

Highlights:

- Neutral and/or acidic XOS were obtained depending on the GH family of xylanase used
- XOS produced from glucuronoxylans have high antioxidant activity
- MeGlcA ramifications of XOS enhance their antioxidant effect
- Antioxidant power of XOS depends on the degree of polymerisation and substrate used
- High antioxidant activity was found on the eucalyptus autohydrolysate
- ABTS method is more sensitive method for the XOS antioxidant activity than DPPH

1 **Antioxidant activity of xylooligosaccharides produced from**
2 **glucuronoxyylan by Xyn10A and Xyn30D xylanases and eucalyptus**
3 **autohydrolysates**

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25 **ABSTRACT**

26 Antioxidant activity of xylooligosaccharides (XOS) released from beechwood
27 and birchwood glucuronoxylans by two different xylanases, one from family
28 GH10 (Xyn10A) and another from family GH30 (Xyn30D) was examined. The
29 ABTS (2, 2'-Azino-bis(3-ethylbenzothiazoline-6-sulfonic acid)) method was
30 used, since it resulted more accurate for the antioxidant activity determination of
31 XOS. Thin layer chromatography and MALDI-TOF MS analysis showed that
32 Xyn10A produced a mixture of neutral and acidic XOS whereas the XOS
33 produced by Xyn30D were all acidic, containing a methylglucuronic acid
34 (MeGlcA) ramification. These acidic XOS, MeGlcA substituted, showed a
35 strongly higher antioxidant activity than the XOS produced by Xyn10A (80% vs.
36 10% respectively, at 200 $\mu\text{g mL}^{-1}$). Moreover, the antioxidant activity increased
37 with the degree of polymerization of XOS, and depended on the xylan substrate
38 used. The antioxidant capacity of eucalyptus autohydrolysates after xylanase
39 treatment was also analysed, showing a decrease of their antioxidant activity
40 simultaneous with the decrease in XOS length.

41

42 **Keywords:**

43 Xyn10A, Xyn30D, antioxidant activity, xylooligosaccharides, glucuronoxylan,
44 eucalyptus autohydrolysate

45

46 1. INTRODUCTION

47 New interest has been aroused to search natural and safe antioxidant
48 agents from natural sources. An antioxidant compound can be defined as a
49 substance whose action can inhibit oxidation rate of a free radical. Although
50 synthetic antioxidants seem to be promising, their toxicity and side effects rule
51 out their extensive use. Plant biomass, which is the main source of renewable
52 materials on earth, consists largely of two polysaccharides (cellulose and
53 hemicelluloses) and an aromatic polymer (lignin). Hemicelluloses account for
54 15-25% of all lignocellulose. In hardwood species, grasses and agro-industrial
55 by-products (cereal straws, sugarcane bagasse, corn stover and sisal, among
56 others), xylans are the most abundant hemicelluloses (Aracri & Vidal, 2011;
57 Valls, Gallardo, et al., 2010; Valls, Cadena, & Roncero, 2013) whereas in
58 softwoods, mannans are more abundant (Scheller & Ulvskov, 2010). The term
59 arabinoxylan is used to describe xylan of cereals and grasses which shows
60 abundant decorations of arabinose and ferulic acid (Ebringerová & Heinze,
61 2000). On the other hand, xylan from hardwood species is denominated
62 glucuronoxylan as is highly substituted with methylglucuronic acid decorations
63 (MeGlcA), and can be heavily acetylated while it does not contain arabinose
64 and ferulic acid decorations (Teleman, Tenkanen, Jacobs, & Dahlman, 2002).

65 Endoxylanases (EC 3.2.1.8) randomly hydrolyse the β -(1,4) glycosidic
66 bonds of the xylose backbone of xylan (Biely, Vrsanská, Tenkanen, & Kluepfel,
67 1997; Pollet, Delcour, & Courtin, 2010). These enzymes are glycosyl hydrolases
68 (GH) grouped in families GH10, GH11 and GH30 (previously classified in family
69 GH5) (Lombard, Golaconda Ramulu, Drula, Coutinho, & Henrissat, 2014). The

70 hydrolysis products of these enzymes are a mixture of xylooligosaccharides
71 (XOS) of different degrees of polymerisation and substitution, depending on the
72 xylanase GH family. XOS show a structure of around 2–7 xylose units linked by
73 β -(1,4) bonds, variably decorated with different substituents depending on the
74 source. For example, XOS from hardwoods contain acetyl groups and MeGlcA
75 ramifications whereas XOS from agricultural residues contain arabinose units
76 and ferulic acid. While XOS released from glucuronoxylans by GH10 and GH11
77 xylanases are a mixture of linear (neutral) and MeGlcA substituted (acidic)
78 oligomers, XOS released by xylanases of family GH30 are all MeGlcA
79 substituted xylooligomers (Kolenova, Vrsanska, & Biely, 2006; Vrřanská,
80 Kolenová, Puchart, & Biely, 2007). Antioxidant activity of XOS from
81 arabinoxylans has been reported in several articles (Bian et al., 2013;
82 Gowdhaman & Ponnusami, 2015; Mandelli et al., 2014; Veenashri &
83 Muralikrishna, 2011), although in these works the activity of different types of
84 XOS has not been evaluated.

85 Xylanases from families GH10 (Xyn10A), and GH30 (Xyn30D) have been
86 previously isolated from *Paenibacillus barcinonensis* and characterized
87 (Valenzuela, Diaz, & Pastor, 2012; Valenzuela, Díaz, & Pastor, 2010). These
88 xylanases were previously tested for their bleach boosting ability on pulp from
89 eucalyptus and agricultural fibres, showing different effectiveness (Valenzuela
90 et al., 2013; Valls, Vidal, et al., 2010). We have applied Xyn10A and Xyn30D on
91 glucuronoxylans for XOS production in order to analyse their antioxidant effect.
92 A comparison between the antioxidant activity of XOS produced by xylanases of
93 different GH families has been evaluated in this work for the first time.

94 Additionally we have analysed the antioxidant activity of XOS released from
95 eucalyptus, the most important raw material for pulp production in Spain, a
96 process that generates a large amount of residues that can be upgraded to
97 added value products.

98 **2. MATERIALS AND METHODS**

99 **2.1. Raw material**

100 Birchwood, oat spelt xylans, D-Glucuronic acid, gallic acid and Trolox
101 were purchased from Sigma Aldrich (Germany). Beechwood xylan was
102 purchased from Roth (Germany). Autohydrolysates from eucalyptus wood were
103 a gift of J. Parajó (Gullón et al., 2011).

104 **2.2. Xylanases**

105 The xylanases assayed were Xyn10A from family GH10 (Valenzuela et
106 al., 2010) and Xyn30D from family GH30 (Valenzuela et al., 2012). Xyn10A is a
107 single-domain enzyme and thus comprised of a sole catalytic module whereas
108 Xyn30D is a modular enzyme comprised of a GH30 catalytic module linked to a
109 CBM35 carbohydrate binding module. These xylanases were recombinant
110 enzymes from *P. barcinonensis* that were previously cloned in *Escherichia coli*
111 and characterized. They were purified as described (Valenzuela et al., 2012;
112 Valenzuela et al., 2010).

