd between AlyA1_{PL7} and AlyA5 is 2.02), but even closer to both A1-II' (2Z42) and Aly-1 (1UAI) with an rmsd of 1.4 (Figure 8B). The structural determinant leading to the exolytic activity can easily be identified as three large additional loops, Trp197-Asp217, Ser257-Glu284 and Gly304-Asp318 (Figure 7C, D; Figure 8B, Figure 9) that close up at one end the catalytic groove of the concave β -sheet. In particular, Trp313 is located close to the three conserved catalytic residues, Gln147, His149 and Tyr299, hydrophobic constituting a wall obstructing the continuation of the groove (Figure 10C). As a consequence the catalytic active site of AlyA5 is located at the bottom of a pocket (Figure 7D), in contrast to the wide-open cleft observed in AlyA1_{PL7}. Interestingly, although our structure does not contain any bound ligand, the loop between β 4 and β 5 (Ser92-Asn101 in AlyA5, equivalent to Gly263-Gln271 in AlyA1_{PL7}, Figures 8 and 9) displays exactly the same closed tunnel conformation as the equivalent loop in A1-II' with trapped substrate (closed form).

Discussion

Heterotrophic bacteria have evolved diverse strategies to efficiently assimilate the polysaccharides available in their habitat. In general, polysaccharide degrading organisms will produce several enzymes that together, synergistically or by complementary actions, will break down the complex polymers into smaller mono- and oligosaccharide building-units. degradation These products are subsequently taken and further up catabolized. A large number of studies are available on such enzyme systems for

microorganisms degrading terrestrial plant cell walls. On the contrary, only few studies are available on the molecular bases of marine algal cell wall utilization. For example, work on catabolic systems targeting the polysaccharide alginate has in general focused on enzymatic pathways for the complete degradation of bacterial alginate, rather then on alginate present in the brown algal cell wall. These studies shown that bacterial have alginate breakdown by terrestrial species such as Pseudomonas sp. (38, 57), Sphingomonas sp. (58) or Agrobacterium (59) relies on a system that utilizes a particular ABCtransporter complex to channel the polymer to cytoplasmic lyases. In contrast, as evidenced by our recent study, the marine flavobacterium Z. galactanivorans produces a series of intra-cellular, outer membrane bound and secreted alginate lyases, most of which are encoded by PUL operon structures comprising SusCD-like genes coding a putative oligo-alginate import system (20). Interestingly, in the context of a bioengineering project aiming at building an E. coli strain able to produce bioethanol from brown seaweed, Wargacki et al. (60) have shown by an indirect genetic approach, that the outer membrane porin KdgMN (V12B01 24269 and V12B01_309) and the sodium/solute symporter V12B01 24194 (plasma membrane) are responsible for the uptake of oligoalginates in V. splendidus 12B01, pointing towards the hypothesis that alginate uptake systems in Flavobacteriaceae and Vibrionaceae are unrelated.

Enzymes specifically recognizing and cleaving D-mannuronate or L-guluronate units in alginate have been identified in following PL families: PL5, PL7, PL14, PL15, PL17, PL18 and PL20 (**31**). The substrate specificities and modes of action of these alginate lyases vary between families. Namely, the family PL14 enzyme shows a pH dependant exo- and endo-lytic activity (**61**), while the PL5 enzyme with known structure is a M-M specific lyase

efficiently liquefies acetylated that (62), and the structurally alginates characterized enzyme belonging to family PL15 cleaves off the unsaturated sugar in an exo-lytic manner (63). Most examples of characterized enzymes come from family PL7, where G-, M- and MGspecific enzymes have been described (52, 64-66).Albeit the wealth of biochemical data and the occurrence of numerous annotated gene sequences, only few structural studies of alginate lyases are available, among which three family PL7 enzymes from different terrestrial bacteria (65-67). All PL7 enzymes for which structures have been determined so far are endo-lytic and they randomly degrade alginate chains to produce unsaturated oligo-alginates with various chain lengths. The genome of Z. galactanivorans contains seven putative alginate lyase genes, three of which are members of family PL7. The transcription of the PL7 coding genes (*alyA1*, *alyA2* and *alyA5*) was strongly induced when Z. galactanivorans used alginate as sole carbon source (20). phylogenetic analyses The of the sequences of the three PL7 enzymes show that they fall into three distinct subfamilies, namely SF3, SF5 and the new SF6, which supports the hypothesis that these lyases play different roles and have complementary activities. In the present study, the biochemical and structural characterization of two family PL7 alginate lyases from the alginolytic system of Z. galactanivorans, the secreted AlvA1_{PL7} and the outer membrane bound AlyA5, reveals that at least two of these PL7 enzymes have very complementary activities.

