

# In vitro degradation of covalently cross-linked arabinoxylan hydrogels by Bifidobacteria

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#### 1 Highlights

- 2 Covalently crosslinked arabinoxylans (AX) gels with different structure were obtained
- 3 AX gels were biodegradable by mixture of two *Bifidobacterium*
- 4 Changes in microstructure of AX gels allowed different degradation profiles
- 5 AX gels could be used as microflora-activated system for colon delivery
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# 21 Bifidobacteria 22 23 Ana L. Martínez-López<sup>a</sup>, Elizabeth Carvajal-Millan<sup>a,\*</sup>, Valérie Micard<sup>b</sup>, Agustín Rascón24 Chu<sup>a</sup>, Francisco Brown-Bojorquez<sup>c</sup>, Norberto Sotelo-Cruz<sup>c</sup>, Yolanda L. López-Franco<sup>a</sup>,

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#### 37 Abstract

Arabinoxylan gels with different cross-linking densities, swelling ratios, and rheological properties were obtained by increasing the concentration of arabinoxylan from 4 to 6% (w/v) during oxidative gelation by laccase. The degradation of these covalently cross-linked

gels by a mixture of two Bifidobacterium strains (Bifidobacterium longum and 41 Bifidobacterium adolescentis) was investigated. The kinetics of the evolution of structural 42 morphology of the arabinoxylan gel, the carbohydrate utilization profiles and the bacterial 43 production of short-acid fatty acid (SCFA) were measured. Scanning electron microscopy 44 analysis of the degraded gels showed multiple cavity structures resulting from the bacterial 45 action. The total SCFA decreased when the degree of cross-linking increased in the gels. A 46 slower fermentation of arabinoxylan chains was obtained for arabinoxylan gels with more 47 48 dense network structures. These results suggest that the differences in the structural features 49 and properties studied in this work affect the degradation time of the arabinoxylan gels.

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51 Keywords: cross-linked arabinoxylans, gels, biodegradability, bifidobacteria, fermentation

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#### 53 1. Introduction

Polysaccharide-based hydrogels have been extensively studied for their potential as colon-54 55 specific drug delivery systems due to their chemical and three-dimensional structures, biocompatibility properties and swelling (Peppas, 1997; Sinha & Kumria, 2001). However, 56 most of these hydrogels are susceptible to changes according to the low gastrointestinal pH 57 or the duration of transit time, thus leading to premature drug release. For these reasons, 58 some polysaccharides such chitosan, alginate and pectin have been chemically modified or 59 cross-linked or combined with hydrophobic polymers for use in the development of 60 systems that are specifically biodegradable by colonic bacteria (Brøndsted, Andersen, & 61 62 Hovgaard, 1998; Chambin, Dupuis, Champion, Voilley, & Pourcelot, 2006; Tozaki et al.,

olymers, 144 (76

1997). Covalent cross-linking has been used to enhance the mechanical strength and 63 chemical stability of the hydrogels. However, the chemical cross-linking agents used are 64 often toxic compounds that require harsh reaction conditions. Another alternative is to use 65 polysaccharides, which can form covalent hydrogels via enzymatic cross-linking. 66 Biological enzymes can catalyze polysaccharide cross-linking into gels via strong covalent 67 bonds under mild conditions in the absence of organic solvents (Figueroa-Espinoza et al., 68 1999; Micard & Thibault, 1999). Unlike most polysaccharides, arabinoxylans (AX) can 69 form covalent hydrogels via enzymatic cross-linking of ferulic acid esterified to the 70 71 polysaccharide (Izydorczyk & Biliaderi, 1995; Vansteenkiste, Babot, Rouau, & Micard, 2004). Covalently cross-linked AX gels are thermo-irreversible (Carvajal-Millan, 72 Guigliarelli, Belle, Rouau, & Micard, 2005), generally present an absence of pH or 73 electrolyte susceptibility and exhibit no syneresis after long periods of storage (Izydorczyk 74 75 & Biliaderis, 1995).

