

Functional exploration of Pseudoalteromonas atlantica as a source of hemicellulose-active enzymes: Evidence for a GH8 xylanase with unusual mode of action

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1	Functional exploration of Pseudoalteromonas atlantica as a source of hemicellulose-
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3	
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15	Running title: Pseudoalteromonas atlantica as a source of hemicellulose-degrading enzymes

17 Abstract

To address the need for efficient enzymes exhibiting novel activities towards cell wall 18 polysaccharides, the bacterium *Pseudoalteromonas atlantica* was selected based on the 19 20 presence of potential hemicellulases in its annotated genome. It was grown in the presence or 21 not of hemicelluloses and the culture filtrates were screened towards 42 polysaccharides. P. 22 atlantica showed appreciable diversity of enzymes active towards hemicelluloses from 23 Monocot and Dicot origin, in agreement with its genome annotation. After growth on beechwood glucuronoxylan and fractionation of the secretome, a β -xylosidase, a α -24 25 arabinofuranosidase and an acetylesterase activities were evidenced. A GH8 enzyme obtained in the same growth conditions was further cloned and heterologously overexpressed. It was 26 shown to be a xylanase active on heteroxylans from various sources. The detailed study of its 27 mode of action demonstrated that the oligosaccharides produced carried a long tail of un-28 29 substituted xylose residues on the reducing end. 30

Keywords: CAZymes; Glycoside-hydrolases; enzyme screening; hemicellulases; GH8 xylanase
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33 1- Introduction

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34 Hemicelluloses are among the most abundant polysaccharides present in the plant cell wall after cellulose. They play an important role in the regulation of cell wall mechanical 35 properties during plant growth, which impacts the various applications of plant crops quality 36 37 and their agro-industrial processes **[1]**. Hemicelluloses include several polysaccharides, such 38 as xyloglucans, heteroxylans, heteromannans, mix linked β -glucans [2]. Xyloglucans, 39 heteroxylans, heteromannans are made of (1.4) linked β -D-glucose, xylose and mannose, 40 respectively, on which several other sugars or non-sugar substituents can be linked. In 41 monocot endosperm, xylans carry (1,3)- and/or (1,2)-linked α -L-arabinofuranosyl units as the 42 main substituents to form arabinoxylans (AX) [3]. Heteromannans include linear mannan, 43 glucomannan (GlcM), galactomannan (GalM), and galactoglucomanan (GgM) [4]. Mix linked βglucans consist of a (1,4)-linked β -D-glucose backbone in which (1,3) linkages are 44 45 interspaced. A deeper knowledge on the fine structural details of hemicelluloses is required 46 not only to improve their degradation in biorefinery process, but also for a better 47 understanding of the impact of their versatile decorations on their functions in the cell wall. Degradation with carbohydrate active enzymes, followed by chemical, chromatographic and 48 49 spectrometric analyses is one of the leading tools to solve the fine structure of 50 polysaccharides [5-8]. However to address the structural complexity of hemicelluloses, there 51 is a need for enzymes exhibiting novel specificity. The CAZy database (http://www.cazy.org) 52 classified the Carbohydrate Active enZymes in structurally-related families [9]. To date the 53 database includes 156 families of glycoside hydrolases (GH), of which 43 are involved in the 54 degradation of hemicelluloses. 55 Due to the structural complexity of hemicelluloses, their enzymatic degradation requires a large panel of enzymes working in synergy. For all the hemicelluloses cited above, this 56 57 includes the enzymes able to split linkages in the backbone and enzymes specific for the side

chain residues. When focusing on backbone, splitting the β -(1,4)-linkage between two glucose

59 residues in XyG requires either broadly specific endo- β -1,4-glucanases (EC 3.2.1.4), belonging 60 to 13 different GH families (5, 6, 7, 8, 9, 12, 44, 45, 48, 51, 74 and 124), or highly specific endo-61 acting xyloglucanases (EC 3.2.1.151), classified in 6 GH families (5, 9, 12, 16, 44 and 62 74)(http://www.cazy.org). Complete degradation of heteroxylan requires the co-working of 63 endo- β -1,4-xylanase, β -xylosidase, α -glucuronidase, α -arabinofuranosidase and esterase **[10]**. 64 Most of the microbial β -1,4-xylanases (EC 3.2.1.8) are grouped into 7 GH families: GH5, 8, 30, 65 43, 51, where a bi-functional cellulase/xylanase stands, and GH10 and 11, where the most studied β -1,4-xylanases are gathered **[11–14]**. The endo- β -1,4-mannanases required for the 66 67 hydrolysis of heteromannan backbone are classified into the GH families 5, 26 and 113. 68 As part of a comprehensive screening of bacteria to highlight enzymatic activities towards a 69 large number of polysaccharides, *Pseudoalteromonas atlantica* was shown to produce an 70 interesting diversity of enzyme repertoire. Its genome contains several GH genes (Table 1) 71 but there is a lack of information on most of the corresponding proteins. Therefore it was 72 grown in the presence or not of different hemicelluloses to study the potential inducer effect 73 on the enzyme secretion. Its secretome was fractionated to identify the proteins involved in 74 the observed activities. One of these proteins annotated as belonging to GH8 family was 75 studied in more details.

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77

78 2- Material and methods

79 2-1 Culture conditions

The strain *Pseudoalteromonas atlantica* T6c (ATCC, USA) was grown in its optimum growth conditions (Table 1). A pre-culture was carried out by inoculating frozen cells in 10 mL of specific culture medium (Table 1) in a 100 mL Erlenmeyer flask. After 36 h incubation at the suitable temperature under shaking at 150 rpm in a New Brunswick incubator, 1 mL of the culture was inoculated in 50 mL of fresh medium and incubated for additional 8 h under 85 similar conditions. Then, 50 mL were inoculated in 1 L of fresh medium in 5 L shaker flask. After 36 h, the culture was centrifuged at 5,000 × g for 30 min at 4°C. Culture supernatant and 86 87 cell pellet were treated separately prior to the screening assay. The cell free culture 88 supernatant was filtered overnight on a 300 kDa membrane (PES, Millipore) in a 400 mL 89 Amicon system 8400 (Millipore). The filtrate was then diafiltered and concentrated to 20 mL 90 with 50 mM Tris-HCl buffer pH 8.5 containing a protease inhibitor cocktail (Roche, 91 Indianapolis, IN, USA), on a 10 kDa membrane (PES, Millipore) under compressed air (0.5 bar). 92 The cell pellet was washed twice with 500 mL of 50 mM Tris-HCl buffer pH 8.5 and re-93 suspended in 10 mL of the same buffer containing anti-protease mixture. Bacterial cells were 94 disrupted by Constant cell disruption system (Constant systems Ltd, Northants, UK) and the 95 membrane fragments were removed by centrifugation at 20,000 ×g for 60 min at 4°C. The 96 clarified lysate was diluted 10 times, filtered overnight on a 300 kDa membrane and finally 97 concentrated, diafiltered with 50 mM Tris-HCl buffer pH 8.5 on a 10 kDa membrane to a final 98 volume of 20 mL. Culture supernatant and cell lysate were further used for activity screening.

99

100 2-2 Preparation of polysaccharides for enzyme screening

101 The forty-two polysaccharides used as substrates for the screening are shown in Table 2. All 102 the polysaccharides were dissolved in deionized water (4 mg/mL). Solutions of commercial 103 polysaccharides were diafiltered on a 10 kDa membrane (PES, Millipore, Billerica, MA, USA) 104 using a 50 mL Amicon system 8050 (Millipore). Polysaccharides purified from apple and 105 tomato and belonging to the laboratory collection were directly dissolved in deionized water.