113 Protein concentration was determined using the Bradford method
114 (Bradford, 1976). Xylanase activity was assayed by measuring the amount of
115 reducing sugars released from xylan hydrolysis by the dinitrosalicylic (DNS)
116 reagent method (Miller, 1959). DNS was purchased from Sigma Aldrich

117 (Germany). The 100 μ l volume standard assay contained the enzyme samples
118 in 50mM sodium-phosphate buffer pH 6.5 with 1.5% birchwood xylan, final
119 concentration. The reaction mixtures were incubated at 50° for 15 min. Then,
120 temperature was increased at 100°C for 2 min to inactivate the enzymes. To
121 quantify the reducing sugars released from substrate, samples were chilled, 100
122 μ L of DNS reagent were added and the reaction mixtures were incubated for 5
123 min at 100°C. Finally, 40 μ L of samples were mixed with 260 μ L of distilled
124 water in ELISA plates and the absorbance at 540 nm was measured. A
125 standard curve of xylose was used to calculate activity units. One unit of
126 xylanase activity was defined as the amount of enzyme that releases 1 μ mol of
127 xylose reducing sugar equivalent per min under the assay conditions described.
128 All determinations of enzyme activity were made in triplicate.

129 **2.3. Enzymatic treatments**

130 XOS were obtained incubating 2% xylans (wt/vol) (2.22 mL of a xylan
131 solution at 4.5%) with Xyn10A or Xyn30D at 10 U g⁻¹ of substrate in 50 mM
132 sodium-phosphate buffer at pH 6.5 for 2 h at 50°C in a final volume of 5 mL.
133 Incubations were performed in a thermostatic bath. These conditions
134 correspond to the optimum pH for the activity of both enzymes and optimum
135 temperature for Xyn30D activity. Xyn10A shows optimal activity at 60°C while at
136 50°C shows more than 85% of maximum activity (Valenzuela et al., 2012;
137 Valenzuela et al., 2010). After incubation, unhydrolysed and partially hydrolysed
138 xylans were separated from XOS by step precipitation with three volumes of
139 ethanol. For this purpose, 5 mL of ethanol were added to the 5 mL volume

140 reaction and samples were centrifuged at 4500 rpm (revolutions per min) for 10
141 min. The supernatants were collected, mixed with additional 10 mL of ethanol
142 and centrifuged again at 4500 rpm for 10 min. The resulting supernatants of this
143 second centrifugation were collected, ethanol was eliminated by evaporation,
144 and dried samples obtained were considered as purified XOS. They were
145 dissolved in 1.5 mL of 200 mM sodium-phosphate buffer pH 6.5, analysed for
146 their content in reducing sugars, and kept for the determination of antioxidant
147 properties.

148 **2.4. Determination of reducing sugar**

149 The reducing sugar content of the XOS released from xylan hydrolysis
150 was measured according to the dinitrosalicylic (DNS) reagent method (Miller,
151 1959). 100 μ l of DNS were added to 100 μ l of XOS samples and mixtures were
152 incubated at 100°C during 5 min. Then, 40 μ L of reaction samples were placed
153 in ELISA plates, 260 μ L of distilled water were added and the absorbance at
154 540 nm was measured. Samples were analysed by triplicate. A standard curve
155 of xylose was used to calculate the xylose reducing sugar equivalent of the
156 different samples.

157 **2.5. Antioxidant activity (%)**

158 The antioxidant activity was assessed by a procedure consisting in the
159 quantification of the ABTS^{•+} radical decoloration described by several authors
160 (Cusola, Valls, Vidal, & Roncero, 2015; Re et al., 1999; Valls & Roncero, 2013)
161 with some modifications. The method consists in the addition of the antioxidant
162 compound to a pre-formed ABTS^{•+} radical solution and quantifying the

163 remaining ABTS^{•+} after a fixed time period, by means of UV spectrophotometry.
164 Firstly, the ABTS^{•+} radical was pre-formed adding 44.5 μL of 140 mM potassium
165 persulfate to 2.5 mL of 7 mM ABTS and keeping at darkness for 16h. Due to
166 the precipitation of XOS of high molecular weight with ethanol, ABTS^{•+} was
167 dissolved in water instead of in ethanol until the absorbance was 0.7±0.1 at 730
168 nm. 900 μL of the ABTS^{•+} solution and 100 μL of XOS (at several
169 concentrations) were mixed in a methacrylate cuvette. The reaction was left in
170 the darkness during 10 min. Then, the final absorbance was measured at 730
171 nm. Complete decoloration meant 100% of antioxidant activity, while the
172 percentage of decoloration was equivalent to the % of antioxidant activity. It was
173 calculated as inhibition % as follows:

$$174 \text{ ABTS}^{\bullet+} \text{ inhib. (\%)} = 100 [(A_i - A_f)/A_i]$$

175 where A_i is the ABTS^{•+} absorbance value of the blank, and A_f is the ABTS^{•+}
176 absorbance value after contact with the antioxidant compound. The % of
177 inhibition or decoloration of ABTS by a compound was considered the % of its
178 antioxidant activity. As positive control, the potent antioxidant agents Gallic acid
179 and Trolox were used. Experiments were performed by triplicate on a T92+UV
180 Spectrophotometer (PG Instruments).

181 **2.6. Thin-layer chromatography**

182 Purified XOS were analyzed by thin-layer chromatography (TLC) as
183 previously described (Gallardo et al., 2010). 10-15 μL of XOS were applied on a
184 silica gel plate (Merck, Germany) constituting the solid phase. 10 μL of an
185 oligomer standard mixture containing neutral xylooligosaccharides (Megazyme,
186 Ireland) at a concentration of 20 mg mL⁻¹ were applied as migration standards.

187 The mobile phase was a mixture of chloroform, acetic acid and H₂O in a 6:7:1
188 ratio, respectively. The migration was repeated twice and the silica gel plate
189 was then sprayed (Fungilab S.A., Spain) with a developing solution, consisting
190 of 5% H₂SO₄ in ethanol. Finally, the plate was heated in the oven at 100°C for 5
191 min, where the spots corresponding to the different XOS were visualized.

192 **2.7. MALDI-TOF MS**

193 For the analysis by MALDI-TOF MS, 1 µl of XOS was mixed with 1 µl of
194 matrix solution (10 mg/ml 2, 5- dihydroxybenzoic acid dissolved in acetonitrile-
195 water [1:1, vol/vol], 0.1% [wt/vol] trifluoroacetic acid). One microliter of the
196 mixture was spotted onto the MALDI-TOF MS plate and allowed to dry before
197 the analysis. Positive mass spectra were collected with a 4800 Plus MALDI
198 TOF/TOF (ABSciex 2010) spectrometer with an Nd:YAG 200-Hz laser operated
199 at 355 nm.