AX are non-starch polysaccharides that resist digestion and absorption in the human small 76 intestine and are fermented in the large intestine by microbiota in the colon, especially 77 Bacteroides and Bifidobacterium (Hopkins et al., 2003; Hughes et al., 2007). The basic 78 structure of AX consists of a linear backbone chain of xylose units containing arabinose 79 substituents attached through O-2 and/or O-3 (Izydorczyk & Biliaderis, 1995). Some of the 80 81 arabinose residues are ester-linked to ferulic acid (FA) through (O)-5 (Smith & Hartley, 1983). FA oxidation by enzymatic free radical generating agents (laccase or peroxidase/ $H_2O_2$ ) 82 allows for the coupling of AX chains through the formation of FA dimers and trimers (di-83 FA, tri-FA) as covalent cross-linking structures (Carvajal-Millan, Landillon, et al., 2005; 84 Izydorczyk & Biliaderis, 1995). Recent studies have demonstrated that AX gels formed via 85 oxidative cross-linking can be used for the controlled release of model proteins (Carvajal-86

Millan, Guilbert, Doublier, & Micard, 2006), methylxanthine (Iravani, Fitchett, & Georget,
2011), diclofenac sodium (Iqbal, Akbar, Hussain, Saghir, & Sher, 2011), lycopene
(Hernández-Espinoza, Piñón-Muñiz, Rascón-Chu, Santana-Rodríguez, & Carvajal-Millan,
2012) and probiotic bacteria (Morales-Ortega et al., 2014).

The degradation of the AX structure results from the activity of different bacterial enzymes, 91 92 which mainly include endo-xylanases, beta-xylosidases, reducing end xylose-releasing exooligoxylanases and alpha-L-arabinofuranosidases (Pollet et al., 2012). The mechanism of 93 AX degradation depends on the degree of polymerization, crosslinking and branching 94 present in the AX structure (Hughes et al., 2007; Rose, Patterson, & Hamaker, 2010). 95 Notably, bifidobacteria comprise up to 25% of the cultivable gut microflora, with 96 97 Bifidobacterium longum and Bifidobacterium adolescentis predominating in adults. 98 Mixtures of these strains have been associated with the complete fermentation of low molecular weight arabinoxylans (Pastell, Westermann, Meyer, Tuomainen, & Tenkanen, 99 2009). Though the fermentation properties of arabinoxylans have been researched 100 extensively, only a few studies have reported the degradation of the covalently cross-linked 101 arabinoxylan network, and these studies did not report the structural features, cross-linking 102 103 density and mechanical properties of the gel (Hopkins et al., 2003). These structural 104 parameters of AX matrix gel could influence the ability of colonic microflora to degrade polysaccharides. In the present study, the ability of a mixture of *Bifidobacterium longum* 105 and Bifidobacterium adolescentis to degrade covalent arabinoxylan hydrogels with 106 different swelling and structural parameters was examined. 107

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#### 109 **2. Experimental**

#### 110 *2.1. Materials*

111 Maize bran arabinoxylans (MBAX) were obtained and characterized as previously reported

112 by Martínez-López et al. (2012). MBAX contained 85% dry basis (d.b.) of pure AX,

contained ferulic acid (FA), di-FA, and tri-FA contents of 0.25, 0.14, and 0.07 µg/mg of
MBAX, respectively, and an A/X ratio of 0.72. Laccase (benzenediol: oxygen
oxidoreductase, E.C.1.10.3.2) was isolated from *Trametes versicolor* and all other chemical

116 products were purchased from Sigma Chemical Co. (St. Louis, MO, USA).

117 2.2. Preparation of MBAX gels

118 MBAX solutions at 4% (MBAX-4) and 6% (MBAX-6) (w/v) were prepared in 0.1 M 119 acetate buffer (pH 5.5). The buffer was filtered through a 0.2  $\mu$ m filter to prevent microbial 120 contamination. Laccase (1.675 nkat/mg of MBAX) was used as a cross-linking agent. The 121 gels were allowed to develop for 6 h at 25 °C.