106

107 2-3 Screening assay

The 42 polysaccharide solutions (150 μL) and 5 controls containing water were dispensed
into a 96-well filter microplate (10 kDa, PES, Pall) with an equal volume (150 μL) of culture
supernatant or cell lysate from bacterial culture [17]. Polysaccharide controls were done by

incubating 150 µL of polysaccharide solutions with 150 µL 50 mM buffer solution (Tris-HCl,
pH 8.5 or acetate, pH 5.2) instead of culture filtrate or cell lysate. Microplates were sealed
with a plastic film and incubated overnight with shaking at 34°C. The incubation medium was
filtered on a multiscreen HTS vacuum manifold (MSVMHTS00, Millipore) connected to a highoutput vacuum pressure pump (Millipore) for 3 h under about 0.5 bar pressure.

116 For the colorimetric assay, 40 µL of each incubation filtrate were transferred to a microplate 117 (Dutscher, Brumath, France). Then, 200 µL of ferricyanide solution [18] were added to the 118 sample and the microplate was sealed with plastic film. The plate was heated at 95°C for 119 15 min in a thermocycler (GenAmp PCR system 2700, Applied Biosystems France, Villebon-120 sur-Yvette, France) and cooled to room temperature. The occurrence of reducing ends was 121 revealed by reading the absorbance at 420 nm of 200 µL of the samples in a microplate reader 122 (Nunc, Roskilde, Denmark) using a Wallac 1420 multilabel counter (PerkinElmer Life Sciences, 123 Boston, MA, USA). The amount of reducing ends was expressed in absorbance units. The 124 samples with absorbance higher than the corresponding bacterial filtrate or lysate alone 125 (control) minus 0.15 were considered as negative hits. The mean (X) and the standard 126 deviation (σ) of the optical density (OD) of these negative hits were calculated. When the OD 127 of the sample was lower than X- 2σ , the hit was considered as positive. The polysaccharides were thus classified according to four intervals based on the control X and its σ . 128

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130 *2-4 Time course of growth of* P. atlantica

Two pre-cultures of *P. atlantica* were carried out by inoculating frozen cells in 10 mL of sterilized bactomarine broth culture medium in a 100 mL Erlenmeyer flask with 12 h time lag. After 36 h incubation at 20°C with shaking at 180 rpm in a New Brunswick incubator, 1 mL of the culture was inoculated in 250 mL of fresh medium. The growth of the bacteria was followed by the absorbance of 1 mL culture at 600 nm at 1 h of time intervals. Similarly the time course of growth was followed using konjac GlcM or beechwood glucuronoxylan (GuX)added at 1 g/L in the culture medium.

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139 *2-5 Fractionation of the culture supernatant of* P. atlantica

Similar cultures were carried out at larger scale by inoculating 50 mL of pre-culture in 1 L offresh medium in 5 L shaker flask in the same conditions.

P. atlantica was cultivated in sterilized bactomarine broth without any added polysaccharide or in the presence of konjac GlcM or beechwood GuX at 1 g/L. After 16 h, 18 h and 16 h of respective growth for the three conditions, the culture was centrifuged at 5000 × g for 30 min at 4°C. The cell free supernatants were recovered and complemented with a protease inhibitor cocktail.

The culture supernatants were filtered overnight on a 300 kDa membrane (PES, Millipore) 147 148 using a 400 mL Amicon system 8400 (Millipore). The filtrates were diafiltered and 149 concentrated to 20 mL, with deionized water on a 10 kDa membrane (PES, Millipore) under 150 compressed air (0.5 bar) to remove small molecules. The concentrated filtrates were 151 separated into 2 x 10 mL and the pH of each half was adjusted at 8.5 or 5.2 by adding 100 mM 152 Tris-HCl or 100 mM acetate buffer, respectively. By this way, 6 culture filtrates were 153 obtained: without any added polysaccharide or with konjac GlcM or beechwood GuX, and for 154 each, at pH 8.5 and 5.2. All culture filtrates were stored at 4°C before enzyme assays.

The culture filtrate obtained in the presence of beechwood GuX was fractionated by ion exchange chromatography on a Mini QTM column (4.6 x 50 mm, GE Healthcare, Uppsala, Sweden) installed on an AKTA purifier (GE Healthcare) and equilibrated with 50 mM Tris-HCl buffer at pH 8.5. Elution was performed at a flow rate of 0.8 mL/min with 50 mM Tris-HCl buffer pH 8.5 (7 column volumes) to recover the unbound fraction. Bound proteins were eluted by a gradient on 10 column volumes from 35 to 75% of NaCl 1 M in 50 mM Tris-HCl buffer pH 8.5. The fractions of interest were dialyzed in a 10 kDa cut off dialysis tube 162 (Nanosep 10K Omega, Pall Life Science, USA) against 50 mM Tris-HCl buffer at pH 8.5,
163 adjusted to a final volume of 3 mL and stored at 4 °C.

164

165 *2-6 Enzymatic activities in* P. alteromonas *culture filtrates and after anion* 166 *exchange chromatography*

167 Two hundred μL of polysaccharide solutions were incubated overnight with 200 μL of 168 bacterial culture filtrates or protein fractions eluted from ion exchange chromatography. After 169 withdrawing 100 μL aliquot, the amount of reducing ends produced in the rest of the reaction 170 medium was quantified by Nelson method **[19]** adapted to microplate **[20]**, and read on a 171 microplate reader (Multiskan Ex). The amount of reducing ends was expressed in μg/mL 172 using standard curve prepared with appropriate sugar. Controls were prepared similarly with 173 previously heat-inhibited culture filtrates and polysaccharide solutions.

Glycosidase activities were assayed by incubating 100 μL culture filtrates with 100 μL 4 mM para-nitrophenyl (*p*-NP) derivatives of α-L-arabinofuranoside, α-D-galactopyranoside, β-Dmannopyranoside, β-D-glucopyranoside, α-D-xylopyranoside, β-D-galactopyranoside, α-Lfucopyranoside, acetate, and *ortho*-nitrophenyl (*o*-NP) β-D-xylopyranoside for 16 h at 40°C. The incubation was stopped by adding 0.6 mL of 1 M sodium carbonate solution **[21]**. The liberation of *p*-nitrophenol was reported as nmol/mL. Control tests were prepared by mixing the culture filtrate with sodium carbonate before adding the substrate.

181

182 *2-7 Protein analysis in* P. alteromonas *culture filtrates*

183 The concentration of protein was determined in the culture filtrate by using the Bradford

184 reagent (Bio-Rad, Marnes-la-Coquette, France) and bovine serum albumin (Sigma-Aldrich,

185 L'Isle d'Abeau, France) as a standard $(0-25 \mu g/mL)$ [22].

186 *Electrophoresis*: polyacrylamide gel electrophoresis was performed under denaturing

- 187 conditions (sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE)). It was
- 188 carried out in a MiniProtean 3 apparatus (BioRad) using a continuous 10–20%

189 polyacrylamide gel in the presence of 0.1% sodium dodecyl sulfate.