200 **3. RESULTS AND DISCUSSION**

201 **3.1. Xylooligosaccharides (XOS) production**

202 Firstly, we analyzed the production of XOS from oat spelt arabinoxylan
203 by a xylanase of GH10 family, Xyn10A from *Paenibacillus barcinonensis*. Xylan
204 concentration was set at 2% according to Akpinar, Erdogan, Bakir, & Yilmaz,
205 2010 and Bian et al., 2013, which showed that higher substrate concentration
206 decreased the yield of XOS, due to the increase in viscosity and density of the
207 reaction mixture. We treated with several enzyme doses (0.1, 1, 10 and 40
208 Xylanase Units g⁻¹ of xylan) at pH 6.5 and 50°C for 2h and analyzed the
209 production of XOS. To quantify them, a mixture of oligomers of different

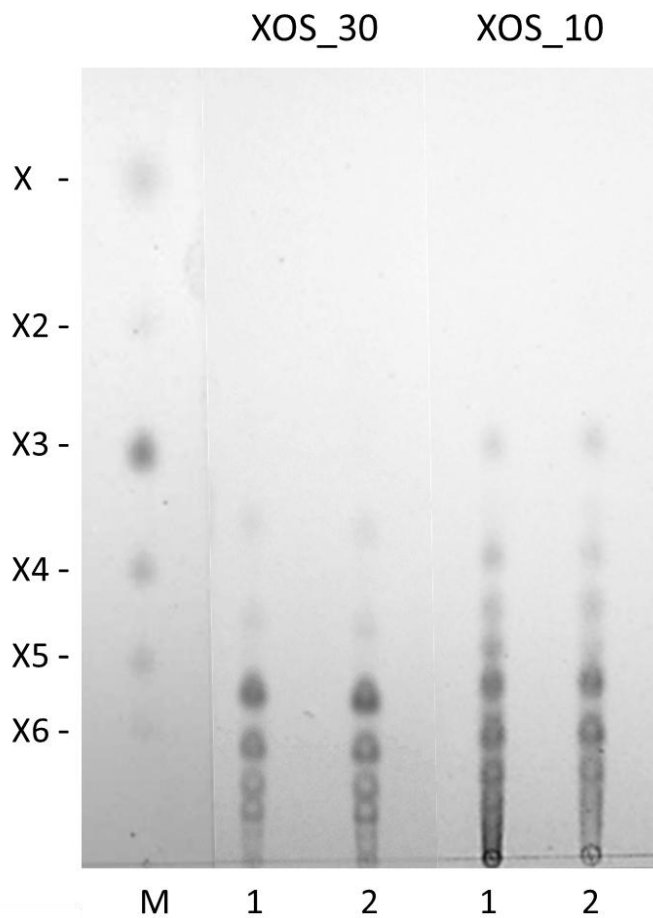
210 polymerization degree, we determined the release of reducing sugars. As
211 xylooligomers have a unique reducing group in their terminal xylose residue, we
212 assumed that one mol of xylose reducing sugar would be equivalent to one mol
213 of released XOS. We believe that this is a more accurate way to quantify XOS
214 that measuring in g the amount of products released from xylan, comprised by
215 XOS of quite different size. The doses of 0.1 and 1 U g⁻¹ of xylan produced low
216 amount of XOS, while with 10 U g⁻¹ the XOS production was increased until 2.1
217 g of xylose reducing equivalent L⁻¹. Higher enzyme concentrations (40 U g⁻¹)
218 gave rise to lower XOS production, probably because of poor xylan conversion
219 by high concentrations of enzyme and substrate. For this reason, the xylanase
220 dose of 10 U g⁻¹ was chosen for the following experiments.

221 A different type of xylan, beechwood glucuronoxytan, was tested for XOS
222 production. Beechwood xylan was treated with xylanase Xyn10A or with a
223 family GH30 xylanase, Xyn30D from *Paenibacillus barcinonensis*, in the same
224 conditions as above and the release of XOS at different intervals of incubation
225 was analyzed. Both xylanases released a similar amount of XOS (around 500
226 mg L⁻¹) at short incubation time (5 min), in accordance to their similar specific
227 activity. However, in prolonged incubations (120 min), while the amount of XOS
228 released by Xyn30D remained constant, higher XOS production (up to 1745 mg
229 L⁻¹) was obtained with Xyn10A, probably reflecting the higher thermostability of
230 this enzyme (Valenzuela et al., 2012; Valenzuela et al., 2010). To calculate the
231 yield of enzymatic treatments, the dry weight of the XOS produced by the
232 enzymes was determined. Treatment for 120 min with Xyn10A gave rise to 0.22

233 g of XOS from 1 g of birchwood xylan (22% yield) while Xyn30D gave rise to a
234 lower production (13%).

235 The XOS released from beechwood xylan were purified and analyzed by
236 thin layer chromatography (TLC). They showed a different pattern of oligomers
237 (Fig. 1). Hydrolysis products of Xyn10A (XOS_10) contain a mixture of
238 oligomers with the mobility of neutral XOS, such as xylotriose (X3), xylo-tetraose
239 (X4) xylopentaose (X5) and xylohexaose (X6), accompanied by oligomers of
240 intermediate mobility, indicating they are methylglucuronic acid (MeGlcA)
241 substituted oligomers (acidic XOS). Moreover, longer XOS were found, which
242 were less abundant in long term incubations (120 min). On the contrary, all
243 oligomers produced by Xyn30D (XOS_30) show mobility corresponding to
244 acidic XOS, in accordance to the mode of action of GH30 xylanases, that
245 exclusively release MeGlcA substituted xylooligomers from glucuronoxylans
246 (Kolenova et al., 2006; Vrřanská et al., 2007). Moreover, no differences in
247 oligomer size was found in long term incubations with Xyn30D.

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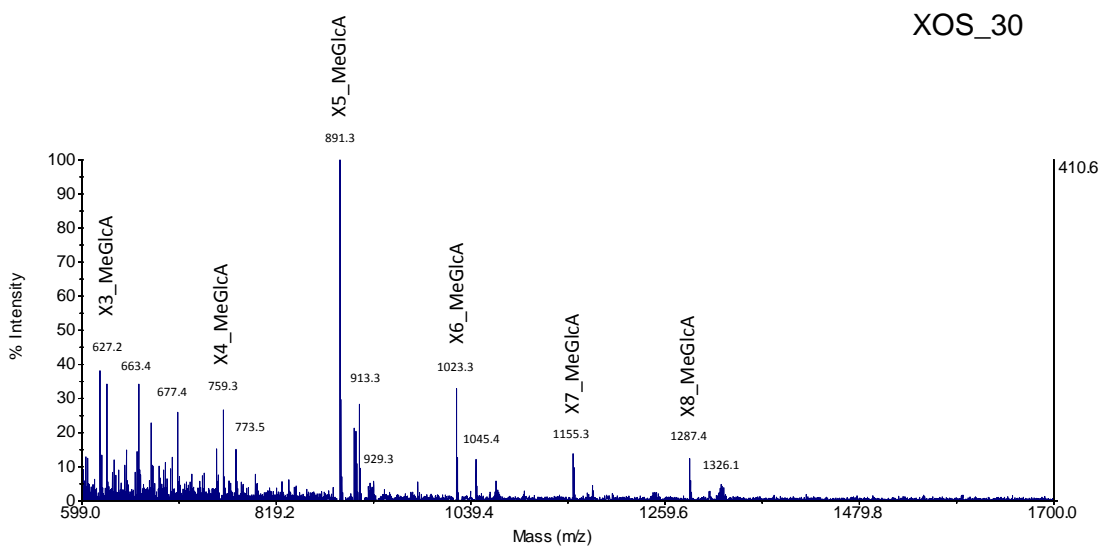
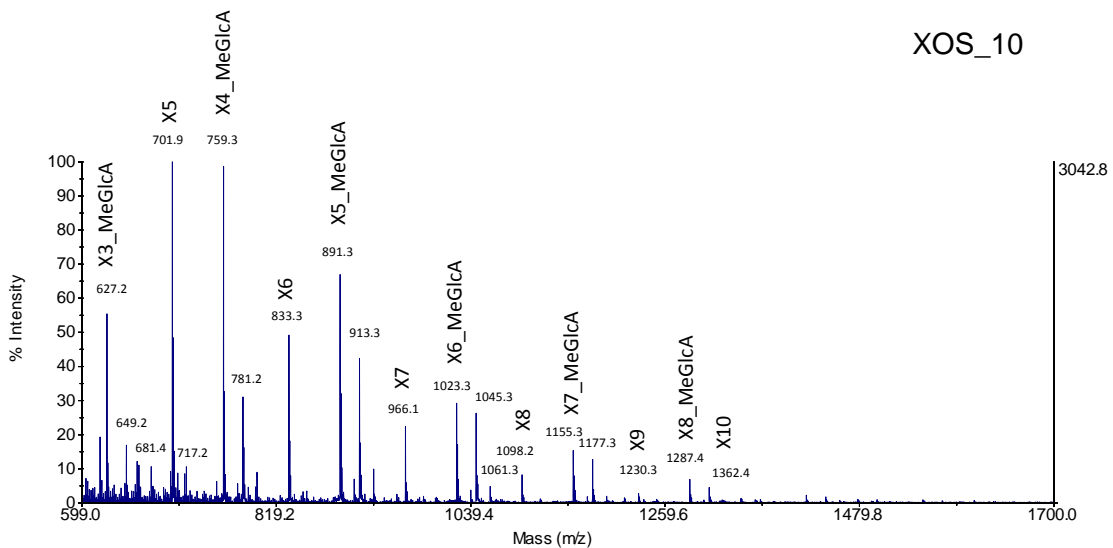
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250 **Figure 1.** TLC analysis of XOS released from beechwood xylan by Xyn10A
 251 (XOS_10) or Xyn30D (XOS_30). Xylan was incubated with enzymes for 5 min
 252 (1) or 120 min (2). M) size markers of xylose (X), xylobiose (X2), xylotriose (X3),
 253 xylotetraose (X4), xylopentaose (X5), and xylohexaose (X6)

