Enzymatic depolymerization of alginate by two novel thermostable alginate lyases from *Rhodothermus marinus*

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Alginate (alginic acid) is a linear polysaccharide, wherein (1→4)-linked β-D-mannuronic acid and its C5 epimer, α-L-guluronic acid, are arranged in varying sequences. Alginate lyases catalyze the depolymerization of alginate, thereby cleaving the (1→4) glycosidic linkages between the monomers by a β-elimination mechanism, to yield unsaturated 4-deoxy-L-erythro-hex-4-enopyranosyluronic acid (1) at the non-reducing end of resulting oligosaccharides (α-L-erythro configuration) or, depending on the enzyme, the unsaturated monosaccharide itself. In solution, the released free unsaturated monomer product is further hydrated in a spontaneous (keto-enol tautomerization) process to form two cyclic stereoisomers.

In this study, two alginate lyase genes, designated *alyRm3* and *alyRm4*, from the marine thermophilic bacterium *Rhodothermus marinus* (strain MAT378), were cloned and expressed in *Escherichia coli*. The recombinant enzymes were characterized, and their substrate specificity and product structures determined. At higher temperatures, the lyases were used as catalysts of depolymerization reactions in the generation of hex-4-enopyranosyluronic acid derivatives.

**KEYWORDS**
- alginate oligosaccharides
- thermophilic alginate lyase
- guluronic acid
- mannuronic acid
- *Rhodothermus marinus*
- NMR spectroscopy
Introduction

Alginate (alginic acid) is the most abundant carbohydrate in several types of macroalgae, including marine brown seaweed (Phaeophyceae), and a major constituent of the algal cell wall and intracellular material (approximately 22–44% of the total cell dry weight) (Senturk Parreidt et al., 2018). It is a linear anionic polymer (molecular mass range 10–600 kDa) composed of (1→4)-linked β-D-mannopyranosyluronic acid (β-D-ManpA = M; 4C1 conformation) and (1→4)-linked α-L-gulopyranosyluronic acid (α-L-GulpA = G; 3C4 conformation) units (Figure 1), arranged in varying sequences, i.e., block-regions of both homo- (M- or G-blocks) and hetero- (MG- or GM-blocks) polyuronic acid sequences (Penman and Sanderson, 1972; Gacesa, 1992; Pawar and Edgar, 2012). The constituting monomers β-D-ManpA and α-L-GulpA are C-5 epimers. Alginate oligosaccharides, derived from brown seaweeds, are generally recognized as safe (GRAS) and non-toxic, and are widely used in the food, cosmetics, biochemical, and pharmaceutical industries due to their stabilizing, dehydrating, viscifying, and gelling (with Ca2+) properties (Wong et al., 2000; Draget and Taylor, 2011; Zhang et al., 2021). Moreover, alginate oligosaccharides were demonstrated to have prebiotic activity (Wang et al., 2006). Alginate oligosaccharides and their derivatives have also been attracting attention due to their anti-inflammatory and anti-coagulant activities, and anti-tumor effects (Lee and Mooney, 2012; Chen et al., 2017; Xing et al., 2020). The constituting monomeric products of the lyase reactions may also be considered as important catalytic products as they can be converted into value-added chemicals, e.g., 2-keto-3-deoxy-2-gluconate (KDG). The application potential and the importance of the catalytic products of alginate is reviewed by Zhu and Yin (2015).

The enzymes, capable of catalyzing the depolymerization of algicnic acid by splitting the glycosidic bonds through a β-elimination reaction, are generally known as alginate lyases (also called alginases, alginate depolymerases, transeliminases) (Gacesa, 1987; Wong et al., 2000; Kim et al., 2011; Ertesvåg, 2015). Three types of alginate-depolymerizing enzymes have been defined, i.e., poly(α-L-guluronate) lyase (EC 4.2.2.11), poly(β-D-mannuronate) lyase (EC 4.2.2.3), and bifunctional alginate lyase (Garron and Cygler, 2010; Zhu and Yin, 2015). Additionally, alginate lyases can have either endo- or exo-cleaving specificity. Cleavage of the R1-(1→4)-β-D-ManpA-R2 or R1-(1→4)-α-L-GulpA-R2 glycosidic linkages (R1 and R2 = M- and/or G-containing saccharide) in the alginate polymers leads to a C-4,5 double bond in the D-mannuronate and/or the L-guluronate residue, yielding in both cases Δ-R2 with the new non-reducing terminal residue 4-deoxy-α-L-erythro-hex-4-enopyranosyluronic acid, designated by the symbol Δ (Figure 1; Gacesa, 1987, 1992).

A large number of alginate lyases, from sources ranging from various types of soil and marine bacteria, algae, and fungi to marine animals, have been characterized (Wong et al., 2000). In the Carbohydrate-Active Enzymes (CAZY) database (Drula et al., 2022),1 alginate lyases are assigned to different polysaccharide lyase (PL) families, based on amino acid sequence similarities. Several bacterial alginate lyase genes have been identified, cloned, and sequenced. Both native and recombinantly expressed enzymes have been characterized, giving insight into alginate lyase structure and function (Kim et al., 2009, 2011, 2012; Tondervik et al., 2010; Hamza et al., 2011; Park et al., 2012). Few thermophilic alginate lyases have been described. A structure-function analysis of an alginate lyase from the marine thermophile Defluviitales phaphyphila was recently reported. The enzyme, which was affiliated with a new PL family, PL39, had temperature optimum of activity at 65°C and pH optimum of activity at 5.8 (Wang et al., 2018; Ji et al., 2019). Another alginate lyase from Nitratiruptor sp. SB155-2, affiliated with family PL7, had a temperature optimum of activity at 70°C and pH optimum 6 (Inoue et al., 2016). Further, a recently described thermophilic exolytic alginate lyase of PL family 17, derived from an Arctic Mid-Ocean Ridge (AMOR) metagenomics data set, had pH optimum of activity at pH 6 and retained 100% activity after a 24-h incubation at 60°C (Arntzen et al., 2021).

Alginate lyases have been used for production of defined alginate oligomers and for protoplasting seaweed (Inoue et al., 2007; Chen et al., 2018). Their application for degradation of alginate polymers produced by Pseudomonas aeruginosa in cystic fibrosis patients has been studied (Mrsny et al., 1994; Alipour et al., 2009). Alginate lyases may also be used for the degradation of algal biomass in the production of biofuels or renewable commodity compounds (Jones and Mayfield, 2012; Wargacki et al., 2012; Doshi et al., 2016; Li et al., 2016, 2020). For utilization in those various industrial processes, robust alginate lyases which can function at extreme circumstances, such as elevated temperatures, may be of interest. Thermostability enables purification of recombinant enzymes from a mesophilic host by heat precipitation of the host’s proteins, which consequently lowers downstream production.