- 190 *In-gel digestion*: Bands of interest were excised from the SDS-PAGE and subjected to in-gel
- reduction, alkylation and trypsic digestion essentially as described [23].
- 192 *LC–MS/MS analysis*: Mass spectrometry analyses were conducted by the platform
- 193 "Biopolymers-Interaction-Structural Biology" located at the INRA Center of Angers-Nantes
- 194 (www.bibs.inra.fr). Nanoscale capillary liquid chromatography-tandem mass spectrometry
- 195 (LC-MS/MS) analyses of the digested proteins were performed using an Ultimate U3000 RSLC
- 196 system (Dionex) coupled with an LTQ-Orbitrap VELOS mass spectrometer (Thermo Fisher).
- 197 Details of the analytical procedure are given in **[23]**.
- 198 *Databank search and protein identification:* The LC-MS/MS collected data files were processed
- 199 into mgf format using Proteome Discoverer (Thermo-Fisher). The mgf files were then 200 searched against the Uniprot databank (http://www.uniprot.org, November 2014) restricted 201 to the taxonomy Pseudoalteromonas atlantica T6c (Taxon ID# 342610) using the Mascot 2.2 202 program (Matrix Science). Enzymatic cleavage was declared as a trypsic digestion with one 203 possible missed cut event. The fixed modifications of Cys residues by iodoacetamide and the 204 possible oxidation of Met residues were considered. Precursor mass and fragment mass 205 tolerance were set at 5 ppm and 0.5 Da, respectively. Proteins were validated when they had 206 an e-value below 0.01, and when they were identified with a minimum of two peptides of 207 individual score above the significance threshold provided by Mascot (p < 0.01).
- 208

209 2-8 Cloning and expression and purification of rPaGH8

- 210 The gene *Patl_1069* (GenBank accession number ABG39595.1), coding for the glycosyl
- 211 hydrolase of *P. atlantica* T6c grouped in the GH8 CAZy family was amplified from genomic
- 212 DNA using the following primers:

213 5-GTGAGCATTGATCACTTAACGCTAACAAAACGCGGG-3' (forward) and

5'-GCGCGGCCGCTTCAGGCTCGTTTTCATTTGGCTTATAG-3' (reverse) encompassing BamH1
and EcoR1 restriction site respectively. The gene was cloned in pET28a which contains an Nterminal fused six-histidine-tag (6His-tag) using BamH1 and EcoR1 restriction sites (pET28a-

GH8 plasmid) and transformed in *E. coli* strain BL21 (DE3). *E. coli* BL21 (DE3) harbouring the

218 pET28-GH8 plasmid was pre-cultured overnight in Luria-Bertani medium containing 50

µg/mL of kanamycin before being diluted 20 times in the same growth medium and incubated
under shaking at 20°C overnight.

221 Cells were harvested by centrifugation, washed with 50mM Tris-HCl NaCl 100 mM (pH 7.5),

and resuspended in one-tenth of the culture volume of cold Tris-HCl 50 mM pH7.5, NaCl 100

223 mM, imidazole 20 mM (pH 7.5) containing protease inhibitor (Pierce[™] Protease Inhibitor

224 Mini Tablets, ThermoFisher, Illkirch, France), 5 µL DNAse (Invitrogen, ThermoFisher, Illkirch,

France) and 20µg/mL RNAse (Invitrogen). Cells were disrupted using a French press at 2000
bars twice for 1 min, centrifuged 10 min at 4°C to recover a supernatant and a pellet.

227 The supernatant was loaded on a HisTrap[™] FF column (1 mL) mounted on a ÄKTA Primer 228 system (GE Healthcare) to purify the His-tag GH8 protein. The recombinant protein was 229 eluted with 20 mM Tris-HCl buffer pH 8 containing 200 mM NaCl, 300 mM imidazole. The 230 supernatant and pellet from the French press lysate, and the fractions eluted from the 231 HisTrap column were subjected to a polyacrylamide gel electrophoresis under denaturing 232 conditions (sodium dodecyl sulfate polyacrylamide gel electrophoresis (SDS-PAGE)), together 233 with a PageRuler[™] Unstained Protein Ladder (Fisher). SDS-PAGE was carried out into a 234 MiniProtean 3 apparatus (BioRad) using a continuous 10% w/v polyacrylamide gel in the 235 presence of 0.1% sodium dodecyl sulfate that was further stained with Coomassie-HCL [24]. 236 The fractions containing the purified rPaGH8 were pooled, dialyzed against Tris-HCl 50 mM 237 pH 8, and concentrated twenty times using a VIVASPIN 500 concentrator (PES membrane 10 238 kDa, Sartorius).

240 *2-9 Analysis of the oligosaccharides produced by rPaGH8*

241 The recombinant xylanase rPaGH8 was incubated overnight with xylan from Palmaria 242 *palmata* [25], beechwood GuX, water-extractable (WEAX) and water un-extractable 243 arabinoxylan (WUAX) from wheat. The results were compared with the hydrolysis by the 244 commercial GH11 xylanase M1 from *Trichoderma viride* (Megazyme, Wicklow, Ireland). 245 Reaction media aliquots were withdrawn and precipitated in 4 volumes of ethanol 96° at 246 +4°C. After centrifugation 5 min at 20,000 g and evaporation under vacuum, samples were 247 dissolved in 500 µL water and filtrated on 0.45 µm membrane. They were analysed by HPAEC 248 on a CarboPac PA 200 column (3 x 250 mm, Dionex, Sunnyvale, USA) thermostated at 25°C. 249 Samples were eluted at 0.4 mL/min with a linear gradient of sodium acetate from 0 to 170 250 mM in 100 mM NaOH **[26]**. The nomenclature for AX oligosaccharides is according to **[27]**: X: 251 D-Xylp (xylose in the β -(1-4)-linked backbone); A2: α -L-Araf-(1-2)- β -D-Xylp; A3: α -L-Araf-(1-3)- β -D-Xylp; A2,3: α -L-Araf-(1-2)-[α -L-Araf-(1-3)]- β -D-Xylp. As an example, XA3X describes a 252 253 main chain of 3 xylose residues of which the middle one is substituted at 0-3 by an arabinose 254 residue.

255 Unknown peaks were purified on the same Carbopac PA 200 column equipped with on-line 256 desalting by the Carbohydrate Membrane Desalter (CMD300, Dionex) to remove sodium ions. 257 The CMD unit was fed with dilute sulfuric acid (75 mM) at 3 mL/min using an AXP pump 258 (Dionex). The separated unknown oligosaccharides were collected (200 μ L) and further 259 analysed by electrospray ionization mass spectrometry. ESI-MS experiments were performed 260 on a Synapt G2Si high-definition mass spectrometer (Waters Corp., Manchester, UK). Two 261 types of mass measurements were performed on the samples: firstly, a mass profile was done 262 on a mass range of 400-2000 m/z. Ions of interest were further fragmented by collision-263 induced dissociation in the transfer cell of the instrument, using an appropriate collision 264 energy to obtain numerous fragments. Samples were mixed volume/volume with acetonitrile

265 and infused at 5 μ L/min in the instrument. The instrument was operated in negative

ionization mode in the so-called 'sensitivity' mode, with an ESI capillary voltage of 2.2 kV and
a sampling cone voltage of 60 V. Data acquisition was carried out using MassLynx software

268 (V4.1).

- 269
- 270
- 271 **3- Results**

3-1 Enzymatic screening after cultivation in the presence or not of polysaccharides 272 273 To highlight enzymes active on various polysaccharides, a preliminary screening was 274 conducted on four bacteria from marine and terrestrial origin whose genome carried at least 275 10 sequences of hemicelluloses-degrading enzymes. As a larger number of sequences did not 276 necessarily give more positive responses for substrate degradation, *P. atlantica* was selected 277 from this screening since it offered the best performance and the greatest diversity on 278 polysaccharides. Its genome contains genes coding for GH enzymes specific for hemicellulose 279 backbone, *i.e.* β-glucanase, xyloglucanase, β-mannanase and β-xylanase (GH3, 5, 8, 10, 16 and 280 43, Table 1).

281 P. atlantica was first grown in the basal medium recommended (Table 1) and in the absence 282 of added polysaccharide. Forty-two terrestrial or marine polysaccharides of diverse origins 283 and structures (Table 2) were tested as substrates to screen enzymatic activities in the 284 culture supernatant and in the cell lysate (Table 3). By comparing the results, it was obvious 285 that the culture filtrate was more active and more diverse than the cell lysate. The culture 286 filtrate of *P. atlantica* was highly, moderately or slightly active on 15 substrates among the 42 287 tested, including 10 hemicellulose-like substrates, while the cell lysate was highly active on 288 only 2 substrates and slightly active on 4 substrates.