254

255 In order to better elucidate the structure of the XOS obtained, they were
 256 analysed by MALDI-TOF MS (Fig. 2). The mass spectrum showed the presence
 257 of molecular ions of linear and substituted oligomers identified as sodium and
 258 potassium adducts (Valenzuela, Lopez, Biely, Sanz-aporicio, & Pastor, 2016).

259 XOS released by Xyn10A were linear oligomers consisting of 5 to 10
 260 xylopyranosyl (X5-X10) residues (Fig. 2), although X3 was also found.
 261 Moreover, acidic XOS containing 3 to 8 xylopyranosyl residues and a single
 262 MeGlcA residue (X3_MeGlcA to X8_MeGlcA), were also observed. In long term
 263 incubations xylobiose (X2) was also detected. The pattern found is in
 264 accordance with products pattern of family 10 xylanases, where the smaller
 265 products obtained are X2 and X3_MeGlcA (aldotetrauronic acid) (Biely, Singh,
 266 & Puchart, 2016).



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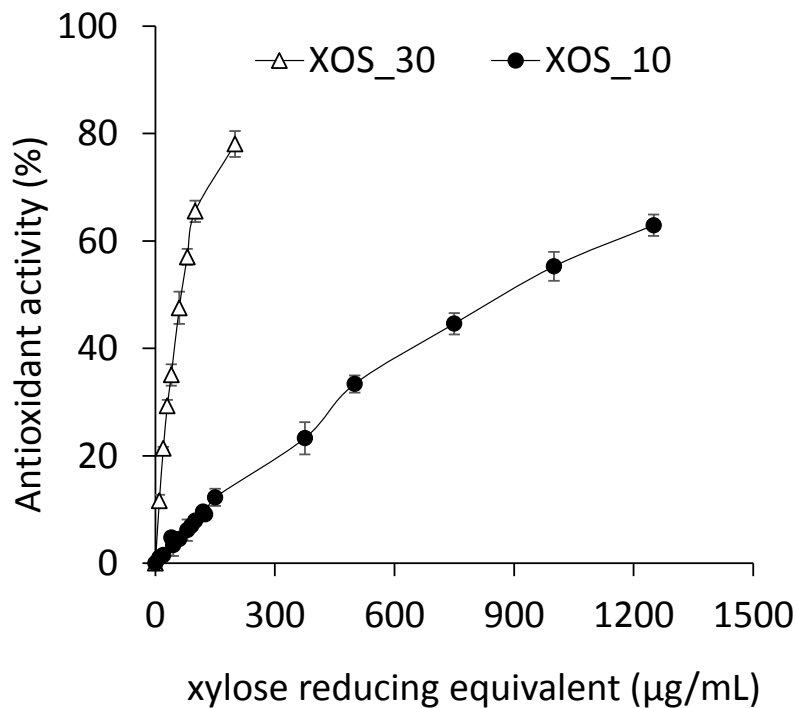
268 **Figure 2.** MALDI-TOF MS spectra of XOS_10 and XOS_30 from beechwood
269 xylan.

270 The mass spectra of products released by Xyn30D showed that all of
271 them are acidic oligomers containing a single MeGlcA residue (Fig. 2).
272 Substituted xylooligosaccharides consisting of 3 to 8 xylopyranosyl residues
273 (X3_MeGlcA to X8_MeGlcA) are shown, in accordance with TLC analysis.
274 Moreover, thin layer chromatograms shown in Fig. 1 revealed that X5_MeGlcA
275 and X6_MeGlcA are produced in higher amounts, followed by X7_MeGlcA and
276 X8_MeGlcA. The pattern found is in agreement with the mode of action of
277 family 30 xylanases, which have an absolute requirement of MeGlcA side
278 chains for hydrolysis of xylan, and release XOS substituted in the penultimate
279 xylose from the reducing end (Biely et al., 2016; Vršanská et al., 2007). The
280 structure of the main XOS_10 and XOS_30 obtained from beechwood xylan are
281 shown in supplementary Fig. S1.

282 **3.2. Antioxidant activity of XOS from beechwood xylan**

283 We analyzed the antioxidant capacity of the XOS produced from
284 beechwood xylan by Xyn10A and Xyn30D. As the methodology to evaluate
285 antioxidant activity we used quantification of the ABTS^{•+} radical discoloration.
286 This method confirmed the high antioxidant effect of well known compounds
287 such as Trolox and gallic acid, which exhibited a powerful antioxidant activity,
288 100% decoloration of ABTS^{•+} at 20 and 10 μM respectively (supplementary Fig.
289 S2). We measured the antioxidant activity of the XOS produced by Xyn10A and
290 Xyn30D at several concentrations (Fig. 3). In both cases the antioxidant activity
291 increased with XOS concentration, although the antioxidant activity of the two

292 types of XOS was clearly different. XOS_30 showed a much higher
293 effectiveness in reducing the ABTS^{•+} radical, much higher antioxidant activity
294 than XOS_10. At a concentration of 200 μg xylose reducing equivalents mL^{-1}
295 the antioxidant activity was around 80% in XOS_30 and 10% in XOS_10.
296



297

298 **Figure 3.** Antioxidant activity of XOS_10 and XOS_30 from beechwood xylan.