1 http://www.cazy.org/
costs. Temperature is a major factor in defining the application range of an enzymatic activity, but optimum temperature of an industrial process is dependent on both substrate and products. High temperature (50–80°C) is optimal for polysaccharide degradation as it increases solubility, reduces viscosity of polysaccharides and facilitates enzymatic access. Thermostable enzymes have longer shelf lives and will tolerate prolonged reaction times and harsh conditions in bioconversion processes. Thermophilic enzymes may also streamline the integration of degradation processes with prior pre-treatment of seaweeds which is optimally carried out at elevated temperatures. Thermophilic and thermostable enzymes reduce the need for cooling from often high pre-processing temperatures of algal biomass that would be required for mesophilic enzymatic alginate degrading processes. High temperature also prevents contamination by spoilage bacteria (Turner et al., 2007).

The genome of the marine thermophile bacterium *Rhodothermus marinus* (strain MAT378) contains four alginate lyase genes, designated *alyRm1* to *alyRm4*. Two of the genes, encoding alginate lyases AlyRm1 and AlyRm2 belong to the CAZY polysaccharide lyase (PL) family PL6 subclass 2 and have been shown to function as endo-poly-MG lyases (Mathieu et al., 2016). This report describes the second two thermophilic alginate lyases, AlyRm3 and AlyRm4, their substrate specificity and products. AlyRm4 belongs to the PL17 and AlyRm3 belongs to a newly described polysaccharide lyase family, PL39. The encoding genes were identified in *Rhodothermus marinus*, a Gram-negative, aerobic, marine thermophile belonging to a sister phylum to *Bacteroidetes*, *Rhodothermaeota* (Munoz et al., 2016). The species which has been isolated from marine habitats around the world in the proximity of hot spring vents, is a known producer of various robust polysaccharide-degrading enzymes (Bjornsdottir et al., 2006; Ara et al., 2020). The thermophilic *R. marinus* alginate lyases with complementing activity add to the previously described activity range of alginate lyases and may have valuable industrial application properties.

The encoding genes *alyRm3* and *alyRm4* were amplified by PCR, cloned into expression vector and expressed in *E. coli*. The resulting recombinant enzymes, AlyRm3 and AlyRm4, were characterized with regard to thermostability, and temperature and pH optimum of activity and subsequently investigated in more detail for their depolymerizing activity on mannanuronan (M-block alginate), guluronan (G-block alginate), and low-viscosity alginate from *Macrocystis pyrifera*. The complete reaction mixtures and the products, isolated by size-exclusion chromatography (SEC), were analyzed by TLC, MALDI-TOF-MS, GLC-EI-MS, and one-/two-dimensional $^1$H and $^{13}$C NMR spectroscopy.

**Materials and methods**

**Cultivation, genome sequencing and analysis**

*R. marinus* strain MAT378/R-21 is a close relative of the type strain *R. marinus* DSM4252. Both strains were originally isolated from a shallow submarine hot spring in SW Iceland (Alfredsson et al., 1988). The MAT378 strain was cultivated in RSC selective medium (Bjornsdottir et al., 2007), containing 0.2% soluble starch (Merck). Pure high-molecular-weight DNA and plasmid DNA was isolated using the MasterPure Gram Positive DNA Purification kit (Lucigen) according to manufacturer's instructions and used for genome sequencing. The genome was originally sequenced by Sanger sequencing with approximately 5x coverage. With the emergence of high throughput sequencing, the genome was re-sequenced. Libraries were prepared using the Nextera Flex and Nextera MatePair methods (Ilumina) and sequenced on the Illumina MiSeq platform. The generated sequence reads were assembled by using SPAdes assembler (Banoschvich et al., 2012) and annotated by using the RAST annotation server (Overbeck et al., 2014).
Identification of genes encoding alginate lyase

Putative alginate lyase encoding genes were identified by the NCBI BLAST program (non-redundant protein sequences database). Sequence Alignments were performed using the EBI ClustalW2—Multiple Sequence Alignment tool. Further protein domain prediction and classification into families was done using InterPro (EMBL-EBI) and SMART (EMBL) databases. Signal peptides were predicted using the SignalP-4.0 Server (CBS).

Cloning, expression in *Escherichia coli* JM109 and purification

The putative *alginate lyase* genes, designated *alyRm3* and *alyRm4*, were amplified from the presented genome of strain MAT378 without the predicted signal peptide sequences (aa 1–17 and 1–22, respectively). For heterologous expression in *E. coli*, all the alginate lyases genes were modified with an N-terminal hexa-histidine tag. Primers were designed as shown in Supplementary Table 1 to amplify the coding regions of the respective genes and introducing the restriction sites *Bam* and further grown for 4 h at 30 °C till cell density reached OD600 of 0.6 then induced by adding of 0.1% rhamnose and further grown for 4 h at 30°C. The cells were harvested by centrifugation at 4,500 × *g* for 20 min at 4°C, washed, resuspended in 10 mM potassium phosphate buffer, pH 6.5 and disrupted by passing them twice through a French press cell. After centrifugation (13,000 × *g* for 15 min at 4°C), the supernatants of the crude cell extracts and the cell pellets were analyzed by SDS-PAGE.

The purifications of recombinant alginate lyases proteins were performed by immobilized metal affinity chromatography (IMAC). The supernatant of the respective crude cell extract, containing approximately 25 mg *E. coli* protein, was applied onto 2 mL Talon<sup>®</sup> metal affinity resin (Clontech) in a column using gravity flow. The resin was washed with 10 mL of washing buffer (50 mM potassium phosphate, 300 mM NaCl, 5 mM imidazole, pH 7.0). Bound protein was eluted with 3 mL of elution buffer (50 mM potassium phosphate, 300 mM NaCl, 150 mM imidazole, pH 7.0). (His)6-alginate lyase-containing fractions were combined and applied onto an NAP10 column (GE Healthcare), equilibrated with 50 mM potassium phosphate, 300 mM NaCl, pH 7.0 to remove imidazole and stored at 4°C.

Protein determination and protein electrophoresis

Protein concentration was determined using Bradford reagent (Bio-Rad) and BSA standards for preparation of a standard curve. Sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE) was performed using the method of Laemmli (1970), standard gels and a protein standard (Fermentas). Gels were stained using Coomassie Brilliant Blue R-250 (Sigma).

Alginate substrate material

Sodium M- and G-block alginates with a broad range of degrees of polymerization (DP) [M-block (ALG500), averaged DP 20, M/G ratio > 4; G-block (ALG300), averaged DP 20, G/M ratio > 3] were purchased from Elicityl Oligotech<sup>®</sup> (Crolles, France). Low-viscosity sodium alginate from *Macrocystis pyrifera*, Kelp (Brookfield Viscosity 100–300 cps; 2% in H<sub>2</sub>O at 25°C) was purchased from Sigma-Aldrich (St. Louis, MO, United States).