289 To enhance the enzyme production, different hemicelluloses were added to the culture

290 medium, as the addition of polysaccharides can play an important role in the induction of

291 enzymatic activity **[28-31]**. The polysaccharides tested were chosen with respect to the GH 292 enzymes annotated in the genome. *P. atlantica* genome possesses one sequence in GH5, one in 293 GH8, one in GH10, and three sequences in GH43 (Table 1) potentially active on β-linked-1,4-294 mannan, glucan and xylan. For this reason, *P. atlantica* culture broth was complemented with 295 either konjac GlcM or beechwood GuX or a mixture of CMC, tara GalM and beechwood GuX 296 (abbreviated as '3P', Table 3). Beechwood GuX added alone or in admixture with CMC and 297 tara GalM (3P) in the culture broth reinforced the activities against itself, and against rye and 298 wheat AX, while the activities towards apple and tomato XyG disappeared. 299 The improvement of the secretion of hemicellulolytic enzymes by adding polysaccharide to P.

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300

302 *3-2 Effect of polysaccharides on growth kinetics and enzyme production*

atlantica culture broth was further investigated.

303 Growth of *P. atlantica* without any added polysaccharide was followed over 48 h (Figure 1). 304 Growth was slow for the first 5 h and increased rapidly to reach a plateau after 20 h culture. 305 Adding konjac GlcM did not lead to a significant change in the bacterial growth in comparison 306 to culture in the absence of added polysaccharide. Conversely beechwood GuX positively 307 affected the growth rate of *P. atlantica* as it induced faster growth from 10 to 20 h. 308 The culture filtrates were recovered from each culture and tested for their enzymatic activity 309 on 12 different hemicelluloses and 8 nitrophenyl-derivatives at pH 5.2 and 8.5 to identify the 310 secreted activities (Figure 2). The culture filtrates obtained without added polysaccharide and 311 in the presence of konjac GlcM exhibited roughly similar activity profiles on hemicelluloses 312 (Figures 2A and 2C, respectively). Activities on xylan-like polysaccharides were favoured at pH 8.5. Also the activities towards nitrophenyl-derivatives were similar in the two culture 313 314 filtrates (Figures 2B and 2D). The activities towards *o*-NP β-D-xylopyranoside, *p*-NP α-Larabinofuranoside and *p*-NP acetate were higher at pH 5.2 than at pH 8.5. Comparison of 315 316 Figures 2A and 2B and Figures 2C and 2D demonstrated that adding konjac GlcM in the

culture medium did not improve the production of hemicellulases or glycosidases. This result
agreed with the growth kinetics study, which showed that konjac GlcM did not significantly
enhance the bacterial growth.

In the culture filtrate produced in the presence of beechwood GuX (Figures 2E and 2F), the
activities on heteroxylans were reinforced, mostly at pH 8.5. Conversely to what was observed
in the other culture conditions, the activity on *o*-NP xylopyranoside was much higher at pH
8.5.

324

325 *3-3 Fractionation of culture filtrate obtained on beechwood glucuronoxylan*

326 The culture filtrate obtained in the presence of beechwood GuX was fractionated by ion 327 exchange chromatography. The activities were evaluated in the collected fractions on konjac 328 GlcM, CMC, beechwood GuX and wheat AX, and also on *o*-NP β-D-xylopyranoside and *p*-NP α-329 L-arabinofuranoside (Figure 3). The results suggested that enzymes active on konjac GlcM 330 and CMC (Figure 3A) eluted mainly with xylosidase activity (Figure 3B), whereas enzymes 331 active on beechwood GuX and wheat AX eluted later in the gradient (Figure 3A). 332 The gel electrophoresis of the fractions eluted between 7 and 13 mL elution volume of the ion 333 exchange chromatography (Fractions B2 to B12, Figure 4A) showed that the major band in 334 fractions B4 to B6 stand around 45 kDa and could correspond to the GH5 β-glucanase 335 Patl_1404. These fractions also contained a slight band around 55 kDa and a band at 72 kDa 336 possibly corresponding to the GH43 arabinofuranosidase Patl_0842 and the GH3-like protein 337 Patl_4162, respectively. In the fractions B7 to B9, the major band was around 42 kDa and 338 could correspond to the GH10 β-xylanase Patl_2657. Thus the electrophoresis demonstrated 339 the proteins associated with the enzyme activities recovered from the chromatography and is 340 an additional element to suggest the β -glucanase activity of GH5 Patl_1404, the 341 arabinofuranosidases activity of GH43 Patl_0842 and the β -xylanase activity of GH10 342 Patl_2657.

In the fractions B9 to B12 a lighter band around 96 kDa possibly corresponded to the GH8
protein Patl_1069. The band was excised and submitted to trypsic digestion. The LC-MS/MS
analysis of the digests confirmed that it corresponded to Patl_1069 (Supplementary data 1).
As the activity of this GH8 protein was not yet described, it was cloned and expressed in order
to analyse further its activity and specificity.

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349 *3-4 Production and activity of the recombinant enzyme rPaGH8.*

The GH8 protein Patl_1069 was expressed in the heterologous system *E. coli* with a poly-His
tag fused to its N-terminus. The recombinant protein rPaGH8 was purified on a His trap
affinity column. Three fractions eluting in the imidazole gradient and containing a band at 96
kDa (Fractions 4 to 6 on Figure 4B) were pooled and further tested for their enzymatic
activity. The pooled fraction exhibited 289 nkat/mL on soluble mix-linked xylan from *Palmaria palmata* confirming that rPaGH8 is a xylanase. Conversely, rPaGH8 did not degrade
o-NP xyloside, and thus has no β-xylosidase activity.

357

358 3-5 Specificity and mode of action of rPaGH8

359 The hydrolysis products released by rPaGH8 from heteroxylans from various sources were 360 analysed by HPAEC and compared to the hydrolysis products released by the commercial 361 GH11 xylanase M1 (Figure 5A). The first substrate tested was the soluble mix-linked xylan from *Palmaria palmata*, which contains mainly (1,4) linkages but also about 20% (1,3) 362 363 linkages [25]. The GH11 xylanase M1 produced monomer (X1), $\beta(1,4)$ dimer (X2), $\beta(1,4)$ trimer (X3), traces of tetramer (X4), and two peaks eluting at 13.2 min and 14.1 min, 364 365 potentially containing (1,3) linkages. On the contrary, rPaGH8 did not release X1 but released 366 significant amount of X3 and $\beta(1,4)$ hexamer (X6) eluting at 12.4 min. It produced two additional unknown peaks eluted at 13.4 min and 14.5 min, the later being minor (Table 4). 367

368 When beechwood glucuronoxylan was hydrolysed by xylanase M1 or rPaGH8, the later 369 released three unidentified products absent with the former, at 17.4 min, 18.3 min and 19.0 370 min (Figure 5B and Table 4). When water-extractable or water-unextractable arabinoxylans 371 from wheat (WEAX and WUAX, respectively) were submitted to xylanase M1 (Figure 5C and 372 D), many different products appeared including X1 and X2 and many ramified 373 oligosaccharides, of which the majority has been previously identified [32]. rPaGH8 produced 374 X1 to X4 and many ramified oligosaccharides. Some of them were similar to those produced 375 by xylanase M1, but in different amounts. Remarkably rPaGH8 released lower amounts of 376 oligosaccharides with 1 or 2 un-substituted xylose at the reducing end (such as XA3X and 377 XA3XX) and higher amount of oligosaccharides with 3 un-substituted xylose at the reducing 378 end (such as XA2,3XXX). Three additional peaks appeared at 19.0 min, 19.7 min, 23.7 min. The 379 same unidentified peaks were produced from both WEAX and WUAX, and the peak at 19.0 380 min was also produced from beechwood GuX (Table 4).