#### 122 2.3. Rheological properties

The rheological tests were performed by small amplitude oscillatory shearing using a strain-controlled rheometer (Discovery HR-2 rheometer, TA Instruments, New Castle, DE, USA) as reported by Vansteenkiste et al. (2004). MBAX gelation was studied for 6 h at 25 °C. All measurements were carried out at a frequency of 0.25 Hz and 5% strain (linearity range of viscoelastic behavior). The mechanical spectra of gels were obtained by conducting a frequency sweep from 0.01 to 100 Hz at 25 °C.

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#### 131 2.4. Phenolic Acids Content

The FA, di-FA and tri-FA contents in the MBAX gels were quantified by RP-HPLC after a
de-esterification step as described by Vansteenkiste et al. (2004). An Alltima (Alltech,
Deerfield, IL, USA) C18 column (250 × 4.6 mm) and a photodiode array detector Waters
996 (Millipore Co., Milford, MA, USA) were used. Detection was followed by UV
absorbance at 320 nm.

137 2.5. Swelling and structural parameters

The MBAX-4 and MBAX-6 were allowed to swell as described by Carvajal-Millan, Landillon, et al. (2005). The equilibrium swelling was reached when the weight of the samples changed by no more than 3% (0.06 g). The swelling ratio (q) was calculated as follows:

142  $q = (Ws - W_{MBAX})/W_{MBAX}$ ,

where Ws is the weight of swollen gels and  $W_{MBAX}$  is the weight of the MBAX-4 or MBAX-6 (w/v) gels. From the swelling measurements, the molecular weight between two cross-links (M<sub>c</sub>), the cross-linking density ( $\rho_c$ ) and the mesh size ( $\xi$ ) values of the MBAX-4 and MBAX-6 gels were calculated as reported by Carvajal-Millan, Guilbert, Morel, & Micard, 2005, using the model of Flory & Rehner (1943) modified by Peppas & Merrill (1976) for gels in which the cross-links are introduced in solution.

149 2.6. Scanning electron microscopy

The microstructure of the MBAX-4 and MBAX-6 gels and surface morphology of the residual MBAX gels after 18 and 36 h fermentation by a mixture of *Bifidobacterium longum* and *Bifidobacterium adolescentis* were studied by scanning electron microscopy

153 (SEM) (JEOL JSM-7401F, Peabody, MA, USA) without coating at low voltage (10 kV).

154 SEM images were obtained in secondary and backscattered electron image modes.

155 2.7. Degradation of MBAX gels

156 2.7.1. Organism and culture conditions

The bacterial strains used (*Bifidobacterium adolescentis* ATCC 15703 and *Bifidobacterium longum* ATCC 15708) were purchased from American Type Culture Collection (Manassas, VA, USA). The bacteria were maintained at -80 °C in glycerol stock solutions. The bacterial suspensions were thawed, and the bacteria were revived by culturing in the basal media as described previously by Hughes et al. (2007). The strains were incubated in the broth under anaerobic conditions at 37 °C until an optical density of one (OD= 1) was reached. Afterward, they were used as an inoculum for growth studies on MBAX gels.

#### 164 2.7.2. In vitro fermentation

For the fermentation analysis, the MBAX-4 and MBAX-6 gels were formed in Hungate 165 tubes and autoclaved at 121 °C for 15 min. MBAX without gelation was used as a positive 166 control. The MBAX and MBAX gels were mixed with autoclaved basal media to give a 167 final concentration of 5 g/L (w/v) MBAX as a sole carbon source. 100 µL Oxyrase 168 (Oxyrase Inc. Mansfield, OH, USA) was added as the broth to remove oxygen from the 169 microenvironment of the test tube. The tubes were sealed anaerobically by flushing the 170 headspace with carbon dioxide and placed at 4 °C overnight. These Hungate tubes were 171 inoculated with a 1% (v/v) mixture of two bifidobacteria species (B. adolescentis and B. 172 longum; ratio 1:1) and incubated in the dark without shaking at 37 °C for 18, 36 and 72 h. 173 174 Fermentations were carried out in triplicate, and duplicate blank tubes containing no

substrate were included for each time point. The OD (650 nm) was measured at 18, 36 and
72 h after inoculation. The pH was measured at the end of each different growth period, and
random samples were examined microscopically to exclude contamination by the bacterial
cultures.