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300 The antioxidant activity of a compound can be compared to that of
301 Trolox, obtaining the TEAC (trolox equivalent antioxidant activity) value (Cusola
302 et al., 2015; Re et al., 1999). However, this value is determined according to the
303 molar concentration of the compound. Since the evaluated XOS include
304 oligomers of different length and molecular weight, the molar concentration
305 cannot be measured. As detailed above we can associate the antioxidant

306 activity value of a XOS sample to its concentration expressed as $\mu\text{g mL}^{-1}$ of
307 xylose reducing equivalents, and refer this value to the Trolox concentration
308 needed to obtain the same antioxidant activity. Thus, $100 \mu\text{g mL}^{-1}$ of XOS_30
309 produce the same antioxidant activity than Trolox at $13.2 \mu\text{M}$ whereas the
310 antioxidant activity of XOS_10 at the same $100 \mu\text{g mL}^{-1}$ concentration
311 corresponds to $3.2 \mu\text{M}$ Trolox.

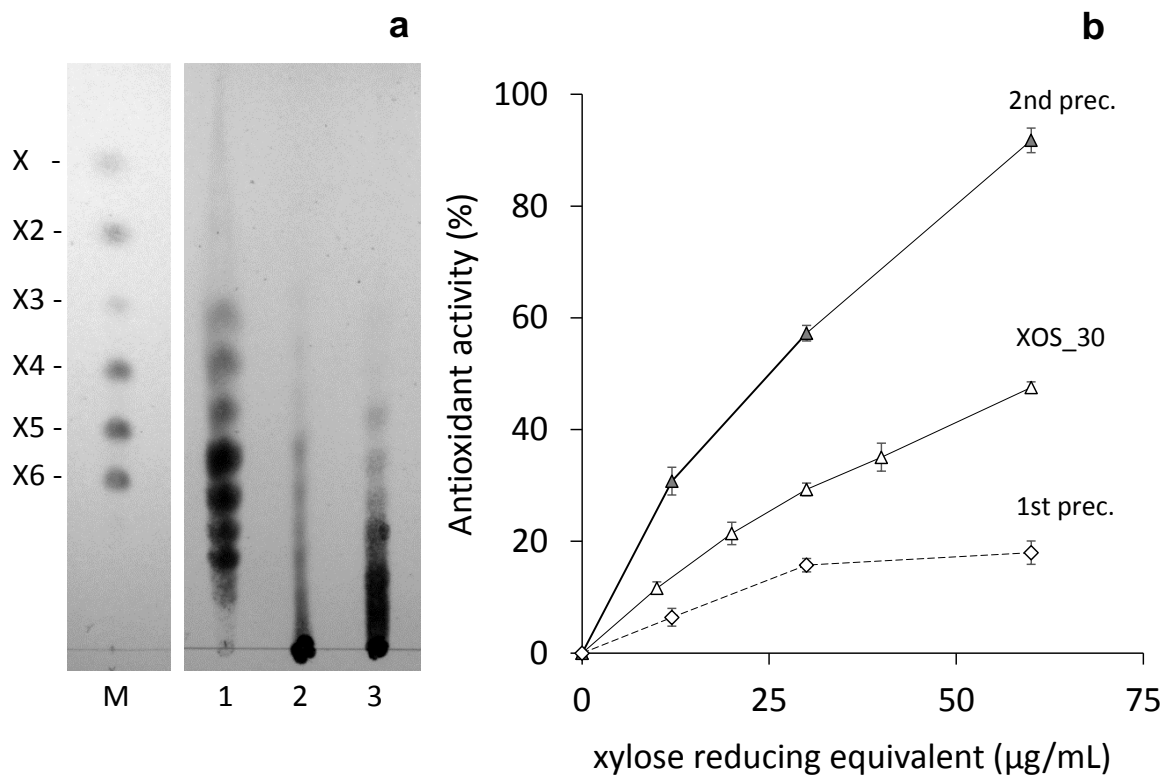
312 Although the antioxidant capacity of XOS produced by xylanases has
313 been reported in several works, to our knowledge, differences in the antioxidant
314 effect of XOS obtained by different families of xylanases has not been
315 described previously. The XOS described in literature come from different raw
316 materials such as sugarcane bagasse (Bian et al., 2013; Mandelli et al., 2014),
317 corncob (Gowdhaman & Ponnusami, 2015), garlic straw (Kallel, Driss,
318 Chaabouni, & Ghorbel, 2014), wheat bran (Lasrado & Gudipati, 2014), wheat
319 aleurone (Malunga & Beta, 2015), sunflower stalk and wheat straw (Akpinar,
320 Erdogan, et al., 2010), or agricultural residues (Rashad et al., 2016). However,
321 to our knowledge this is the first report on the antioxidant properties of XOS
322 from glucuronoxylans, xylans without ferulic acid ramifications.

323 Xylanases randomly hydrolyse xylan chains resulting in XOS of varying
324 degrees of polymerisation and substitution depending on the family of GH used.
325 TLC and MALDI-TOF have demonstrated a different XOS pattern depending on
326 the xylanase used, being those produced by Xyn10A a mixture of neutral and
327 acidic XOS whereas those produced by Xyn30D only acidic XOS. Since neither
328 xylose, neutral XOS (X2-X6) or glucuronic acid, at a concentration range of 200-
329 $1000 \mu\text{g mL}^{-1}$ exhibited antioxidant activity, it could be concluded that the

330 presence of the MeGlcA substitutions determines the antioxidant activity effect
331 found on XOS. In fact, Rao & Muralikrishna (2006) showed that the presence of
332 sugars with uronyl or acetyl groups impart strong antioxidant activity to cereal
333 polysaccharides. It has been also reported that carboxyl groups increase
334 antioxidant activity of cell wall polysaccharides (Pristov, Mitrovi, & Spasojevi,
335 2011). In a similar way, Malunga & Beta, (2015) reported that the antioxidant
336 capacity of XOS obtained from arabinoxylan is correlated to their degree of
337 ferulic acid substitution.

338 **3.3. Effect of the XOS length on the antioxidant activity**

339 Apart of substitution, other factors, such as XOS length, can affect the
340 antioxidant property. XOS samples studied up to this point had been depleted of
341 undigested or partially digested xylans by graded precipitation with three
342 volumes of ethanol. This methodology can also be used to separate XOS of
343 different degrees of polymerisation (Malunga & Beta, 2015). Analysis of a first
344 precipitate obtained by the addition of one volume of ethanol, showed that it
345 mainly contained unhydrolysed xylan and exhibited very low antioxidant activity
346 (Fig. 4a, b). A second precipitate was obtained by the subsequent addition of
347 two more volumes of ethanol. It mainly contained very long xylooligomers,
348 which showed high antioxidant activity. Remarkably this XOS fraction showed
349 considerably higher antioxidant activity than that of the supernatant, which
350 correspond to the purified XOS fraction (XOS_30), which showed lower degree
351 of polymerization (Fig. 4a, b). The results indicate that the antioxidant activity of
352 XOS_30 is also affected by the degree of polymerization. Similar results were
353 obtained by grade ethanol precipitation of XOS_10 (data not shown).



355

356 **Figure 4.** Analysis of XOS_30 from beechwood xylan. a) TLC analysis. 1,
 357 XOS_30; 2, first ethanol precipitate; 3, second ethanol precipitate; M, size
 358 markers of xylose (X), xylobiose (X2), xylotriose (X3), xylotetraose (X4),
 359 xylopentaose (X5), and xylohexaose (X6). b) Antioxidant activity of XOS_30,
 360 first and second ethanol precipitates.