From all these substrates, rPaGH8 produced linear xylooligosaccharides from degree of
polymerisation 1 (as traces) to 4, suggesting that the enzyme is unable to hydrolyse X4. X5
was absent (from WEAX, WUAX) or present in very minor amounts (from beechwood GuX),
suggesting that it was hydrolysed in X3 + X2. X6 accumulated from *Palmaria* xylan, suggesting
that rPaGH8 was not able to degrade it. As the polysaccharide contained (1,3) as well as (1,4)
linkages, it is likely that this accumulating oligosaccharide contained one (1,3) linkage that
prevented its degradation by rPaGH8.

388 The unknown peak eluting at 19.0 min (marked with a star on Figure 5) was chosen to

tentatively elucidate its structure because it was released from wheat arabinoxylans and

beechwood glucuronoxylan. To this aim, the peak was isolated, desalted and analysed by mass

391 spectrometry. The MS analysis showed a m/z of 1073.35, corresponding to an

392 oligosaccharide with a degree of polymerisation of 8 (Figure 6A). The MS/MS analysis did not

allow a complete characterization of this structure. Since all hydroxyl functions could not be

394 isolated via intracyclic fragments, several structures were possible. However, the elution time 395 of the linear X8 may be estimated around 14 min taking into account the elution time of the 396 available linear oligosaccharides (X1 to X6). Thus, it is unlikely that the unknown peak 397 corresponds to the linear oligosaccharide X8. Consequently, it could correspond to a branched 398 oligosaccharide with 1 arabinose and 7 xylose residues (A1X7) or 2 arabinose and 6 xylose 399 residues (A2X6) (Figure 6C). In this last case, the two arabinose residues could be located 400 either on the 03 of the two non-reducing xylose residues or on 02 and 03 of the last non-401 reducing xylose residue (see the orange-marked hydroxyl positions on Figure 6C). The 402 structure presented on Figure 6C was the most probable as the non-reducing xylose residue 403 was free of any substituent. Indeed during catalysis, this xylose was bound to the sub-site +1 404 of the enzyme, which is not supposed to accept a substituted xylose. From the MS/MS 405 spectrum (Figure 6B) it is sure that this oligosaccharide contained at least 4 un-substituted 406 xylose residues at its reducing end.

407

408 **4- Discussion**

409 Hemicellulose-degrading enzymes contribute to the saccharification process of biomass for 410 the production of platform molecules or second-generation bioethanol [34 -39]. They also 411 have a great impact in animal feeding [40]. At an analytical level, polysaccharide-active 412 enzymes are important tools to solve the fine structure of their substrates. They are produced 413 by many microorganisms and bioinformatic tools are essential to select those potentially 414 producing enzymes of interest for their numerous applications. 415 *P. atlantica* is a marine bacterium known to produce enzymes active on seaweeds 416 polysaccharides such as agarases [41], alginate lyase [42] and porphyranase [43]. The 417 genome annotation suggested 14 genes coding for putative hemicellulases (Table 1) [44]. 418 Patl_1404 gene is the only gene from *P. atlantica* T6c coding a GH5 protein. Patl_2657 gene

419 was annotated as belonging to GH10, which suggested it as a xylanase. These predicted endo-

420 glucanase and endo-xylanase could explain the presence of activities on glucan, GlcM and 421 xylan in *P. atlantica* culture filtrate. Three sequences belong to GH43 and 4 to GH16. The 422 substrate specificity of these proteins, together with that belonging to GH8 (Patl 1069), is less 423 straightforwardly predictable, as these families are multispecific (5, 14, and 9 EC numbers in 424 families GH8, GH16 and GH43, respectively). Moreover, the choice of substrates is crucial to 425 succeed in evidencing enzymatic activities. As an example, the activities of *P. atlantica* culture 426 filtrates were much lower on maize AX in comparison to wheat AX and beechwood xylan. In 427 maize AX the xylan backbone can be decorated not only with arabinose but also with xylose 428 and galactose, and the arabinose moiety can be further decorated with xylose, galactose, 429 ferulic acid or one or more arabinose residues [45-46]. This gives it a more complex 430 structure that may make it more resistant to enzymatic hydrolysis than other AX. 431 Consequently, the hydrolysis of maize AX required a wider set of enzymes acting 432 synergistically, some of which may be lacking in *P. atlantica* secretome. 433 A β-xylosidase activity was shown in *P. atlantica* culture filtrates. The genome contains two GH3-like sequences (Patl 3730, Patl 4162) whose family includes β -xylosidases, α -434 435 arabinofuranosidases and β -glucosidases. It also contains 3 putative α -arabinofuranosidases 436 belonging to GH43 family (Patl_0810, Patl_0842 and Patl_3728) of which it is known that 437 many characterized enzymes showed a dual-activity α -arabinofuranosidase/ β -xylosidase 438 [47-50]. Thereby, there is two possible explanations for this β -xylosidase activity, either a 439 GH3 β -xylosidase or a GH43 dual-enzyme. 440 The xylanase activity shown after fractionation of *P. atlantica* secretome could be ensured by 441 two enzymes, the GH10 Patl_2657 and the GH8 Patl_1069. GH10 xylanases are characterized 442 by their high versatility since they hydrolyse many types of heteroxylans in which they can act near a substituted xylose residue [51 – 53]. GH8 family contains xylanases for which little 443

data is available to date, and reducing-end-xylose releasing exo-oligoxylanase (EC 3.2.1.156),

445 which hydrolyses β -(1 \rightarrow 4)-D-xylose residues from the reducing end of xylooligosaccharides

446 (www.cazy.org/GH8) and can be active on branched oligosaccharides [54]. To date, no 447 xylanase was described in Pseudoalteromonas atlantica. As we demonstrated that rPaGH8 448 produced oligosaccharides from wheat arabinoxylans, this ensures that it is an endoxylanase 449 (EC 3.2.1.8) and not a reducing-end-xylose releasing exo-oligoxylanase. In addition, the mode 450 of action of this xylanase on arabinoxylan was unusual since it favoured oligosaccharides with 451 side arabinose residues grouped on the non-reducing end and with an un-substituted tail on 452 the reducing end. Interestingly, two other GH8 enzymes were demonstrated in other 453 *Pseudoalteromonas* strains, the psychrotolerant *P. arctica* [55] and the psychrophilic *P.* 454 haloplanktis [56]. GH8 xylanase from P. haloplanktis has an inverting mechanism and a 455 (α/α) 6-fold. The increase in arabinose substitution on the substrate hindered *P. haloplanktis* 456 xylanase suggesting that the tolerance of GH8 xylanase towards the arabinose may be lower 457 than that of GH10 ones [57] and higher than those of GH11 which can only hydrolyse 458 xylosidic bonds between two un-branched xylose residues [51]. The crystal structure analysis 459 showed a long substrate-binding cleft from +4 to -3 subsites [58], which can be consistent 460 with the structure shown in the present study for the hydrolysis products. Moreover, P. 461 *haloplanktis* xylanase highlights a secondary binding site at the surface of the protein, which 462 favoured significantly the activity towards insoluble substrates [59]. However, its molar mass 463 was 48.4 kDa, two times less than PaGH8 xylanase, suggesting that the latter may be multi-464 modular. Using the genome browser on biocyc.org allowed showing that the corresponding 465 gene may belong to a short operon with a gene whose product was annotated as a 466 hypothetical protein (Supplementary data 2). This gene did not belong therefore to an operon 467 with another GH, or to a polysaccharide-utilizing locus that could have helped the concerted 468 work of polysaccharides-active enzymes.