Indeed, our results show that together and in vitro, these two enzymes can transform natural occurring algal alginate completely into disaccharide units (ΔG or ΔM) plus the unsaturated monosaccharide that spontaneously converts into 4-deoxy-Lerythro-5-hexoseulose uronic acid (DEH).

In agreement with substrate specificities observed in PL7_SF3, AlyA1_{PL7} is a true

G-specific, endo-acting enzyme, that specifically cleaves between two G-units. On the other hand, AlyA5 represents the far first exo-lytic PL7 member so described. Interestingly, AlyA5 appears to be able, albeit having a preference for Δ - or G-units, to cleave any of the three types of non reducing ends, Δ , G or M, from oligoalginate chains. This versatility is rather rare among described alginate lyases, which are in general specific towards one of the three types of block-structures occurring in alginate. One other endo-lytic PL7 alginate lyase, A1-II' (accession code Q75WP3, not included in any attributed subfamily, see Figure 1), has been reported to display broad substrate specificity (68), although these two enzymes do not fall into the same subfamily (54).

Together with PL14 (61, 69), this is the second example of an alginate lyase family enzymes active containing both on polymer chains and on short oligosaccharides. This is in agreement with the general rule that endo- or exo-lytic defined by family activity is not membership (25) but rather by specific structural differences in loops surrounding the active site formed by the conserved scaffold. Here, large loop insertions of up to 27 residues in the sequence of AlyA5 block the catalytic groove at one end leading to a pocket-architecture, whereas most of the secondary structures and residues involved in catalysis are well conserved with the other PL7 members (Figure 9).

A superimposition of the crystal structure of A1-II' in complex with an oligo-alginate allows positioning the substrate with respect to the catalytic residues in AlyA1_{PL7} or AlyA5. In AlyA1_{PL7}, in addition to the conserved catalytic triad (Gln321, His323 and Tyr420), several residues, such as Lys214, Arg274, Arg278, Tyr414 and Lys416 take equivalent positions as those described to be in contact with the ligand molecule in the complex structure of A1-II' (Figure 10). The major differences between AlyA1_{PL7} and the A1-II'-substrate complex are: there is no equivalent to Asn141 (A1-II' numbering) that is part of the first loop (residues Cys260 to Pro273 in AlyA1_{PL7}; Figure 9), secondly Gln97 is replaced by a shorter residue, Asn216 in AlyA1_{PL7}, making the distance to interact with the substrate much longer, and thirdly, in AlyA1_{PL7} an aspartate residue (Asp336, absent in A1-II') is positioned close to the catalytic active site.

But the most striking feature that differentiate AlyA1_{PL7} and AlyA5 from other PL7 alginate lyases is the presence of several more negatively charged aspartate and glutamate residues in vicinity of the site (Figure 1,+1 cleavage 10). In AlyA 1_{PL7} , the equivalent position of Pro202 of A1-II' (Figure 10A) is occupied by Asp336 (Figure 10B), which would be both in sterical clash and electrostatic repulsion with the uronate-group if the +1G-unit would bind exactly in the same deduced position as from the superimposition with A1-II'. In A1-II', Lys218 interacts with the G-unit bound to the -1 binding site. The equivalent region in AlyA1_{PL7} comprises strand β 9 and helix $\alpha 2$. The corresponding strand in A1-II' is badly superimposed with β 9 and there is no equivalent of helix $\alpha 2$. As a consequence, the Ca of Lys218 is roughly substituted by a glycine (Gly359) and its side chain is spatially replaced by the acidic group of Glu363. Again, this conformation appears incompatible with a similar binding of a G unit in -1.

In AlyA5 there are even three acidic residues that would come into sterical clash with alginate G-units if bound as in A1-II' (Figure 10C); here, Pro202 is replaced by Glu157 and again the region including Lys218 in A1-II' is poorly conserved, with strands $\beta 9$ and $\beta 10$ of significantly displaced AlvA5 in comparison to the corresponding strands in spatially A1-II'. Thus Lys218 is substituted in AlyA5 by a cluster of two acidic residues, Glu179 and Asp191, their side chains pointing toward the carboxylic