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362 The time length of the enzyme treatments can affect to the degree of
 363 polymerization of the released XOS. We analyzed the antioxidant activity of the
 364 purified XOS obtained after different incubation time (from 1 to 120 min). Since
 365 XOS content was different at each time, to better visualize the results, the
 366 antioxidant activity of the different samples was divided by the amount (µg) of
 367 xylose reducing equivalents in each of them (Table 1). The values of antioxidant

368 activity oscillated from 0.32 to 0.16 for XOS_10 and from 1.41 to 1.20 for
 369 XOS_30. The results confirm the higher antioxidant activity of the XOS
 370 produced by family GH30 xylanases previously found. Comparing samples of 5
 371 and 120 min enzyme incubation, in the case of Xyn10A, antioxidant activity was
 372 reduced 50% with time. This can be caused by the diminished degree of
 373 polymerization of the XOS_10 obtained after 120 min, as it can be seen by TLC
 374 (Fig. 1). On the other hand, with Xyn30D, the antioxidant activity of XOS_30 did
 375 not show a significant variation with time. In fact, the same pattern of the XOS
 376 obtained after 5 and 120 min incubation with Xyn30D was observed by TLC
 377 (Fig.1).

378

379 **Table 1** Antioxidant activity of XOS obtained after different incubation time.

380 Beechwood xylan was treated with Xyn10A or Xyn30D and the XOS released at
 381 different incubation time were purified and analyzed.

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| | % Antioxidant activity μg^{-1} xylose reducing equivalent | |
|------------|---|--------|
| Time (min) | XOS_10 | XOS_30 |
| 1 | 0.32 | 1.41 |
| 5 | 0.36 | 1.43 |
| 10 | 0.29 | 1.26 |
| 30 | 0.17 | 1.59 |
| 120 | 0.16 | 1.20 |

395

Our results clearly show that apart of ramifications, the degree of

396

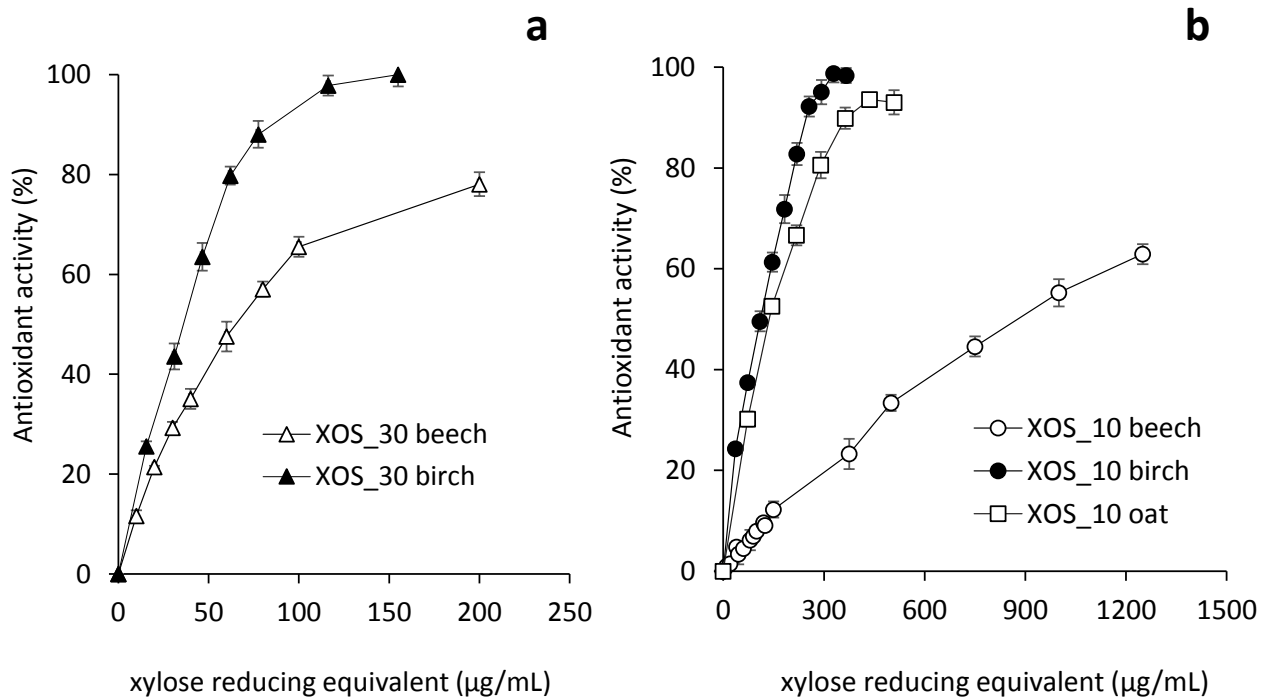
polymerization of XOS obtained from glucuronoxylans affects the antioxidant

397 activity. In fact, the increase of the antioxidant activity with the molecular weight
398 had been previously reported for XOS released from arabinoxylans by Malunga
399 & Beta, (2015) and Rao & Muralikrishna, (2006).

400 **3.4. Antioxidant activity of XOS from birchwood and oat spelt xylans**

401 Once made it evident the antioxidant activity of XOS obtained from
402 beechwood xylan, XOS produced from other glucuronoxylans, such as
403 birchwood xylan, and from arabinoxylans, such as oat spelt xylan, were also
404 evaluated. These xylans were hydrolyzed by xylanases, the XOS obtained after
405 2h of treatment were purified and their antioxidant activity was analysed at
406 increasing concentrations. As shown in Fig. 5 (a and b) the XOS obtained from
407 birchwood xylan showed higher antioxidant activity than those obtained from
408 beechwood xylan for both xylanases, although the difference was higher for
409 Xyn10A. These glucuronoxylans are very similar although they differ in their
410 number of methyl glucuronic acid ramifications, with higher content in birchwood
411 xylan (Hespell & Cotta, 1995). Differences in antioxidant activity may also result
412 from the different size of the XOS produced by the enzymes from the different
413 glucuronoxylans. Since Xyn30D is a glucuronoxylan-specific xylanase, and it
414 does not hydrolyse arabinoxylans (Valenzuela et al., 2012) this enzyme was not
415 tested on oat spelt xylan. XOS produced from oat spelt xylan by Xyn10A
416 showed similar antioxidant activity than those produced from birchwood xylan,
417 and much higher than those produced from beechwood xylan. This effect can
418 be justified by the presence of ferulic acid substitutions in arabinoxylans from
419 oat spelt. In fact, high antioxidant activity has been reported in water-soluble
420 arabinoxylans (feraxans) (Rao & Muralikrishna, 2006), isolated from rice and

421 ragi. The phenolic content of oligosaccharides has been proposed to have an
 422 important effect in antioxidant activity (Bijalwan, Ali, Kesarwani, Yadav, &
 423 Mazumder, 2016; Rashad et al., 2016). On the other hand, in beechwood and
 424 birchwood XOS, since they do not contain ferulic acid, the main reason of their
 425 antioxidant capacity was the presence of methyl glucuronic acid ramifications.