179 2.7.3 Fermentation product analysis

Before analysis, at time 0 of fermentation, one milliliter of each MBAX gel was treated 180 with 0.5 mL 0.5 N NaOH for 15 min at 25°C to disrupt the gels by releasing the phenolic 181 182 bridges. The pH was then immediately adjusted to 7 to prevent MBAX chain degradation. The samples for the other fermentation time points were only centrifuged (15,000 g, 4 °C, 5 183 min) to remove the bacteria. The changes in the molecular weight distribution of the 184 MBAX and MBAX gels at different times of fermentation (0, 18, 36, 72 h) were 185 determined by SE-HPLC at 38 °C using a TSKGel G5000PWXL column (7.8 x 300 mm) in 186 combination with a PWX-guard column (6 x 40 mm) (Tosoh Bioscience, Stuttgart, 187 188 Germany). Molecular weights were estimated after universal calibration with pullulans as standards (P50 to P800). Isocratic elution was conducted at 0.6 mL/min with 0.01 M LiNO<sub>3</sub> 189 filtered through 0.2 µm. Differential refractometry was used for detection. 190

191 Residual carbohydrates in the freeze-dried fermentation residues and monosaccharides in 192 the supernatant of the centrifuged samples were quantified by RP-HPLC as described by 193 Carvajal-Millan et al. (2007). The main monosaccharides of MBAX (arabinose, xylose, 194 galactose and glucose) were used as standards.

Short-chain fatty acids (SCFA) were analyzed based on Pastell, Westermann, Meyer,
Tuomainen, & Tenkanen (2009). The SCFA analysis was carried out using GC (Clarus
580, PerkinElmer, Waltham, MA, USA) with a flame ionization detector and a capillarity

198 column (Elite-FFAP 30 m x 0.50 mm I.D; film thickness, 1  $\mu$ m). The SCFA in the samples 199 was quantified using external calibration curves of acetic, propionic and butyric acids, and 200 the results were expressed in  $\mu$ g/mL.

201 2.8. Statistical analysis

The results are expressed as the means  $\pm$  Standard Deviation (S.D.) from three replicates. The significance of differences was determined by analysis of variance (ANOVA) (OriginPro software, version 8.6. Originlab Corporation, USA) followed by Tukey's test and the results were considered to be statistically significant when p< 0.05.

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#### 207 3. Results and discussion

208 3.1. Features of MBAX gels

#### 209 3.1.1 Rheological properties and covalent cross-links

The formation of gel networks was monitored by storage (G') and loss (G") modulus 210 changes in 4 and 6% (w/v) AX solutions undergoing oxidative gelation by laccase (1.675 211 212 nkat/mg of MBAX) (Fig. 1a). The gelation profile vs. time exhibited an initial increase of 213 the G' modulus, followed by a plateau region resulting from the sol-gel transition. By increasing the MBAX concentration, the gelation time (time G' = G'', i.e., tan delta= 1) 214 decreased and the gelation rate (rate of G' development) increased. This relationship could 215 be associated with a higher density of polysaccharide chains enhancing the probability of 216 contact between ferulic acid free radicals, which is an essential requirement for the 217 establishment of a three-dimensional AX network (Carvajal-Millan, Guilbert, Morel, & 218 219 Micard, 2005; Rattan, Izydorczyk, & Biliaderis, 1994). The G' values (215 Pa) of the