Measurements of lyase activity of *AlyRm3* and *AlyRm4*

The temperature optimum and stability, pH optimum, and impact of NaCl concentration (salinity) were determined for each enzyme using the *M. pyrifera* alginate as substrate and applying the 3,5-dinitrosaliclycic acid (DNS) method for determination of reducing sugar (Miller, 1959). Generally, the reactions were run not more than 5 min of incubation time to minimize color formation from degradation products at temperatures above 50°C, which otherwise affected the results of the DNS assay. The reaction mixture consisted of 10 μL appropriate diluted purified enzyme, 80 μL 1.25% alginic acid and 10 μL 0.5 M potassium phosphate or acetate buffer (pH 5.5 and 6.5 for *AlyRm3* and *AlyRm4*, respectively) with a final concentration of 50 mM. The temperature optimum experiments were performed at a range from 25 to 100°C. The temperature stability was determined by estimating the temperature at which 50% enzyme activity was measured.
after 30 min of incubation (T1/2 temperature). I.e., the pure enzymes (1 µM) were incubated in 0.1 M potassium buffer, pH 5.5 and 6.5 for AlyRm3 and AlyRm4, respectively, during 30 min at temperatures over a range from 40 to 95°C. Following incubation, the enzyme was cooled, and the remaining activity measured at 60°C, applying DNS assay. The effect of pH on enzyme activity was tested with purified enzymes using various buffers at 0.5 M: sodium acetate (pH 4.5–5.5), sodium/potassium phosphate (pH 6.0–8.0), and Tris-HCl (pH 8.5–9.5) under standard DNS assay conditions. The effect of NaCl on the alginate lyase activity of the purified enzymes was tested in 0.1 M potassium phosphate buffer at pH 7.0 for AlyRm4 and at pH 5.5 for AlyRm3, including different concentration of NaCl. All reactions were done in triplicates. The percent activity was determined from the absorption units subtracting the blank.

Isolation and purification of product oligo-uronates

Preparative quantities of incubation mixtures [3 mL solutions, containing 300 µL (0.7 U/mL) enzyme (AlyRm3 or AlyRm4), 300 µL 0.5 M phosphate buffer, pH 5.5, and 2,400 µL alginate substrate (12.5 mg/mL; M-block, G-block, or M. pyrifera alginate), incubated for 24 h at 40°C] were fractionated by size-exclusion chromatography (SEC) on Bio-Gel P-2 (isolation of dimers), P-4 (isolation of trimers) or P-6 (isolation of tetramers) columns (90 × 1 cm; Bio-Rad, Richmond, CA). Elutions, with UV detection at 235 nm, were carried out with 0.2 M NH₄HCO₃ at room temperature and a flow rate of 12 mL/h, and 10-min fractions were collected and lyophilized. Depending on the oligomer series (DP) according to MALDI-TOF-MS, some fractions were pooled for further separation on adequate Bio-Gel columns.

Thin-layer chromatography

Carbohydrate samples were spotted in 1 cm lines on TLC sheets (Merck Kieselgel 60 F254, 20 × 20 cm), which were developed with 1-butanol: acetic acid: water = 2:1:1 (v/v/v). Bands were visualized by orcinol/sulfuric acid staining (10 min for 100°C).

High performance anion-exchange chromatography

The distribution of the different oligomers in M-block and G-block (1.5 mg/mL, 25 µL) were analyzed on a Dionex DX500 workstation (Dionex, Amsterdam, Netherlands), equipped with an ED40 pulsed amperometric detection (PAD) system. The separation of oligosaccharides was carried out on a Ionpac AS4A column (250 × 5 mm; Dionex) by using a linear gradient of 25–300 mM sodium acetate in 100 mM NaOH (1 mL/min).

Matrix-assisted laser-desorption ionization time-of-flight mass spectrometry

MALDI-TOF-MS experiments were performed using an Axima™ mass spectrometer (Shimadzu Kratos Inc., Manchester, United Kingdom), equipped with a nitrogen laser (337 nm, 3 ns pulse width). Negative-ion mode spectra were recorded using the reflector mode at a resolution of 5,000 Full Width at Half Maximum (FWHM) and delayed extraction (450 ns). The accelerating voltage was 19 kV with a grid voltage of 75.2%; the mirror voltage ratio was 1.12, and the acquisition mass range was 200–6,000 Da. Samples were prepared by mixing on the target 0.5 µL sample solutions with 0.5 µL aqueous 10% 2,5-dihydroxybenzoic acid as matrix solution.

Gas-liquid chromatography—Electron impact mass spectrometry

Reaction mixtures, obtained after incubation of M-block alginate, G-block alginate, and M. pyrifera alginate substrates with AlyRm4 were lyophilized, and after trimethylsilylation [pyridine: hexamethyldisilazane: trimethylchlorosilane = 5:1:1 (v/v/v), 30 min at room temperature] analyzed by GLC-EI-MS. Screenings were carried out on a GCMS-QP2010 Plus instrument (Shimadzu Kratos Inc.) using an EC-1 column (30 m × 0.25 mm; Alltech/Grace, Deerfield, IL) and a temperature program of 140–250°C at 8°C/min (Kamerling and Gerwig, 2007).

Nuclear magnetic resonance spectroscopy

Resolution-enhanced one-/two-dimensional (1D/2D) 500-MHz ¹H/¹³C NMR spectra were recorded in D₂O at pH 7.0 on a Varian Inova Spectrometer (NMR Center, University of Groningen) or on a Bruker DRX-500 Spectrometer (Bijvoet Center, Department of NMR Spectroscopy, Utrecht University) at probe temperatures of 300K or 334K. Before analysis, samples were exchanged twice in D₂O (99.9 atom% D, Cambridge Isotope Laboratories, Inc., Andover, MA) with intermediate lyophilization, and then dissolved in 0.6 mL D₂O. Suppression of the HOD signal using the Bruker instrument was achieved by applying a WEFT (water eliminated Fourier transform) pulse sequence for 1D experiments and by a pre-saturation of 1 s during the relaxation delay in 2D experiments. Standard “Presat” solvent signal suppression (WET1D pulse) was used in case of 1D experiments on the Varian instrument. The 2D
gCOSY and TOCSY spectra were recorded using an MLEV-17 mixing sequence with spin-lock times of 20-200 ms. Natural abundance 2D $^{13}$C-$^{1}$H HSQC experiments ($^{1}$H frequency 500.0821 MHz, $^{13}$C frequency 125.7552 MHz) were recorded without decoupling during acquisition of the $^{1}$H Free Induction Decay (FID). All spectra were processed using MestReNova 5.3 (Mestrelabs Research SL, Santiago de Compostella, Spain).