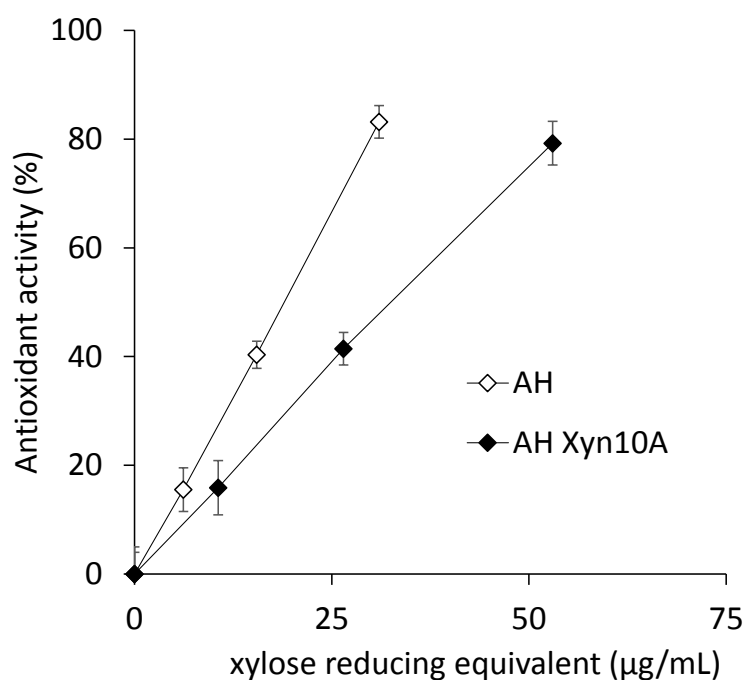


426
 427 **Fig. 5** Antioxidant activity of XOS from beechwood, birchwood and oat spelt
 428 xylans. a) XOS_30. b) XOS_10

429
 430 **3.5. Antioxidant activity of XOS from eucalyptus autohydrolysates**

431 Pulping process in paper industry produces an enormous amount of wastes
 432 containing an important quantity of dissolved xylans among other wood
 433 components. Evaluation of the antioxidant activity of XOS derived from
 434 eucalyptus xylan can make it possible its upgrading to added value products. As

435 a first approach we tested the antioxidant activity of XOS extracted from
436 eucalyptus wood by the autohydrolysis process. In this process, biomass is
437 treated with hot water or steam and hemicelluloses are converted into soluble
438 glucuronoxytan polymers and long oligosaccharides (Gullón et al., 2011).
439 Eucalyptus autohydrolysates showed high antioxidant activity, probably due to
440 the presence of long XOS together with some quantity of lignin (Fig. 6).
441 However, when autohydrolysates were treated with Xyn10A for 1 h antioxidant
442 activity was strongly decreased. A similar value of around 80% antioxidant
443 activity required twice the amount of xylose reducing equivalents when treating
444 with the enzyme. TLC analysis showed a high amount of XOS of high molecular
445 weight (higher than X6) in the initial AH whereas XOS of lower molecular weight
446 appeared after treatment with Xyn10A (data not shown). As previously stated,
447 the XOS length is an important factor for their antioxidant activity. Concerning
448 Xyn30D, the enzyme did not show noticeable enzymatic activity on eucalyptus
449 autohydrolysate, probably as a result of the heavy acetylation of xylan in the
450 autohydrolysate or to the lack of accessibility of the enzyme, of bigger size than
451 Xyn10A, to the substrate. So the effect of Xyn30D could not be tested on
452 eucalyptus hydrolysate.



453

454 **Fig. 6** Antioxidant activity of autohydrolysate liquors from eucalyptus wood.

455 Autohydrolysates without enzyme treatment (AH) or treated with Xyn10A (AH

456 Xyn10A).

457

458 To compare the antioxidant effect of XOS from eucalyptus hydrolysates

459 with that of XOS obtained by enzyme action on xylans, the antioxidant activity

460 per µg of xylose reducing equivalents of the different XOS studied was

461 determined (Supplementary Table S1). Although eucalyptus hydrolysates

462 showed the highest antioxidant activity (2.65), a rather similar activity was found

463 in XOS_30 from birchwood xylan (1.30), while XOS_10 from beechwood

464 showed the lowest activity (0.08).

465 Akpinar, Gunay, Yilmaz, Levent, & Bostanci, (2010) found that liquors

466 from autohydrolysis of agricultural residues had antioxidant activity while

467 González, Cruz, Domínguez, & Parajó, (2004) and Rivas, Conde, Moure,
468 Domínguez, & Parajó, (2013) reported antioxidant effect in liquors from
469 eucalyptus wood autohydrolysis. However, no knowledge exist concerning the
470 effect of xylanases on the antioxidant activity of eucalyptus autohydrolysates.

471 **3.6. Antioxidant activity: ABTS vs. DPPH**

472 Several methods exist for the measure of antioxidant activity. Concerning
473 the antioxidant activity of XOS, several works use the DPPH method (Akpinar,
474 Gunay, et al., 2010; Bian et al., 2013; Gowdhaman & Ponnusami, 2015;
475 Hromádková, Paulsen, Polovka, Košťálová, & Ebringerová, 2013; Kallel et al.,
476 2014; Lasrado & Gudipati, 2014; Veenashri & Muralikrishna, 2011) although
477 other reports use the ABTS method (Malunga & Beta, 2015; Rivas et al., 2013).
478 Both, DPPH and ABTS methods consist on radical scavenging assays whose
479 reaction mechanism involves transfer of electron by the reducing agent to the
480 DPPH/ABTS radical. The discoloration produced by the interaction of this
481 radical with hydrogen or electron donor species is quantified.

482 In this work the antioxidant activity of XOS produced by xylanases from
483 beechwood was also analysed by the DPPH method. However, some difficulties
484 were observed with this method. The DPPH radical has to be firstly dissolved in
485 ethanol or methanol to perform the measurements. When XOS were mixed with
486 this solution, some XOS precipitated giving an erroneous lecture of the
487 absorbance, and making the method less sensitive or not suitable for XOS
488 determination. The concentration that produced the 50% of inhibition of the
489 radicals was measured, being determined as 13157 μg of xylose reducing
490 equivalent mL^{-1} for XOS_10 with DPPH method, or 850 of μg of xylose reducing

491 equivalent mL⁻¹ with ABTS. In the case of XOS_30 these values were 417 or 70
492 µg of xylose reducing equivalent mL⁻¹ with DPPH or ABTS, respectively.
493 Although the higher effectiveness of XOS_30 was shown by both methods, the
494 concentration needed to inhibit DPPH was strongly higher, indicating a higher
495 sensitivity of the ABTS method for XOS studies. Rivas et al., (2013) analysed
496 the antioxidant activity of autohydrolysates liquors and also found higher
497 antioxidant activity with ABTS than with DPPH. In reported works, where the
498 antioxidant activity of XOS from arabinoxylans has been evaluated by the
499 DPPH method, 50% of inhibition has been found with 1000 µg mL⁻¹ of XOS
500 from corncob (Gowdhaman & Ponnusami, 2015) or with 620 µg mL⁻¹ of
501 sugarcane bagasse derived XOS. Although these reported values for producing
502 50% DPPH inhibition cannot be compared to those obtained in our work,
503 obtained from different xylan substrates, our results indicate that the important
504 antioxidant activity we have identified in MeGlcA branched XOS can be
505 compared to that of arabinoxylan derived XOS, ferulic acid branched, of high
506 potential as antioxidants for food industry.