MBAX gel at 4% (w/v) were higher than those reported for laccase-induced maize bran gel 220 (20 Pa) or waste water arabinoxylans gel (2 Pa) at the same concentration, though MBAX-4 221 222 had a similar initial FA content to both of these gels (0.23 and 0.37 µg/mg AX, 223 respectively) (Hernández-Espinoza et al., 2012; Niño-Medina et al., 2009). Such behaviors might have their origin in the structural and conformational characteristics of AX 224 molecules. The increase in the MBAX concentration from 4 to 6% (w/v) in the gel resulted 225 in an approximately two-fold increase in the value of G' (Fig. 1a). These results were in 226 agreement with data reported previously by Rattan et al. (1994), who found linear 227 228 relationships between G' (plateau values) and polymer concentration in samples of wheat AX. This G' vs. polymer concentration trend has also been observed in chemically cross-229 linked alginate gels (Choudhary & Bhatia, 2012) and dextran-lactate gels cross-linked by 230 stereocomplex formation (De Jong et al., 2001). The mechanical strength of the matrix is 231 known to have a direct bearing on drug delivery characteristics (Coviello et al., 2005; 232 Woolfson, Malcolm, Campbell, Jones, & Russell, 2000). The mechanical spectra of the 233 MBAX-4 and MBAX-6 gels after 6 h of gelation are shown in Fig. 1b. Though the MBAX-234 6 gel exhibited a solid-like behavior with G' > G'' and a plateau towards the low-frequency 235 range (0.01 to 10 Hz), G' decreased steeply with increasing frequency (beyond 40 Hz). This 236 behavior is typical of strong gels that present rupture and fail at large deformations. The 237 238 MBAX-4 gel showed a linear G' independent of the frequency of oscillation and a G" that was drastically lower than G' and linearly dependent on the frequency (Ross-Murphy & 239 240 Shatwell, 1993). The dependence of G" on the frequency has been related to the participation of physical entanglement interactions in the maturation of the gel and the 241 various levels of connectivity and junction zones of disparate lengths and strengths within 242 the three-dimensional network (Kasapis, 2008; Picout & Ross-Murphy, 2003). Hence, these 243

differences can be attributed to the covalent cross-linking distribution and the physical
entanglement of AX chains in MBAX gels as a function of AX concentration. Similar
mechanical spectra have been previously reported for strong gels with potential uses as
carriers of proteins and drugs systems (Choh, Cross, & Wang, 2011; Choudhary & Bhatia,
2012; De Jong et al., 2001).



Fig. 1. (a) Monitoring the storage (G') and loss (G'') modulus of MBAX solution at 4 and
6% (w/v) during gelation by laccase (1.675 nKat/mg of MBAX) at 0.25 Hz. (b) Mechanical
spectrum of MBAX gels after 6 h gelation. Rheological measurements were carried out at
25°C and 5% strain.

The extent of covalent cross-linking in the MBAX gels was determined by the content of total di-FA and tri-FA after 6 h of gelation (Table 1). The amounts of di-FA and tri-FA produced during gelation never compensated for the loss of FA monomer. Indeed, at the end of gelation, 80 and 70% of the FA initially present in MBAX was oxidized; by