Chemical shifts (δ) are expressed in ppm by reference to internal acetone (δ 2.225 for $^{1}$H and δ 31.08 for $^{13}$C).

**Results**

**Alginic utilization locus in *Rhodothermus marinus***

An alginic polysaccharide utilization locus (Alg-PUL) was identified following sequencing, assembly and annotation of the *Rhodothermus marinus* strain MAT378 genome as described in section "Materials and methods". Structurally identical locus was identified in the reference strain, DSM4252, following publication of the genome sequence (Nolan et al., 2009). The locus contains three alginic lyase genes, designated *alyRm1* (PL6), *alyRm3* (PL39), and *alyRm4* (PL17) (Figure 2). An additional alginic lyase gene *alyRm2* (PL6) was identified outside the locus. The Alg-PUL also contained genes encoding catabolic enzymes, *kdgF* and *kdgR*, which activity has been verified (unpublished results), necessary for downstream funneling of final alginic degradation products into the Entner Doudoroff pathway (unpublished data) as well as *kdgA* encoding putative 4-hydroxy-2-oxoglutarate aldolase; *kdgK* encoding putative 2-dehydro-3-deoxygluconate kinase. Relevant transporter genes and four hypothetical genes were also detected in the locus (Figure 2). The corresponding Alg-PUL genome sequence has been submitted to GenBank with the accession number ON640824. The enzymes in *Rhodothermus marinus* DSM 4252 homologous to AlyRm4 and the AlyRm3 amino acid sequences have the GenBank numbers ACY48056.1 and ACY48059.1, respectively. Both have been annotated as Heparinase II/III family protein.

The predicted gene *alyRm3* ORF was 2,613 nt, encoding an 870 amino acid long polypeptide. Its calculated MW is 96,623 Da and pI 5.21. A signal sequence was predicted with cleavage site after Gln-17, a heparinase II/III-like protein domain (aa 374–518) and C-terminal associated with sorting to the outer membrane and covalent modification (Muth et al., 1998). A C-terminal sorting is also found in AlyRm1 PL6 alginic lyase encoded from same cluster indicating that both enzymes are extracellular. AlyRm3 can be assigned to a newly described polysaccharide lyase enzymes in the CAZY database, i.e., family PL39 (Ji et al., 2019). Closest relatives are found various lineages of the phylum of *Rhodothermaeota*, homologs having 50–100% aa identity depending on source species relatedness. AlyRm3 homologs are found in other *R. marinus* strains, e.g., the type strain DSM 4252. It is not found in *Rhodothermus bifrosti*, isolated from a terrestrial hot spring. The highest sequence similarity of AlyRm3, outside the genus was found with a heparinase II/III family protein from e.g., *Rhodocaloribacter litoris* (66% aa identity, GenBank: WP_166976759.1). Homologs are found in other phyla albeit with distinctly lower sequence similarity including a homolog from the clostridia species *Defluviitalea saccharophila* with 38% amino acid identity but with similar activity on alginic.

The predicted gene *alyRm4*, encoding AlyRm4, consists of 2,226 nt, which translates into a 742 aa polypeptide with MW of 83,633 Da and pI 6.06. A hydrophobic sequence was detected at the N-terminal, but the C-terminal sorting domain is missing indicating that the enzyme is located in the periplasmic space. Like AlyRm3, the enzyme contains a heparinase II/III-like protein domain (aa 386–539). AlyRm4 belongs to family PL17 of polysaccharide lyases and shows highest identity to homologous proteins in *Rhodothermaeota* and various species from other phyla including e.g., *Rhodocaloribacter litoris* WP166976762, 68% identity and *Vibroniales* bacterium SWAT-3 (45% aa identity, GenBank: ZP_01815528.1). Other thermophilic strains containing a PL17 sequence are *Rhodocaloribacter litoris*, *Spirochaeta thermophila*, *Merioribacter roseus*, and the *Paenibacillus* strain Y412MC10.

**Production and purification of the recombinant AlyRm3 and AlyRm4 enzymes**

The encoding sequences of the mature enzymes were amplified by PCR and inserted into the 1-rhamnose inducible expression vector pJOE5751 with a N-terminal fusion to a hexa-histidine sequence allowing production of His6-fusion proteins. Rhamnose induction of the *E. coli* JM109, harboring the respective plasmids, resulted in the production of high amounts of the recombinant proteins as identified by SDS-PAGE (Supplementary Figure 1). Compared to the uninduced crude extracts, a prominent protein band of the size of ~95 kDa for AlyRm3 and ~81 kDa for AlyRm4, respectively, indicated a tight regulation and high expression level. Recombinantly expressed alginic lyases were purified by IMAC to homogeneity as judged by SDS-PAGE (Supplementary Figure 1).

**Characterization of the recombinant alginic lyases**

The recombinant alginic lyases AlyRm3 and AlyRm4 were active on alginic acid substrates but not on other substrates tested such as chondroitin and chitin (data not shown). The optimal pH, temperature optimum of activity and temperature stability are summarized in Table 1.
The activity was determined applying DNS assay.

Substrate specificity of AlyRm3

In order to determine the substrate-depolymerizing specificity of the AlyRm3 enzyme, incubation experiments (0–24 h, pH 5.5, 40°C) were performed with M-block alginate, G-block alginate, and low-viscosity M. pyrifera alginate as substrates. Calculated by the peak integrals of specific protons in the 1H NMR spectrum, the low-viscosity M. pyrifera alginate contained an M:G ratio of 61:39. The enzymatic depolymerization reactions were followed by TLC, fractionations on Bio-Gel P-2, P-4, and P-6, respectively. The enzymatic depolymerization reactions were followed by TLC, MALDI-TOF-MS, and 1H NMR spectroscopy. As shown in the TLC analysis of the 24-h incubations (Figure 3), the three different substrates were depolymerized into mainly di-, tri-, and tetracarboxydiacids with a 4,5-unsaturated non-reducing-end uronic acid (Δ), indicating that the enzyme can cleave M-M, G-G, M-G, and G-M linkages via a β-elimination mechanism, which was confirmed by 1H NMR spectroscopy of isolated products (see below). Note that the starting materials have completely disappeared. The negative-ion mode MALDI-TOF mass spectra of the three oligosaccharide mixtures (spectra not shown), obtained after a 24-h incubation, showed [M-H]− peaks at m/z 351.0, 526.8, and 703.0, corresponding to di-, tri-, and tetracarboxydiacids, respectively, each with a 4,5-unsaturated non-reducing-end uronic acid (Δ). Furthermore, weak [M-H]− peaks at m/z 369.2, 544.9, and 721.0 were also observed, in agreement with the presence of saturated di-, tri-, and tetracarboxydiacids, respectively. In the case of the G-block alginate incubation, additional small [M-H]− peaks at m/z 879.0, 897.0, and 1073.3 were seen, corresponding with ΔGGGGG, GGGGG, and GGGGG, respectively.