507 **4. CONCLUSIONS**

508 XOS produced by enzymatic hydrolysis of glucuronoxylan showed high
509 antioxidant activity. XOS produced by a family GH30 xylanase (Xyn30D),
510 comprised exclusively of acidic XOS, showed higher antioxidant activity than
511 those produced by xylanase Xyn10A of family GH10, a mixture of neutral and
512 acidic XOS. The MeGlcA ramification contained in acidic XOS seems to be an
513 important determinant of the antioxidant power of xylooligomers. Besides,
514 antioxidant activity increased with XOS degree of polymerization. Eucalyptus

515 wood autohydrolysate showed high antioxidant activity, which was diminished
516 by xylanase treatment. ABTS was a more accurate methodology for the
517 antioxidant activity determination of XOS than DPPH.
518
519

520 **ACKNOWLEDGEMENTS**

521 This work was funded by the Spanish MINECO FILMBIOCEL CTQ2016-77936-
522 R (funding also from the “Fondo Europeo de Desarrollo Regional FEDER”),
523 BIOPAPμFLUID CTQ2013-48995-C2-1-R and CTQ2013-48995-C2-2-R and
524 MICROBIOCEL CTQ2017-84966-C2-1-R projects. Special thanks are also due
525 to the consolidated research group AGAUR 2017 SGR 30 with Universitat de
526 Barcelona and to the Serra Húnter Fellow to Cristina Valls. We thank J.C.
527 Parajó for his generous gift of eucalyptus autohydrolysates and Alejandro Costa
528 for his technical support.

529

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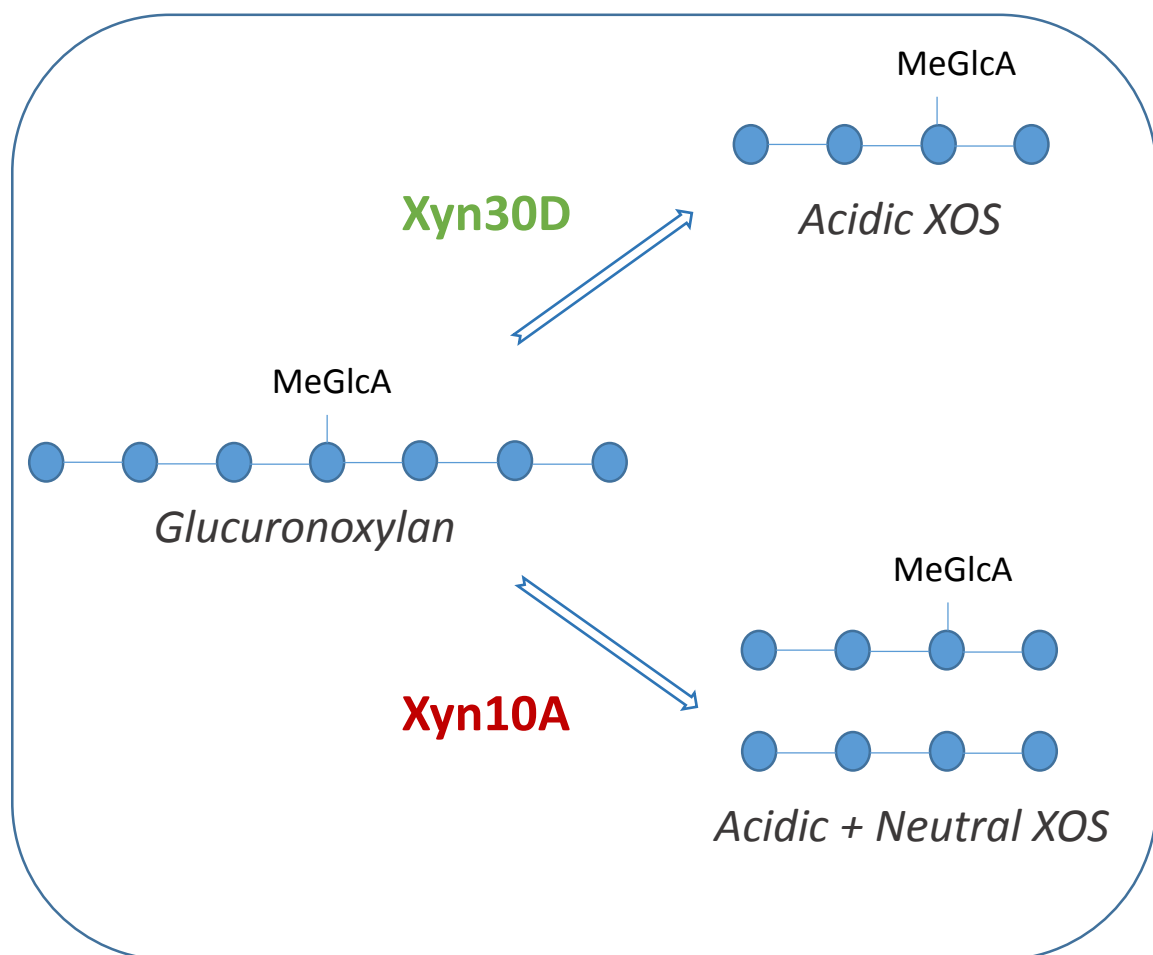
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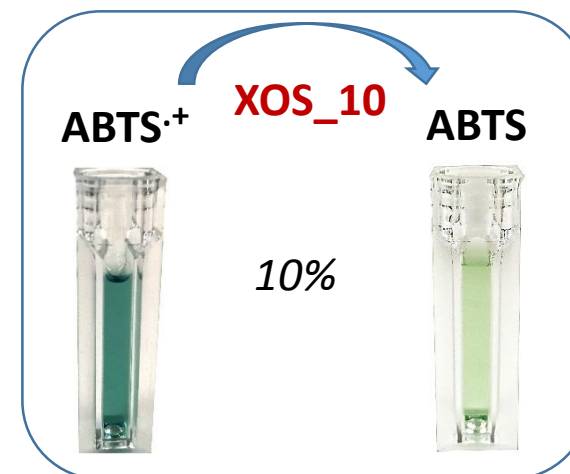
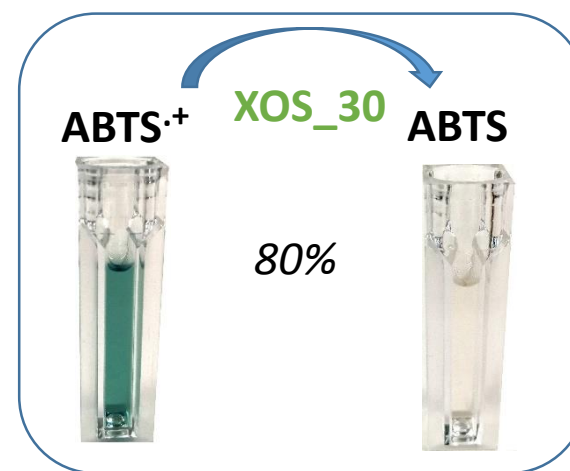
Supplementary data

[Click here to download Supplementary data: SUPPLEMENTARY INFORMATION JC.docx](#)

XOS Enzymatic Production



Antioxidant activity at 200 µg xylose reducing equivalent mL⁻¹



- Increased by:
- *Acidic XOS*
 - *XOS length*
 - *Lignin*

Eucalyptus autohydrolysates (AH)