contrast, only 32 and 34% was recovered as a combination of di-FA and tri-FA in MBAX-6 258 and MBAX-4, respectively. Nevertheless, the tan  $\delta$  (G"/G') values confirmed the formation 259 260 of an elastic covalent system for both gels (Fig. 1a). These behaviors in MBAX gels have 261 been previously reported by several authors (Carvajal-Millan et al., 2007; Lapierre, Pollet, Ralet, & Saulnier, 2001; Niño-Medina et al., 2009). These authors attributed such 262 phenomena to the formation of higher ferulate cross-linking structures unreleased by mild 263 alkaline hydrolysis and/or to the participation of lignin residues coupled with FA 264 monomers in the formation of MBAX gels. At the end of gelation, the 8-5' form (especially 265 266 benzofuran) contributed approximately 68 - 79% of the total di-FA in MBAX-4 and MBAX-6, whereas the 5-5' and 8-O-4' forms contributed 22 - 12% and 10 - 9%, 267 respectively. The predominance of 8-5' di-FA was also observed in wheat flour AX treated 268 with a peroxidase/H<sub>2</sub>O<sub>2</sub> system or laccase (Carvajal-Millan, Landillon, et al., 2005; 269 Figueroa-Espinosa et al., 1999). Nevertheless, the 5-5' di-FA structure has been reported to 270 be predominant in AX gels from maize bran (Carvajal-Millan et al., 2007; Niño-Medina et 271 al., 2009). In a previous study, Hatfield & Ralph (1999) reported that only 5-5' di-FA can 272 be formed intramolecularly in arabinoxylans, i.e., on the same polysaccharide chains. In 273 this study, the greater proportion of the 5-5' di-FA formed in the MBAX-4 gel (22%) 274 compared with the MBAX-6 gel (12%) might have indicated that in MBAX-4, the 275 276 intrachain bonds were favored and thus the polymer network showed decreasing continuity. These results may partly explain some differences in the mechanical properties of the gels. 277

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	q <sup>a</sup>	$Mc^{b} \times 10^{3}$	$\rho_{\rm c}^{\rm c} \times 10^{-6}$	$\xi^{ m d}$	FA	di-FA
	$(g H_2O/g MBAX)$	(g/mol)	(mol/cm <sup>3</sup> )	(nm)	(µg/mg	MBAX)
MBAX-4	18.11 ± 1.23	$36 \pm 0.60$	$39\pm0.05$	$179 \pm 5.23$	$0.054 \pm 0.017$	$0.031 \pm 0.011$
MBAX-6	9.36 ± 1.86	$26 \pm 0.31$	$54 \pm 0.06$	$110\pm0.32$	$0.074 \pm 0.002$	$0.044\pm0.001$

#### **Table 1.** Swelling and structural parameters of MBAX gels.

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<sup>a</sup> Swelling ratio; <sup>b</sup> Molecular weight between two cross-links; <sup>c</sup> Cross-linking density;

<sup>d</sup> Mesh size.

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#### 286 *3.1.2 Swelling and structure of gels*

The swelling ratio (q) of gels is an important parameter that describes the amount of water 287 stored within the gel network. The swelling behavior of the MBAX gels at 4 and 6% (w/v) 288 was assessed in water for 15 h at 25 °C, with the equilibrium being reached between 8 and 289 6 h in MBAX-4 and MBAX-6, respectively. The q value (in g water/g of MBAX dry 290 weight) in the MBAX gels decreased from 18 to 9 when the MBAX concentration in the 291 gel increased from 4 to 6% (w/v) (Table 1). The q value found for the MBAX-4 gel was 292 similar to the value reported for maize bran AX gels (3.5% (w/v), q=20) (Berlanga-Reyes, 293 294 Carvajal-Millan, Caire Juvera, et al., 2009). The lower swelling ratio value registered at the highest MBAX concentration in the gel (MBAX-6) may have been related to the more 295 compact polymeric structure that limited the water absorption (Meyvis, De Smedt, 296 Demeester, & Hennink, 2000; Ross-Murphy & Shatwell, 1993). A similar effect of 297 polysaccharide concentrations on the swelling capacity of a gel has been reported for water-298 299 extractable AX (0.5 to 2 % (w/v), q values ranging from 231 to 69) or sugar beet pectin (2

to 5% (w/v), q values ranging from 120 to 35) after treatment with laccase (CarvajalMillan, Guilbert, et al., 2005; Micard & Thibault, 1999).