To facilitate the NMR analysis of the oligosaccharide mixtures obtained after a 24-h incubation, generated unsaturated di-, tri-, and tetracarboxydiacids were isolated via fractionations on Bio-Gel P-2, P-4, and P-6, respectively. The 1D 1H NMR spectra of ΔM, ΔMM, ΔMΔΔ, ΔG, and ΔGG are presented in Supplementary Figure 4. For detailed 1H and 13C assignments, gCOSY, TOCSY (mixing times 20, 50, 150, and 200 ms), and HSQC experiments were carried out. As an example, the 2D TOCSY (150 ms) and HSQC spectra of ΔM are presented in Supplementary Figure 5. The 1H/13C NMR chemical shift data of ΔG, ΔGG, ΔM, and ΔMM have been collected in Table 2. Note the specific Δ H-1 and Δ H-4 chemical shifts in case of Δ-M-R and Δ-G-R (R = M- and/or G-containing saccharide). Furthermore, it is of importance to note that the pH (pD) can influence chemical shifts, in particular the H-5 proton. Besides our own generated NMR chemical shift database, also previously published NMR data on alginate poly- and oligosaccharides were taken into account for resonance assignments (Grasdalen et al., 1979, 1981;...
Grasdalen, 1983; Steginsky et al., 1992; Heyraud et al., 1996; McIntyre et al., 1996; Zhang et al., 2004; Holtan et al., 2006; Li et al., 2011; Lundqvist et al., 2012; Hu et al., 2013).

The 500-MHz $^1$H NMR spectra at $t = 0$ h and $t = 24$ h of the M-block alginate incubation are presented in Figure 4, and those of the G-block alginate incubation in Figure 5. The average DPs ($\sim$DPs) of the commercial M- and G-block alginites were checked by $^1$H NMR spectroscopy via integration (I) of the anomeric $^1$H signals in the respective spectra ($\sim$DPM$_{-}$block = ($I_{M1} + I_{Mred1}$))$/I_{Mred1}$ and $\sim$DPG$_{-}$block = ($I_{G1} + I_{Gred1}$))/$I_{Gred1}$, yielding $\sim$DPM$_{-}$block 22 and $\sim$DPG$_{-}$block 25 (Figures 4A, 5A). The HPAEC-PAD profiles presented in Supplementary Figure 6, give an impression of the degree of polymerization or chain length distribution in M-block and G-block alginate. As is evident from the changes in the NMR profiles, going from Figure 4A to Figure 4B and from Figure 5A to Figure 5B, the enzyme is clearly acting as a lyase for both poly-M and poly-G homopolymeric regions. The formation of AM from the M-block alginate is demonstrated by the characteristic $\Delta$H-4 resonances at $\delta$ 5.82 (coded $\Delta$M4Ma) and $\delta$ 5.77 (coded $\Delta$M4Mb) (ratio $\sim$ 3:1), whereas the intense signal at $\delta$ 5.76 reflects the presence of AM(M)$_{n}$ (Figure 4B and Table 2; compare Supplementary Figure 4). Note that the MALDI-TOF-MS analysis showed mainly AM, AMM, and AMMM as end products (see above). For the G-block alginate, the appearance of $\Delta$H-4 signals at $\delta$ 5.92 (coded $\Delta$G4G) and $\delta$ 5.86 (coded $\Delta$G4GG) indicated the presence of AG and AGG(G)$_{n}$, respectively (Figure 5B and Table 2; compare Supplementary Figure 4). Remarkably, the reducing end residue G (coded Gred) has mainly $\beta$ configuration, while the reducing end residue M (coded Mred) showed a configuration ratio of $\alpha$/$\beta = 2.1$. Noteworthy, MALDI-TOF-MS analysis showed mainly AG, AGG, and AGGG as end products.

Making use of the NMR data obtained above for the M- and G-block alginate/oligosaccharide, the depolymerization of the low-viscosity M. pyrifera with the AlyRm3 enzyme was studied in more detail. The $^1$H NMR spectrum of the starting material, recorded at 334K instead of 300K to increase solubility and sharper resonance peak recovery (Figure 6A), showed in the $\delta$ 4.4-5.1 region the anameric proton signals. The GulpA H-1 signals resonate at $\delta$ $\sim$5.03 [G or M]G1G + [G or M]G1M and the ManpA H-1 signals at $\delta$ 4.70 [M1G] and $\delta$ 4.66 (M1M). The GulpA H-5 signals are present at $\delta$ 4.47 (G2G2G) with a shoulder at the right side for M2G2G ($\delta$ 4.43) and at $\delta$ 4.73 (M2G2M) with a shoulder at the left side for G2G5M ($\delta$ 4.76). Note that a preceding G residue (G2G2G vs. M2G2G and G2G2M vs. M2G2M) leads to a slightly higher chemical shift for G5-M and that M1-G in the M-M diad resonates at a slightly higher chemical shift than in the M-M diad (Grasdalen et al., 1979; Grasdalen, 1983). The ManpA H-5 signals resonate in the bulk region at $\delta$ $\sim$3.75 (compare Figure 4). The M/G ratio was calculated from the relative integral areas (I) in the anameric region, as described by Grasdalen et al. (1979), Grasdalen (1983) and Zhang et al. (2004). Here, the integration limits applied in the calculations of the area of the peak regions A, B and C were set to 4.97–5.10 ppm, 4.60–4.80 ppm, and 4.40–4.50 ppm, respectively (Figure 6A). Calculation of the percentage G and M monads in the M. pyrifera alginate, using the formula $F_{G} = I_{G}/I_{M} + I_{C}$, resulted in 39% G and 61% M, and therefore a M/G ratio of 1.6. In a similar way, the percentage GG diads followed from the formula $F_{GG} = I_{C}/I_{M} + I_{C}$, yielding 20% GG. As $F_{G} = F_{CG} + F_{GM}$, $F_{M} = F_{MM} + F_{MG}$, and $F_{GM} = F_{MG}$, the other diads comprise 19% GM, 19% MG, and 42% MM.