PaGH8 was active on various xylan-like substrates, originating from the seaweed *Palmaria palmata* or from terrestrial plants, although *P. atlantica* originates from marine environment.
Although it remains unclear why marine bacteria secrete enzymes active towards plant

472 polysaccharides, this observation is consistent with the previous evidence that marine 473 bacteria secrete pectin-degrading enzymes in the presence of pectin. This was demonstrated for bacteria close to *P. atlantica* such as *P. haloplanktis* [60] and *Alteromonas macleodii* [61]. 474 475 The polysaccharides tested as substrates in the present study have rather known structures. 476 The structure of wheat AX was extensively studied and shown to vary largely in its level of 477 arabinose di-substitution [62]. NMR spectroscopy revealed that beechwood GuX contained 478 approximately one 4-0-methylglucuronic acid every 15 xylose residues [63]. From these 479 polysaccharides PaGH8 produced oligosaccharides in which the arabinose substitutions were 480 gathered at the non-reducing side of the molecule, showing that the initial polysaccharides contained such stretches of xylose residues without any substitution. These new 481 oligosaccharides pave the way to elucidate further the chemical structure of the 482 polysaccharides thanks to PaGH8 particular behaviour. 483 484 Altogether the present results highlighted *P. atlantica* as a source of hemicellulose-degrading 485 enzymes. Our data showed that the use of various polysaccharides generated a wide variety of 486 enzymes. Only those produced in the presence of glucuronoxylan were studied, which particularly 487 revealed the new GH8 xylanase. Results obtained with one or more other polysaccharide substrates 488 lead to the production of a wide range of enzymes that remain to be valued in the near future. 489

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497 Authors' contributions

- 498 EB, ML, WH designed the research. SR, JV, AB, AG, MF performed the research experiments.
- 499 SR, MFA, MF, DR, WH, ML and EB analyzed the data. SR and EB drafted the manuscript. All
- 500 authors read and approved the final manuscript.
- 501

502 **Conflict of interest**

- 503 None declared
- 504

505 **References**

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Figure 1: *P. atlantica* growth in the absence of added polysaccharide (black line), and in the
presence of konjac glucomannan, GlcM (green line) and beechwood glucuronoxylan, GuX (red
line).

683

Figure 2: Enzymatic activities measured in the culture supernatants of *P. atlantica* on

hemicelluloses (A, C, E, respectively) and *p*-NP-sugar derivatives (B, D, F, respectively). Three

686 culture conditions were applied, without any added polysaccharide (A, B), in the presence of

687 konjac glucomanann, GlcM (C, D) and in the presence of beechwood glucuronoxylan, GuX (E,

688 **F**). Each activity was measured at pH 5.2 (black bars) and 8.5 (grey bars).

689

Figure 3: Enzyme profile after fractionation by anion exchange chromatography of the

691 culture filtrate of *P. atlantica* grown in the presence of beechwood glucuronoxylan (GuX).

692 (A) Activities measured on carboxymethyl cellulose, CMC (black full line), konjac

693 glucomannan, GlcM (black dotted line), beechwood glucuronoxylan GuX (grey full line) and

694 wheat arabinoxylan AX (grey dotted line).

695 (B) Activities measured on *p*-NP α -L-arabinofuranoside, *p*-NP α -L-Ara (full line), and *o*-NP β-

696 D-xylopyranoside, o-NP β -D-Xyl (dotted line).

697

698 **Figure 4:** SDS-PAGE

699 (A) The labels B2 to B12 referred to the fractions recovered between 7 and 13 mL elution

volume of the anion exchange chromatography of the *P. atlantica* culture filtrate obtained on

701 beechwood glucuronoxylan (see Figure 3).

702	(B) The labels 1	to 8 referred to t	he fractions	eluted from	the His trap	affinity of	column for th	he
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- purification of the recombinant rPaGH8. The fractions 4 to 6 were pooled to get the
- recombinant rPaGH8.
- 705 M: molecular weight markers.
- 706
- **Figure 5:** HPAEC analysis of the reaction products released by rPaGH8 (black line) and the
- commercial xylanase M1 (green line) from different xylan substrates. The labels refer to the
- structure of the oligosaccharides, using the nomenclature of Fauré et al., 2009 [27].
- 710 (A): Xylan from *Palmaria palmata*
- 711 **(B)**: Beechwood glucuronoxylan
- 712 (C): Water extractable arabinoxylan from wheat endosperm
- 713 (D): Water unextractable arabinoxylan from wheat endosperm
- *: oligosaccharide whose structure has been subsequently identified by mass spectrometry
- 715
- **Figure 6**: Mass spectrometry analysis of the HPAEC fraction collected at the retention time of
- 717 19.0 min (see Figure 5).
- (A) Zoom of the MS spectrum with the peak corresponding to an AX DP8 (m/z 1073.35).
- 719 **(B)** MS/MS spectrum of the 1073.35 *m/z* species after further fragmentation.
- 720 (C) One possible structure of observed fragment depicted according to the nomenclature of
- 721 Domon and Costello, 1998 [33]. The two pentoses in orange can be placed in any orange-
- marked position. Blue annotations: intercyclic fragments; Red annotations: intracyclic
- 723 fragments; Stars: fragments with the loss of one water molecule.
- 724

- 726 Supplementary data
- 727 **Supplementary data 1:** Amino-acid sequence of the GH8 protein Patl_1069 excised from the
- 728 PAGE gel (see Figure 4). The matched peptides were identified by LC-MS/MS and are
- highlighted in red. They lead to sequence coverage of 8%.
- 730
- 731 **Supplementary data 2:** *P. atlantica* genome visualized using the genome browser on
- biocyc.org (position 1,214,005 to 1,348,215 bp). The gene PATL_RS05460 (in brown and
- framed in bold in the centre of the figure) corresponds to the protein UniProt Q15WZ3
- 734 (Patl_1069) and its locus is located between position 1,279,741 and 1,282,479 bp. The colours
- refer to putative operons. The gene PATL_RS05460 forms a putative operon with the gene
- PATL_RS05455 located just upstream on the genome. The product of this gene is annotated as
- a hypothetical protein.
- 738

- **Table 1**: Presentation of *Pseudoalteromonas atlantica* T6c strain [15]: origin, culture
- 740 conditions and occurrence of the hemicellulose-modifying GH families.

General information	Strain	Origin				
	ATCC BAA-1087	Marine				
Culture conditions						
Medium	Bactomarine broth (DIFCO 2216)					
рН	7.4					
T (°C)	20					
Occurrence of hemicellulose-related GH families						
GH family	Number of sequences	Gene entry				
		Patl_0825				
GH3	4	Patl_1716				
0115	4	Patl_4162				
		Patl_3730				
GH5	1	Patl_1404				
GH8	1	Patl_1069				
GH10	1	Patl_2657				
		Patl_0805				
GH16	4	Patl_0824				
GHIU	7	Patl_0843				
		Patl_0880				
		Patl_0810				
GH43	3	Patl_0842				
		Patl_3728				

- **Table 2**: List of polysaccharides used as substrates for the screening of polysaccharides-
- 744 degrading enzymes.