The cross-sectional microstructures of the MBAX-4 and MBAX-6 gels before swelling 302 were observed by scanning electron microscopy (SEM). Cross-sectional SEM images of the 303 MBAX gels are shown in Fig. 2. Both gels showed a three-dimensional and heterogeneous 304 305 network structure formed by the aggregation of nodular structures into clusters. MBAX-6 306 (Fig. 2b) presented a more compact microstructure in comparison with the geometries and irregular pores sizes of the MBAX-4 structure (Fig. 2a). These results are consistent with 307 the structural parameters calculated from the equilibrium swelling test. The molecular 308 309 weight between two cross-links (Mc), the cross-linking density ( $\rho_c$ ) and the mesh size ( $\xi$ ) values for MBAX-6 and MBAX-4 are presented in Table 1. When the MBAX 310 concentration increased in the gel from 4 to 6%, Mc and  $\xi$  decreased, leading to an increase 311 in the cross-linking density  $\rho_c$ . Similar Mc,  $\rho_c$  and  $\xi$  values have been reported in 3 to 5 % 312 AX gels (Berlanga-Reves, Carvajal-Millan, Caire Juvera, et al., 2009; Carvajal-Millan, 313 Guilbert, et al., 2005). The structural parameters are important in determining the physical 314 properties of the gels, including their mechanical strength, the diffusivity of the released 315 molecule, and their degradability (Hoffman, 2012). In hydrogels for drugs and protein 316 delivery or cell encapsulation, a typical  $\xi$  range of 5 to 300 nm has been reported for the 317 318 swollen state (Carvajal-Millan et al., 2006; Choh et al., 2011; Choudhary & Bhatia, 2012). The  $\xi$  values found for MBAX-4 and MBAX-6 are within this range. The results discussed 319 above indicate that by modifying the concentration of polysaccharides in the network, gels 320 with different rheological and structural properties can be obtained. 321



Fig. 2. SEM image showing the microstructures of cross-sectional MBAX-4 (a) and
MBAX-6 (b) networks at 350x magnification.

325 *3.2 Degradation of MBAX gels* 

326 The surface structures of degraded MBAX-4 and MBAX-6 hydrogels were studied by SEM and are presented in Fig. 3. At least two independent replicates of each sample were 327 analyzed for their surface morphologies, and similar results were obtained for the 328 replicates. Fig. 3a-b show the surface structures of MBAX-4 and MBA-6, respectively, 329 330 after 18 h of incubation under the same conditions as those used for fermentation but without the bacterial strains. The changes in the surface morphology of the MBAX-4 and 331 MBAX-6 hydrogels were observed as a function of their exposure time to Bifidobacterium 332 culture (a mixture of Bifidobacterium longum and Bifidobacterium adolescentis). SEM 333 images taken after 18 h and 36 h of incubation with the Bifidobacterium culture showed the 334 extent of the gel morphological changes (Fig. 3c-f). For both samples, the undegraded 335 MBAX gels presented relatively smooth and continuous surface structures without holes. 336 The degraded gels lost this structure and exhibited multiple cavities in their microstructure, 337 which increased in size and number with the time of incubation due to the enzymatic 338 activity of B. longum and B. adolescentis. However, at 18 h of incubation, differences in the 339 size of the cavities were observed between the MBAX gels (Fig. 3d vs. Fig. 3c). At 0 h of 340

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incubation, MBAX-6 has more cross-linked networks and a more compact structure 341 compared to MBAX-4 (Table 1 and Fig. 2), which may have reduced the accessibility of 342 the gel to enzymes and thus slowed down its degradation. Nonetheless, at 36 h of 343 344 incubation, the structure of both MBAX gels appeared to be collapsed rather than porous, as shown in Fig. 3e-f. Therefore, it is probable that after an initial surface erosion, the 345 enzymes diffused into the polymeric network and thus caused the degradation of the most 346 internal sites. Similar results were obtained for chemically crosslinked dextran gels that 347 348 were degraded by microbial enzymes (Brøndsted et al., 1998).