The oligosaccharide mixture obtained after a 24-h incubation of the M. pyrifera alginate with the AlyRm3 enzyme was subjected to NMR analyses, and the assignments of resonance peaks were obtained by 2D TOCSY and HSQC experiments. The 1D $^1$H NMR spectrum (Figure 6B) showed anameric signals for both reducing-end G and M residues. The Gred1$\beta$ resonance at $\delta$ 4.88 ($J_{1,2} = 8.5$ Hz) reflects a lyase activity of AlyRm3 on GM and/or GG diads, whereas
the Mred1α and Mred1β resonances at δ 5.25 and δ 4.93, respectively, demonstrate such an activity on MM and/or MG diads (note that the Mred1α signal at δ 5.25 overlapped the very minor Gred1α resonance) (Table 2). Non-reducing-end Δ residues, formed via β-elimination from both M and G residues during the lyase cleavage reaction, were traced by their H-4 signals, thereby giving information about neighboring uronic acid residues. The Δ H-4 signals detected at δ 5.92 (ΔAG) and δ 5.86 (ΔAGG) indicate the presence of ΔG and ΔGGR-type oligosaccharides, respectively, whereas the Δ H-4 signals seen at δ 5.82/5.77 (ΔAMa/ΔAMβ) and δ 5.76 (ΔAMM) revealed the presence of ΔM and ΔMRR-type oligosaccharides, respectively (R = M- and/or G-containing saccharide) (Table 2). The presence of ΔAGM (δ 5.87) and ΔAMG (δ 5.78) signals (Heyraud et al., 1996) could not be positively observed, due to partial signal overlap in the δ 5.75–5.95 region. Furthermore, the G H-5 signals (δ ~4.75) in the 1H NMR spectrum of the alginate substrate stemming from RMG5MR and RG5G5MR parts (Figure 6A) have disappeared in the 24-h incubation mixture (Figure 6B). Therefore, the signal at δ 5.04 belongs to H-1 of internal G residues (G1). The anomic protons stemming from M residues next to the reducing end (e.g., ΔMM and ΔMMM) are observed at δ 4.70 (M1MRedα) and δ 4.65 (M1MRedβ), due to the α/β configuration of the reducing-end M residue. TLC (Figure 3) and MALDI-TOF-MS analyses support the presence of ΔM, ΔMM, ΔMMM, ΔG, ΔGG, and ΔGGG in the product mixture after the 24-h incubation of M. pyrifera alginate with AlyRm3.

The Δ, G and M distribution (F) in the AlyRm3 lyase-degraded M. pyrifera alginate (original distribution: 39% G and 61% M; see above) was calculated by integration (I) of specific NMR peaks, using the following equations: $F_\text{G} = \frac{I_{\text{mm}} + I_{\text{mm\theta}} + I_{\text{gg\theta}}}{I_{\text{mm}} + I_{\text{mm\theta}} + I_{\text{gg\theta}} + I_{\text{mm\beta}} + I_{\text{mm\beta\theta}}}$ and $F_\text{M} = \frac{I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}}}{I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}}}$ and $F_\text{G} = \frac{I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}}}{I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta}} + I_{\text{mm\beta\theta}} + I_{\text{mm\beta\theta}}}$, respectively. This resulted in 49% Δ, 9% G, and 42% M.

The larger decrease of G residues (39% → 9%) compared to the decrease of M residues (61% → 42%) might indicate a higher conversion of G units into non-reducing Δ residues. This could mean a preferential cleavage between RM-GR and/or RG-GR (R = M- and/or G-containing saccharide) by the AlyRm3 enzyme, despite the majority of M-M bonds in M. pyrifera alginate. The integrals of the Mred1α/β signals (δ 5.25/4.93) predominated over those of the Gred1α/β signals (δ 5.23/4.88), indicating a preferred splitting of the RM-GMR bonds over the RG-GMR bonds. Moreover, the integrals ratio 4:1 of (ΔAMa + ΔAMβ + ΔAMM) and (ΔAG + ΔAGG) supported the preferential cleavage between RM-GMR over RM-GGR, and eventually RG-GMR over RM-GGR (R = M- and/or G-containing saccharide). However, a higher rate of cleavage of RM-GMR bonds (yielding ΔG at the non-reducing ends and Mred at the reducing ends) is also possible, in view of the low presence of only 20% GG in M. pyrifera alginate (see above). Nevertheless, the results from the incubation of the M-block alginate (~DP 22) with AlyRm3 (Figure 4) showed the enzyme's capability of cleaving M-M bonds (yielding oligosaccharides having AM at the non-reducing ends and Mred at the reducing ends). The average degree of polymerization (DPn) of the products from the AlyRm3-degraded alginate
 substrate was estimated to be 2.3, using the following equation: 

\[ \text{DP}_n = \frac{I_{11G} + I_{11M} + I_{M1} + I_{G1} + I_{M_{\text{red}}} + I_{G_{\text{red}}}}{I_{11G} + I_{11M} + I_{M_{\text{red}}} + I_{G_{\text{red}}}} \]  

(Ertesvåg et al., 1998). This is in reasonable agreement with the products, ranging from disaccharides to tetrasaccharides, as observed by TLC and MALDI-TOF-MS analysis.

The different observations indicated that within 24-h AlyRm3 can completely depolymerize M-block alginate (\( \sim \text{DP 22} \)) into mainly \( \Delta M \), \( \Delta MM \), and \( \Delta MMM \) (di-, tri-, and tetrasaccharides) and G-block alginate (\( \sim \text{DP 25} \)) into \( \Delta G \), \( \Delta GG \), and \( \Delta GGG \) (di-, tri-, and tetra-saccharides). So, for both M- and G-block alginates the high content of di-, tri-, and tetrascarhides seems to indicate that the smallest substrate of the AlyRm3 is a tetrasaccharide, acting as a minimal recognition pattern. As deduced from the results of the incubations of AlyRm3 with \( M. \ pyrifera \) alginate, for alternating GMGM blocks, depolymerization mainly occurs into \( \Delta GR \) due to the preferential cleaving of the RM-GMR...
bond \((R = \text{GGMG-saccharide})\). So far, the following degradation pattern of alginic acid by AlyRm3 lyase can be proposed:

...M\(\downarrow\)GMM\(\downarrow\)GG\(\downarrow\)GGG\(\downarrow\)GG\(\downarrow\)GM\(\downarrow\)G\(\downarrow\)GM\(\downarrow\)GMM\(\downarrow\)M...

...M, \(\Delta\)MM, \(\Delta\)G, \(\Delta\)GG, \(\Delta\)GGG, \(\Delta\)M, \(\Delta\)MM, \(\Delta\)MMM, \(\Delta\)M, \(\Delta\)M, \(\Delta\)MM, \(\Delta\)...