group of the modelled G-unit in subsite +1. Remarkably, these residues are situated in the extended loops that are found in variable conformations in the different PL7 alginate lyase structures. In our crystal structure of AlyA5 the loops are closed above the active site groove, although closed forms in other PL7 enzymes are only observed in presence of substrate. Overall, the different loop conformations observed in the various crystal structures are indicative of flexible loops, and most probably loop movements assist substrate binding and release. Together with the finding that chelating agents inhibited the lyase activities, these structural features lead us to hypothesize that substrate binding and recognition in AlyA1_{PL7} and AlyA5 includes interactions mediated by calcium ions. Loop-movements displacing the acidic residues together with calciumbinding could bring the residues from their actual conformations into positions for productive interactions of the enzyme with its natural, calcium-chelated alginatesubstrate. A similar substrate binding mode, involving acidic amino acids and calcium ions to a negatively charged substrate is observed in other

polysaccharide lyases, such as chondroitin B lyase (70) and pectate lyases (71). The rationale of such a variation in the binding and recognition mode of the here studied PL7 alginate lyases, as compared to that of the previously described enzymes, would be found in the different nature of alginate originating from bacteria or from brown seaweeds. Indeed, the negative charges of algal alginates are not masked by acetyl group as in bacterial alginates. Moreover, alginate chains have multiple interactions other cell wall compounds, with polysaccharides, phlorotannins, proteins but also inorganic ligands such as calcium or iodine ions (2, 13). In conclusion, our biochemical and structural analyses of first marine alginate these lyases demonstrate that they have indeed complementary roles in the degradation of algal alginate. Moreover, these results shed light on the molecular basis for adapted specificity and binding mode to their natural substrate. Further work is under way to confirm the here described hypothesis and to proceed with the characterization of all the other enzymatic players of the complete alginolytitc system of Zobellia galactanivorans.

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Tables

Table 1: Kinetic parameters of $AlyA1_{PL7}$ activity on alginate with different guluronate contents.

Guluronate content in alginate	K_{m} (μM)	\mathbf{k}_{cat} (s ⁻¹)	$k_{cat}/K_m (nM^{-1}.s^{-1})$
66.7%	1753	12.66	7.22
52.6%	3087	17.89	5.79
33.3%	6180	19.51	3.16

Table 2: Amount of different oligosaccharides produced by $AlyA1_{PL7}$ assessed by SEC and ¹H NMR.

Oligomers	UV	NMR	Total contribution
AlyA1 _{PL7}			
Dimers	19%	-	
ΔG		100%	19%
Trimers	41%	-	
ΔGG		88%	36%
ΔMG		12%	5 %
Tetramers	36%	-	
∆GXG		84%	30%
ΔMXG		16%	6%
Pentamers	2%	-	2%
Hexamers	1%	-	1%
> dp6	1%	-	1%

	AlyA1 _{PL7}	AlyA5 dimer			
Data collection					
Beamline	PROXIMA 1				
Wavelength	0,98 Å				
Space group	P1	P2 ₁ 2 ₁ 2 ₁			
Unit cell	a=43,86 Å; b=50,39 Å; c=55,52 Å α =69,14°; β =90,02°; γ =84,92°	a=93,42 Å ; b=93,91 Å; c=130,15 Å α=β=γ=90,00°			
Resolution range ^a (Å)	43,66-1,43 (1,47-1,43)	41,95-1,75 (1,80-1,75)			
Total data	358902	967134			
Unique data	78881	114168			
Completeness (%)	96,1 (94,2)	99,4 (93,3)			
Mean I/ $\sigma(I)$	14,16 (7,79)	16,53 (2,30)			
R_{merge} (%)	7,8 (21,9)	9,6 (84,4)			
Redondancy	4,55	8,47			
Refinement statistics					
Resolution range	40,66-1,43 (1,47-1,43)	41,95-1,80 (1,85-1,80)			
Unique reflexions	74933 (5454)	101057 (7343)			
Reflexions R _{free}	3944 (287)	5319 (386)			
R / R_{free} (%)	14,6/20,1 (11,0/19,4)	15,4/17,8 (21,1/24,6)			
RMSD bond lengths	0,024 Å	0,039 Å			
RMSD bond angles	2,00°	3,02°			
Overall B factor (Å ²)	12,74	29,16			
B factor: Molecule A (Å ²)	10,39	21,74			
B factor: Molecule B (Å ²)	10,32	21,31			
B factor: Solvant (Å ²)	25,00	22,67			
B factor: Ligands (Å ²)	-	31,56			

Table 3: Data collection and refinement statistics for the crystal structures of the native $AlyA1_{PL7}$ and AlyA5.

^aValues in parentheses concern the high resolution shell.

Table 4: Pentasaccharides that can be cleaved by $AlyA1_{PL7}$ and the possible degradation products.