	Polysaccharide	Biological origin	Source
1	Arabinoxylan (AX)	Maize	Lab collection
2	Arabinoxylan (AX)	Rye	Megazyme
3	Arabinoxylan (AX)	Wheat	Megazyme
4	Glucuronoxylan (GuX)	Beechwood	Sigma
5	Glucuronoxylan (GuX)	Tomato	[16]
6	Glucuronoarabinoxylan (GAX)	Apple	[8]
7	Glucomannan (GlcM)	Amorphophallus konjac (Konjac powder)	Dextra Lab
8	Galactomannan (GalM)	Caesalpina spinosa (Tara gum)	Dextra Lab
9	Galactomannan (GalM)	Cyamopsis tetragonolobus (Guar gum)	Dextra Lab
10	Galactomannan (GalM)	Ceretonia siliqua (Locust bean gum)	Dextra Lab
11	β (1,4) Mannan	Carob	Megazyme
12	Galactoglucomannan (GgM, E3)	Apple	[8]
13	Galactoglucomannan (GgM, A1)	Apple	[8]
14	Galactoglucomannan (GgM)	Tomato	Lab collection
15	CarboxyMethyl Cellulose (CMC)		Sigma
16	Avicel		Sigma
17	Xyloglucan (XyG)	Tamarindus indica (Tamarind gum)	Dextra Lab
18	Acetylated xyloglycan (AcXyG)	Apple	[8]
19	Xyloglucan (XyG)	Apple	[8]
20	Xyloglucan (XyG)	Tomato	[16]
21	Partially depectinated AIR* (pDAIR)	Apple	[8]
22	Arabinan	Sugar beet	Megazyme
23	Welan gum	Alcaligenes sp.	Dextra Lab
24	Tragacanth gum	Astragalus	Dextra Lab
25	Arabic gum	Acacia sp.	Dextra Lab
26	Amylopectin	Waxy corn	TCI
27	Amylose	Potato	Lab collection
28	Pullulan	Scerotium rolfsii	Dextra Lab
29	High methylated Pectin (HMPectin)	Apple	[8]
30	Low methylated Pectin (LMPectin)	Citrus	Cargill
31	Rhamnogalacturonan (RG)	Carrot	Lab collection
32	Rhamnogalacturonan (RG)	Apple	[8]
33	Xanthan	Xanthomonas campestris	Dextra Lab
34	Dextran	Leuconostoc mesenteroides	Dextra Lab
35	α, α Trehalose		Carbosynth Ltd
36	ι–ν Carragenan	Kappaphycus alvarezzi	CP-Kelco
37	κ–μ Carragenan	Eucheuma denticulatum	CP-Kelco
38	κ-Carragenan	Kappaphycus alvarezzi	CP-Kelco
39	Porphyran	Porphyra umbilicalis	Lab collection
40	Mannuronan		Lab collection
41	Ulvan	Ulva rotundata	Lab collection
42	Alginate	Ascophyllum sp.	Lab collection

* AIR = Alcohol Insoluble Residue

Table 3: Colorimetric profiling of the hemicellulose-degrading enzymes on 42 747 748 polysaccharides in the extracellular (E) and intracellular (I) culture filtrates of 749 *Pseudoalteromonas atlantica* T6c grown without or with added polysaccharides: mixture of 750 carboxymethyl cellulose + tara galactomannan + beechwood xylan (abbreviated as '3P'), 751 konjac glucomannan (GlcM) and beechwood xylan. Positive hits of polysaccharide 752 degradation were classed according to 4 intervals based on the control mean (x) and standard 753 deviation (σ) of negative hit polysaccharides. The extent of degradation was indicated according to grey intensity, $x - 2\sigma < absorbance < x - 4\sigma$; 754

755

 $x - 4\sigma$ absorbance $< x - 6\sigma$; $x - 6\sigma < absorbance <math>< x - 8\sigma$; $x - 8\sigma < absorbance$.

Added Polysaccharides	No		3P	GlcM	Xylan
Culture filtrate*	E	1	Е	E	E
Protein concentration (µg/mL)	1265	1470	821	606	660
Screening substrate					
Maize AX	1	1			
Rye AX					
Wheat AX					
Beechwood xylan					
Tomato GuX					
Apple GAX	1				
Konjac GlcM					1
Tara GalM					
Guar GalM					
Locust GalM					
β (1,4) Mannan					
Apple GgM		1			
Apple GgM					
Tomato GgM					
CMC		1			Í
Avicel					
Tamarind XyG					
Apple AcXyG					
Apple XyG					
Tomato XyG					
pDAIR					
Arabinan					
Welan gum					
Tragacanth gum					
Arabic gum					
Amylopectin					
Amylose					
Pullulan					
Apple HM Pectin	1				
Citrus LM Pectin	1				

	l	l	1
Carrot RG			
Apple RG		1	
Xanthan			
Dextran			
α,α Trehalose			
ι–ν Carragenan			
κ–μ Carragenan			
к-Carragenan			
Porphyran			
Mannuronan			
Ulvan			
Alginate			
* Two type of culture filtrates: E	= extrace	ellular, I :	= intracellular

- **Table 4**: Hydrolysis products released by rPaGH8 from various hemicelluloses. Crosses (+) in
- bold indicate oligosaccharides produced exclusively by rPaGH8 and not by xylanase M1. The
- number of + is used as an indication of the peak size.

		Xylan	Beechwood		
		Palmaria	GuX	WEAX	WUAX
Ara	3.3				+
X1	3.7		+	+	+
X2	5.4	+	+++	+	+
ХЗ	8.1	++	+++	+	+
X4	10	+	+++	+	+
X5	11.2		+		
X6	12.4	++			
X7 ?	13.2	+			
Unknown	13.4	+			
XA3X	13.8			++	++
Unknown	14.2				
XA3XX	14.5			++	++
Unknown#	14.5	+			
XXA3XX	15.3		+	+	+
Unknown	16.3		++		
Unknown	17.4		++		
XA2,3XXX	18			+	+
Unknown	18.3		++		
Unknown*	19.0		++	+	+
Unknown	19.7		+	+	+
Unknown	20.1				
Unknown	21.4			+	+
XA3A3XX	21.7			+	+
XA3XA3XX	22.2			+	+
Unknown	23.7			+	+
XA3A2,3XX	24.2			+	+

761 #this peak eluted at the same time as XA3XX but corresponds to another oligosaccharide

as the xylan from *Palmaria* does not contain any arabinose.

763 * this peak was further identified by mass spectrometry

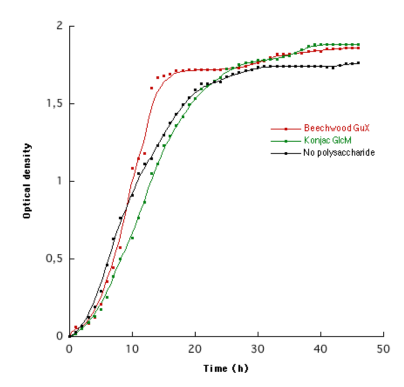


Figure 1: *P. atlantica* growth in the absence of added polysaccharide (black line), and in the presence of konjac glucomannan, GlcM (green line) and beechwood glucuronoxylan, GuX (red line).

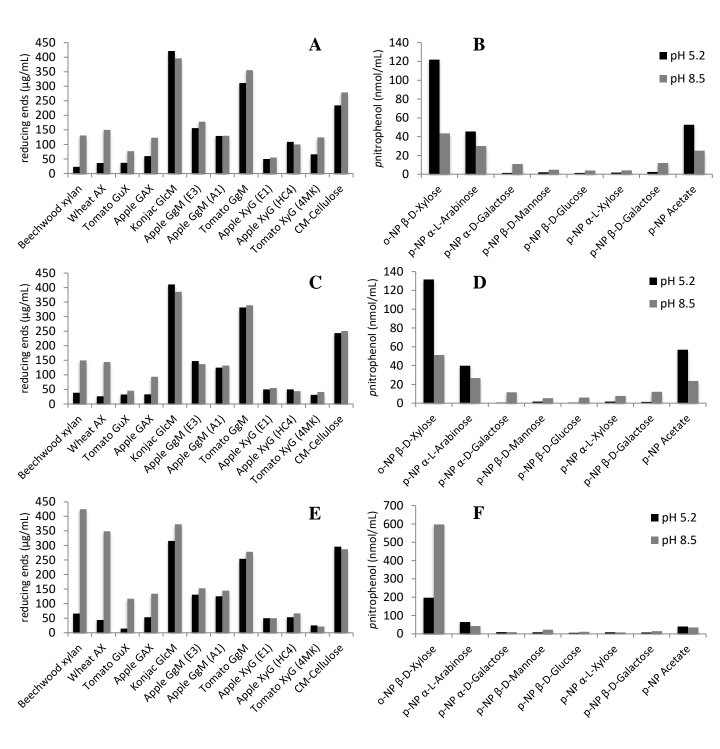


Figure 2: Enzymatic activities measured in the culture supernatants of *P. atlantica* on hemicelluloses (**A**, **C**, **E**, respectively) and *p*-NP-sugar derivatives (**B**, **D**, **F**, respectively). Three culture conditions were applied, without any added polysaccharide (**A**, **B**), in the presence of konjac glucomanann, GlcM (**C**, **D**) and in the presence of beechwood glucuronoxylan, GuX (**E**, **F**). Each activity was measured at pH 5.2 (black bars) and 8.5 (grey bars).