**Substrate specificity of AlyRm4**

In a similar way, as described for the AlyRm3 enzyme, for the determination of the substrate specificity of the AlyRm4 enzyme, incubation experiments (0–24 h, pH 5.5, 40°C) were performed with M-block alginate (\(~\text{DP 22}\) ), G-block alginate (\(~\text{DP 25}\) ), and low-viscosity *M. pyrifera* alginate (M:G ratio of...
FIGURE 6
1D $^1$H NMR spectra of (A) (334K, D$_2$O) M. pyrifera alginate substrate before addition of the AlyRm3 enzyme, and (B) (300K, D$_2$O) the oligosaccharide mixture generated after a 24-h incubation of the alginate with the AlyRm3 enzyme. For the coding system in panel (A), see text; for the coding system in panel (B), see Figures 4, 5. M$_{1}$M$_{red}^α$ and M$_{1}$M$_{red}^β$/M$_{1}$M mean M H-1 of internal M-residues.

61:39) as substrates. The enzymatic depolymerization reactions were followed by TLC and $^1$H NMR spectroscopy.

TLC analysis of the 24-h incubations revealed that the enzyme can almost completely depolymerize the M-block and M. pyrifera alginites into monomers, as shown by the intense decrease of the spot at the origin on the TLC plate (Figures 7A,C). The G-block alginate, however, was only partially degraded into monomers after 24-h, as demonstrated by the left-over spot at the origin on the TLC plate (Figure 7B). Interestingly, no intermediate oligosaccharide bands could be detected, suggesting an exclusive exolytic activity of AlyRm4 on the three substrates. The assignment of the spots (Figure 7) correlated with isolated and identified products (see below).

The product mixtures, obtained after the 24-h incubations of the M-block, G-block, and M. pyrifera alginites, were further studied by $^1$H NMR spectroscopy (Supplementary Figure 7). In all three cases, the typical signals reflecting the presence of 4,5-unsaturated uronic acids (e.g., the Δ H-4 resonances in the δ 5.7–5.9 region) were absent, indicating, when compared with the AlyRm3 product mixtures (see Figures 4–6), a different depolymerization mechanism exists for AlyRm4.
Inspection of the 1D 1H NMR spectra of the product mixture of the three alginates (Supplementary Figure 7) showed only minor signals of residual free or reducing M (Gred1α, δ 5.22; Gred1β, δ 4.90) and/or free or reducing G (Gred1α, δ 5.23; Gred1β, δ 4.88). However, in the case of the G-block alginate incubation, intense signals (G1, δ 5.03; G2, δ 3.90; G3, δ 3.99; G4, δ 4.10; G5, δ 4.45) of remaining polymeric/oligomeric material were visible (Supplementary Figure 7B), supporting the TLC indications by the residual spot at the origin (Figure 7). To generate information about the origin of the intense signals, coded A and B in each NMR spectrum, the product mixture of the M. pyrifera alginate incubation was fractionated on Bio-Gel P-2, yielding two fractions denoted I and II. Fraction I contained free ManpA (M) and free GulpA (G) (NMR analysis; spectrum not shown), which could be explained as derived from non-reducing ends of different DP chains of the alginate substrate. Fraction II was studied by 1D/2D NMR spectroscopy (Figure 8 and Supplementary Figure 8) and GLC-EI-MS (Supplementary Figure 9), showing that the fraction contained two saturated cyclic monomers identical with 4-deoxy-1-erythro-5-hexosulose uronic acid [2,4,5,6-tetrahydroxy-pentahydro-pyan-2-carboxylic acid / 1,2,3,5-tetrahydro-4H-pyran-5-carboxylic acid (TPC)]. As depicted in Figure 9, it is suggested that free A (4-deoxy-1-erythro-hex-4-enoxyranosyluronic acid), released from alginic acid via an exolytic AlyRm4-catalyzed β-elimination reaction, is converted into its open chain form, which is further hydrated in a spontaneous process in aqueous solution to form the cyclic hemiacetal stereoisomers (Enquist-Newman et al., 2014). Endolytic activity is excluded because oligomers with A in a non-reducing position were not detected, which means that AlyRm4 cleaves monomers one by one from the reducing end. In this context, it should be noted that many lyases catalyze the hydration of bound terminal A units, with subsequent rearrangements, resulting in glycosidic bond cleavage (Enquist-Newman et al., 2014; Hobbs et al., 2016). If the enzyme would have a reductase activity, then the released A is non-enzymatically converted into 4-deoxy-1-erythro-5-hexosulose uronic acid (DEHU), then reduced to 2-keto-3-deoxy-D-gluconate (KDГ), and further metabolized through the Entner-Doudoroff pathway (Preiss and Ashwell, 1962a,b). However, in our case, the hypothesis that AlyRm4 has reductase activity can be excluded. The proton at C4 is added in a later spontaneous process (keto-enol tautomerization) in solution.

Note that, in theory, any stereochemical combination at C1 and C5 can take place during the ring closure process, yielding four products. As is clear from the 1H NMR spectrum (Figure 8), only two sets of ring protons are seen in a peak ratio of 1:0.8, indicating the presence of two main cyclic structures A (H1: δ 5.01, J1,2 3.5 Hz) and B (H1: δ 4.98, J1,2 6.0 Hz) in equilibrium. The full 1H and 13C NMR data of both stereoisomers A and B are presented in Table 3, and the TOCSY (200 ms) and HSQC spectra in Supplementary Figure 8.

GLC-EI-MS analysis of the incubation mixture of the M. pyrifera alginate (24 h, 40°C), after direct lyophilization and trimethylsilylation, showed a major GLC peak (Rt 10.2 min) with a shoulder of which the EI-MS fragmentation patterns were in agreement with the trimethylsilylated (TMS) proposed six-membered structures A and B (MW 554 Da) (Supplementary Figure 9). Both stereoisomers could not be separated on the GLC column under the conditions used. The presence of phosphate buffer was reflected by the presence of the dominant GLC peak of trimethylsilylated phosphate [characteristic ions at m/z 314 (M) and 299 (M minus CH3)]. Surprisingly, a series of additional small peaks at the beginning of the gas chromatogram gave mass spectra that could be correlated with small organic acids products, such as glyceric acid, malic acid, (hydroxylated) glutaric acids and others (Supplementary Table 2 and Supplementary Figure 10). When the incubation of the M. pyrifera alginate with the AlyRm4 enzyme was performed at 65°C instead of 40°C, the GLC-EI-MS analysis revealed an increase of organic acid formation. It is suggested that the intermediate open chain 4-deoxy-1-erythro-hexosulose uronic acid (DEHU) in Figure 9 is the precursor of these non-enzymatic generation of organic acid products. The formation of high amounts of dicarboxylic acids was promoted by increasing the incubation temperature (data not shown). However, it still
remains unclear how these products are generated and further study is in progress.