Pentasaccharides		Possible degradation products
GGGGG	\rightarrow	$\Delta GGG + \Delta GG + \Delta G$
GGMGG	\rightarrow	ΔMGG
GGGMG	\rightarrow	$\Delta GMG + \Delta MG$
GGMMG	\rightarrow	ΔMMG

Figure Legends

Figure 1: Unrooted phylogenetic tree of 38 enzymes of the PL7 family.

This phylogenetic tree was calculated by maximum likelihood approach with the program PhyML (34). Uniprot accession numbers are given. Numbers indicate the bootstrap values in the ML analysis. Values lower than 30 were not indicated. Red dots indicate enzymes from Z. galactanivorans. Pink triangles indicate enzymes characterized biochemically. Blue squares indicate that the structure of the protein has been solved. SF: subfamily. A: Azotobacter. Am: Amycolatopsis. C: Cellulophaga. Ca: Catenulispora. Cr: Croceibacter. G: Gramella. K: Klebsiella. Kr: Kribbella. P: Pseudomonas. R: Rhodopirellula. S: Streptomyces. Sa: Saccharophagus. V: Vibrio.

Figure 2: Schematic representation of the catalytic mechanism of G-specific alginate lyses.

The catalytic mechanism in G-specific alginate lyases is postulated to be an anti-elimination. The polysaccharide is cleaved to produce a 4-deoxy-L-erythro-hex-4-enepyranosyluronate moiety (Δ) at the newly formed non-reducing end of the chain; due to loss of the asymmetric center at C-4 and C-5, guluro- or mannurono-configured substrates yield essentially the same product (depending on the stereochemistry at C-1). As a prelude to chain scission, the C-5 proton adjacent to the carbonyl group is abstracted by a suitably poised basic amino acid sidechain (B:). Departure of the glycosidic oxygen is likely to be facilitated by proton donation from a catalytic acid (B:H). Co-ordinating and charge-stabilizing cations, Ca²⁺ or a positively charged amino-acid side chain, are also a common feature of PL active sites.

Figure 3: Effect of temperature (A), concentration of NaCl (B) and pH (C) on AlyA1_{PL7} and pH (D) on AlyA5 activity, (E) C-PAGE analysis of AlyA1_{PL7} degradation products and (F) Degradation kinetics of alginate by AlyA5

A, B. Experiments were conducted in Tris-HCl buffer pH 7.5 using sodium alginate 0.05% as substrate and 12 nM of the purified enzyme. Values are mean \pm S.D. (n = 3). A, Effect of temperature. Activity at 30°C was taken as 100%. B, Effect of [NaCl]. Reactions were conducted at 30°C. Activity with 200 mM NaCl was taken as 100%. C. Effect of pH on AlyA1_{PL7} activity. Experiments were undertaken at 30°C in 100 mM buffer using sodium alginate 0.05% as substrate and 27 nM of the purified enzyme. Sodium citrate (open rhombuses), MOPS (open circles), Tris-HCl (open triangles), HEPES (closed squares) and glycine-NaOH (closed circles) buffers were used in the assay. Activity in Tris-HCl pH 7.0 was taken as 100%. D. Effect of pH on AlyA5 activity. Experiments were done as above except that the protein concentration of purified AlyA5 was 56 nM. Activity in MOPS pH 7.0 was taken as 100%. E. The reaction was conducted at 30°C in 100 mM Tris-HCl buffer pH 7.5 using sodium alginate 0.05% as substrate and 12 nM of the purified enzyme. Aliquots were sampled before adding the enzyme (lane 1) and after 2 min (lane 2), 5 min (lane 3), 10 min (lane 4), 30 min (lane 5), 1 h (lane 6), 2h (lane 7) and 14 h (lane 8). F. Degradation kinetics of G- (black), MG- (dark gray) and M-blocks (light gray) by AlyA5. A235 was measured at 30°C for reaction mixtures (500 µl) composed of 100 mM Tris-HCl buffer pH 7.5, 4 mM CaCl₂, 50 μ g of substrate and 27 nM of enzyme.

Figure 4: Proton NMR spectra of alginate degradation by AlyA1_{PL7}

Proton NMR spectra at 70°C of 0.5 ml solutions (5 mg.ml⁻¹ alginate in 100 mM deuterated phosphate buffer, pH 7.6, 200 mM NaCl and 5 mM CaCl₂) of crude alginate, (A) alone and (B) in the presence of AlyA1_{PL7}.