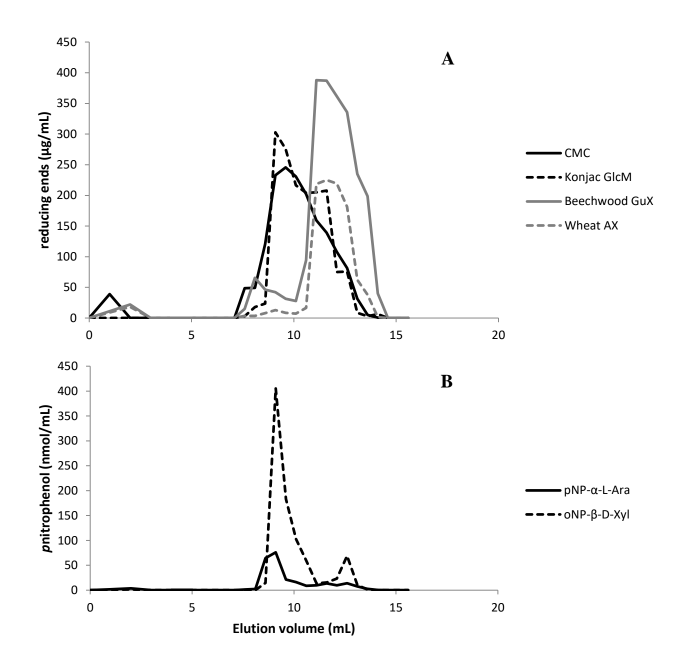


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(A) Activities measured on carboxymethyl cellulose, CMC (black full line), konjac glucomannan, GlcM (black dotted line), beechwood glucuronoxylan GuX (grey full line) and wheat arabinoxylan AX (grey dotted line).

(**B**) Activities measured on *p*-NP α -L-arabinofuranoside, *p*-NP α -L-Ara (full line), and *o*-NP β -D-xylopyranoside, *o*-NP β -D-Xyl (dotted line).

А

B2 Μ B3 B4 B5 B6 B7 B8 B9 B10 B11 B12 Μ 2 3 5 6 7 8 1 4 170 kDa 200 kDa 130 kDa 150 kDa 95 kDa ←Patl 1069 120 kDa 72 kDa -Patl_4162 100 kDa 85 kDa -Patl 0842 55 kDa 70 kDa ←Patl 1404 60 kDa 43 kDa ← Patl_2657 collected 50 kDa 34 kDa fractions 40 kDa 26 kDa 30 kDa 25 kDa 17 kDa 10 kDa 20 kDa

Figure 4: SDS-PAGE

(A) The labels B2 to B12 referred to the fractions recovered between 7 and 13 mL elution volume of the anion exchange chromatography of the *P. atlantica* culture filtrate obtained on beechwood glucuronoxylan (see Figure 3).
(B) The labels 1 to 8 referred to the fractions eluted from the His trap affinity column for the purification of the recombinant rPaGH8. The fractions 4 to 6 were pooled to get the recombinant rPaGH8.
M: molecular weight markers.

В

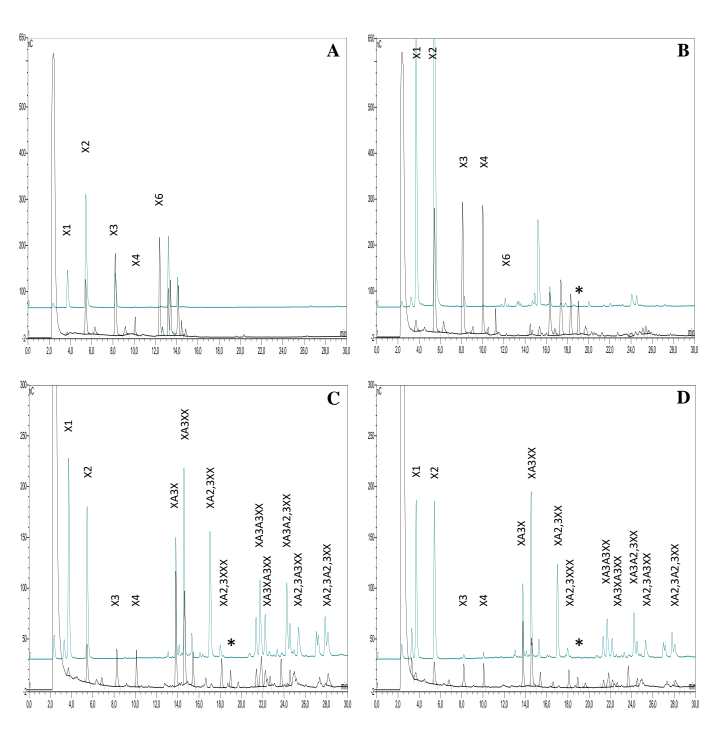


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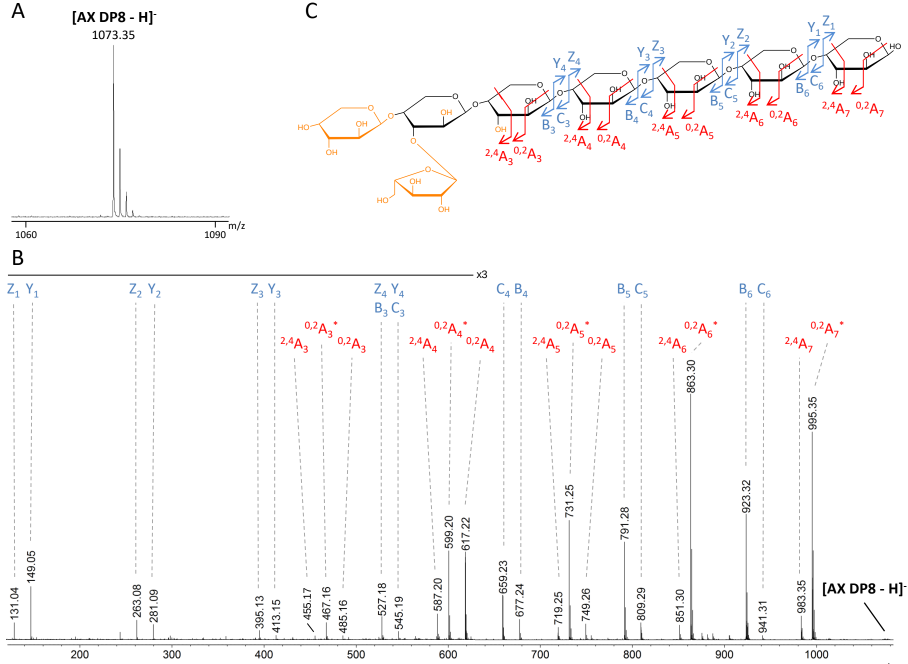


Figure 6: Mass spectrometry analysis of the HPAEC fraction collected at the retention time of 19.0 min.