Finally, the $^1$H NMR spectrum of the G-block alginate, after a 24-h incubation with AlyRm4 at pH 5.5 and 40°C, showed the uncomplete depolymerization of the G-block by a broad signal of anomeric protons of internal G residues (G1) at δ 5.03, together with minor signals at δ 5.23 (Gred1α) and δ 4.87 (Gred1β) of Ga/β of non-degraded
G-block alginate (Supplementary Figure 7B). Nevertheless, an exolytic activity of AlyRm4 is demonstrated by the presence of the signals belonging to the cyclic, saturated stereoisomers A and B, stemming from released reducing ends initially present at different length chains of the substrate (~DP25).

Taking together, the absence of Δ in all three incubations with AlyRm4 clearly demonstrates that the lyase activity occurs only from the reducing end. Even when the G-block alginate was not completely degraded by the AlyRm4 enzyme, unsaturated oligomers with a Δ unit at the non-reducing end (e.g., ΔGGG) were not detected. Evidently, the free unsaturated monomers, released via β-elimination, are immediately converted to the two cyclic saturated monomers (A and B, Figure 9) and, eventually, into small organic acids, depending on the incubation temperature. As observed so far, all substrates were degraded, suggesting that the AlyRm4 enzyme has a broad substrate tolerance and can cleave M-M, M-G, M-G and G-G bonds, with a higher specificity/depolymerizing rate for MM homopolymeric sequences than for GG homopolymeric sequences. From the obtained analysis data, it can be concluded that the AlyRm4 lyase completely depolymerizes alginic acid into unsaturated monomers, which are mostly converted into cyclic monomers A and B and small organic acids, under the used conditions at elevated temperature:

\[
G\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta
\]

\[
\downarrow
\]

\[
G\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta\Delta
\]

\[
\downarrow
\]

cyclic monomers A and B.

**Discussion**

Based on sequence similarities, alginate lyases have been assigned to the polysaccharide lyase families PL5, PL6, PL7, PL14, PL15, PL17, and PL18 \(^3\) (Garron and Cygler, 2010; Zhu and Yin, 2015; Mathieu et al., 2016; Drula et al., 2022). Most bacterial alginate lyases, according to their primary structure, belong to the polysaccharide lyase families PL5 and PL7, and their molecular masses range from 25 to > 60 kDa (Lombard et al., 2010). The enzymes are generally classified according to their dominant typical cleaving action as poly-M lyase [(1→4)-β-D-mannuronic lyase; EC 4.2.2.3; PL5 (PL15 and PL17)] and poly-G lyase [(1→4)-α-1-guluronan lyase; EC 4.2.2.11; PL7] (Wong et al., 2000; Yamasaki et al., 2005; Kim et al., 2011). Furthermore, their three-dimensional arrangements allow grouping them into three structural classes, displaying either an α/β helix-barrel fold (Yoon et al., 1999) or a β-sandwich (jelly-roll) fold (Osawa et al., 2005). Several alginate lyases have been cloned and characterized from various sources (Zhu and Yin, 2015).

In case of the alginate degrading thermostable bacterium *R. marinus* strain MAT378/R-21, the (recombinant) AlyRm4 enzyme belongs to the PL17 polysaccharide lyase family, whereas the (recombinant) AlyRm3 enzyme belongs to the recently defined family PL39 (Li et al., 2019).

Compared to other reported thermophilic alginate lyases, the recombinant AlyRm3 and AlyRm4 are highly thermostable, with T1/2, in absence of substrate, (50% activity after 30 min incubation in buffer) of 77°C and 78°C for AlyRm3 and AlyRm4, respectively. This makes them among the most thermostable alginate lyases that have been described to date, along with AMOR_PL17A, derived from an Arctic Mid-Ocean Ridge (AMOR) metagenomics data set (Arntzen et al., 2021).

Temperature and pH optimum, stability and optimum salinity determinations were carried out applying DNS assays, measuring the release of reducing sugars quantified as glucose equivalents. At higher temperatures (above 55°C), a dark brown color was formed in the reaction solutions and formation of dicarboxylic acids was detected. The DNS assay was adopted accordingly by limiting the time of the enzyme reaction. However, due to the color formation at high temperatures, measuring the specific enzyme activities close to temperature optimum applying DNS assay was considered imprecise, and not reported here.

The thermophilic AlyRm3 and AlyRm4 alginate lyases differ in depolymerizing activity, but together they cover a wide activity range. Our results showed that the endolytic AlyRm3 enzyme can cleave any of the four types of bonds (M-M, G-G,
M-G, and G-M), with a slight preference for M-M bonds, and completely degrade alginate into unsaturated di-, tri-, and tetrasaccharides. Interestingly, the exolytic AlyRm4 lyase appears to cleave any of the two reducing end types (G and M) from alginate chains, releasing the unsaturated monosaccharide 4-deoxy-L-erythro-5-hexosulose uronic acid (DEHU), Δ, which spontaneously converts into cyclic monomers. Increasing the incubation temperature resulted in the formation of carboxyl organic acid products (Aoyagi et al., 2008). It must be noted that poly-MG lyases, showing exolytic activity, are very rarely found in nature.

The enzymes, described in this report, can be used as biocatalyst for saccharification of alginate, since they can efficiently degrade polyM, polyG, polyMG blocks, alginate oligosaccharides, and alginate to produce specific molecules with defined properties optimally suited for a given application ("tailor-made alginate"). The enzymatic degradation of alginate can be either selective or complete depending on the choice of enzymes. For instance, the recombinant AlyRm3 converted alginate into unsaturated di-, tri- and tetrasaccharides, while AlyRm4 depolymerized ~95% of alginate into monomers. High efficiency of the reactions could be explained by the fact that the incubations were performed at elevated temperatures, where the solubility of alginate increases and viscosity decreases, facilitating enzymatic access. Robust alginate lyases, active in such extreme circumstances, may be of great interest for biofuel and chemical industries.

Conclusion

The two alginate lyases described in this report are highly thermostable. Detailed NMR analysis of the incubation products demonstrated that AlyRm3 is an endolytic lyase, while AlyRm4, as an exolytic lyase, cleaves monomers from the non-reducing end of oligo/poly-alginates. The two enzymes can be used separately for selective partial degradation of alginate, or in combination for complete degradation of alginate for production of monomer residues. Their application in industrial processes, where higher processing temperatures are required or preferred, is of substantial interest.

Data availability statement

The data presented in the study are deposited in the GenBank repository, accession number: ON640824.

Author contributions

All authors listed have made a substantial, direct, and intellectual contribution to the work, and approved it for publication.

Funding

This work was supported by EU FP7 grant 265992, AMYLOMICS; BBI grant MACROCASCADE 720755 and the BlueBio Cofund consortium grant, MARIKAT, Grant Icelandic Technology Development Fund 2011273-0611.

Conflict of interest

The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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Supplementary material

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/fpls.2022.981602/full#supplementary-material

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