Figure 5: Proton NMR spectra of defined alginate samples after degradation by $AlyA1_{PL7}$

Proton NMR spectra of (A) poly-MG (B) poly-M and (C) poly-G before and after incubation with AlyA1_{PL7}.

Figure 6: Proton NMR (TOCSY) spectra of defined tetrasaccharides and tetraoligoalginate samples after degradation by AlyA1_{PL7}. A. ¹H NMR spectra of selective TOCSY on Δ H4(G) signal including a magnification of the Δ H2(G) region. B. ¹H NMR spectra of selective TOCSY on Δ H4(M) signal including a magnification of the Δ H2(M) region. C. Tetrasaccharide fraction containing Δ GGG, Δ GMG, Δ MGG and Δ MMG. It is seen that two Δ H2(G) signals are present demonstrating the existence of Δ GGG and Δ GMG, in the same way the two Δ H2(M) signals show that both Δ MGG and Δ MMG are present.

Figure 7: Crystal structure representations of AlyA1_{PL7} and AlyA5

Ribbon and surface representations of the crystal structures of $AlyA1_{PL7}$ and AlyA5. (A) The conserved jelly roll scaffold of $AlyA1_{PL7}$ catalytic module is coloured in green and the flexible, additional loops are coloured in red (B) Surface representation of $AlyA1_{PL7}$. (C) The conserved jelly roll scaffold of AlyA5 is coloured in pink and the additional loops are coloured in blue. (D) Surface representation of AlyA5. A tetrasaccharide GGGG from A1-II' was superimposed to both surface representations.

Figure 8: Tube representation of the superimposition of the 3D structure of PL7 alginate lyases.

(A) Superimposition of AlyA1_{PL7} (green tube) onto AlyPG from *Corynebacterium sp.* ALY-1 (blue tube, PDB code: 1UAI) and A1-II' from *Sphingomonas sp.* A1 (yellow tube, PDB code: 2ZAA). (B) Superimposition of AlyA5 (magenta tube) ontoAlyPG from *Corynebacterium sp.* ALY-1 (blue tube, PDB code: 1UAI) and A1-II' from *Sphingomonas sp.* A1 (yellow tube, PDB code: 2ZAA). Loops that have different configurations or large loop insertions with respect to 1UAI and 2ZAA are highlighted by the arrows and coloured in red for AlyA1_{PL7}and cyan for AlyA5. Residues in the coloured loops all display significantly higher B-factors by 2 to 10 Å² than the mean B-factor of the respective molecules.

Figure 9: Structure-based sequence alignment of AlyA1_{PL7} and AlyA5.

The sequences are aligned with those from two other PL7 alginate lyases with known structure, AlyPG from *Corynebacterium sp.* ALY-1 (PDB code: 1UAI) and A1-II' from *Sphingomonas sp.* A1 (PDB code: 2ZAA). The secondary structures of AlyA1_{PL7} and AlyA5 are shown above and below the alignment, respectively. Conserved amino acids in white highlighted in the red background are identical and those in red are similar. Alpha helices are represented as helices, and β -turns are marked with TT. Green triangles indicate the conserved residues involved in the catalytic machinery. This figure has been generated using the program ESPRIPT (**72**).

Figure 10: Residues surrounding the conserved catalytic active site in AlyA1 $_{\rm PL7}$ and AlyA5.

Close-up view into the active site of (A) A1-II' (pdb code: 2Z42) in complex with an oligoguluronate tetrasaccharide, (B) AlyA1_{PL7} and (C) AlyA5. The potential positions of oligotetrasaccharide molecules were obtained by superimposition with A1-II' in complex with substrate. The respective residues at equivalent positions are labelled. Both AlyA1_{PL7} and AlyA5 contain additional negatively charged residues close to the carboxylate groups of the substrate.

Figure 11: Proton NMR spectra of alginate degradation by AlyA5. Proton NMR spectra (A) poly-M alone, (B) poly-M in presence of AlyA5, (C) poly-MG, (D) poly-MG after addition of AlyA5, (E) poly-G, (F) poly-G after addition of AlyA5.

Figure 12: Proton NMR spectra of alginate degradation by AlyA5

¹H-NMR spectra monitoring the degradation of G-blocks by AlyA5 as a function of time.

Figure 13: Proton NMR spectra of the degradation of a defined alginate-oligosaccharide

by AlyA5. ¹H NMR spectra showing the degradation of Δ GG by AlyA5 as a function of time in minutes and days. Only the region corresponding to unsaturated H4 protons at the non-reducing end is shown. Bottom spectra: Δ GG before addition of the enzyme.

Figure 1



























Figure 9