Fig 3. Scanning electron micrographs showing the surface morphologies of covalently
cross-linked MBAX-4 (a,c,e) and MBAX-6 (b,d,f) gels: (a-b) without degradation and
after; (c-d) 18 and (e-f) 36 hours of incubation with a mixture of *Bifidobacterium* strains (*B. longum*, *B. adolescentis*). All samples are at 350x magnification.

The degradation of MBAX gels with different structures was assessed by measuring the 354 residual AX molecular weights (Mw) and free monosaccharides in the fermentation media. 355 During the degradation experiments, MBAX chains with different Mw (Fig. 4) and a high 356 357 amount of free monosaccharides were released (data not shown). The changes in the residual AX Mw in ungelified MBAX and MBAX gels at different fermentation times were 358 registered by gel permeation chromatography (Fig. 4). The AX elution profile at time 0 359 represents the Mw distribution of ungelified MBAX and MBAX gels without bacterial 360 chain degradation. The Mw loss of MBAX gels was observed only in the presence of both 361 362 strains of Bifidobacterium. These results indicated that the presence of both bacteria is necessary to break down the gel structure. Previous research has suggested a possible cross-363 synergy between B. longum and B. adolescentis in the degradation of low molecular weight 364 arabinoxylans (Pastell et al., 2009; Wang, Sun, Cao, & Wang, 2010). The ungelified 365 MBAX showed over 80% of molecular weight loss after 18 h incubation and Mw 366 distributions of less than 50 kDa. This pattern was similar for different times of incubation 367 of ungelified MBAX (Fig 4a). Fig. 4c shows that MBAX-6 was slowly degraded and had a 368 Mw distribution of less than 50 kDa after 72 h of incubation, whereas MBAX-4 presented 369 this pattern after 36 h of incubation (Fig. 4b). However, in the first 18 h of incubation, a 370 fraction of both gels had Mw greater than 110 kDa. Hopkins et al. (2003) reported that the 371 372 breakdown of the xylan backbone in crosslinked AX was more resistant to degradation compared with the uncrosslinked AX due to the restricted access of xylanolytic enzymes. 373 Thus, the degradation of the gel in the early hours of incubation was restricted by the ability 374 of the enzymes to access their target sites. 375



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Fig. 4. HPSEC elution profiles of hydrolysis products of (a) ungelified MBAX, (b) MBAX-4 gel and (c) MBAX-6 gel degradation by a mixture of *Bifidobacterium* strains (*B. longum* and *B. adolescentis*) after incubation for 0 ( $t_0$ ), 18 ( $t_1$ ), 36 ( $t_2$ ) and 72 ( $t_3$ ) h. The elution profiles of pullulan molecular weight markers (kDa) were used for calibration and are shown at the top.

Changes in the degree of substitution were also observed during the fermentation, as shown 382 in Fig. 5a. The A/X ratio decreased with time in MBAX solution, MBAX-4, and MBAX-6 383 but increased after 36 h for the MBAX solution only. The differences observed after 36 h 384 385 between the MBAX solution and the MBAX gels could be associated with a more rapid initial removal of the arabinose side chains from the ungelified MBAX compared to the 386 MBAX gels. Once the branched regions of MBAX were removed, the xylan backbone was 387 hydrolyzed by bacterial xylanases, as reflected the increase in the A/X ratio. Hence, the 388 MBAX solution, MBAX-4, and MBAX-6 were degraded by a debranching mechanism 389 390 (Rose et al., 2010). The MBAX-4 and MBAX-6 gels showed different slopes for the linear decrease of the A/X ratio, in which the A/X ratio decreased much more rapidly in the 391 MBAX-4 gel than in the MBAX-6 gel. This difference was mainly related to the low 392 average mesh size (Table 1), compact microstructure (Fig. 2) and probably higher content 393 of intrachain di-FA in MBAX-6 compared to MBAX-4. Rosa et al. (2013) observed an 394 effect of the pore size on the in vitro fermentation of wheat aleurone with a high content of 395 insoluble AX. Moreover, Park et al. (2014) reported that the main factor causing the high 396 resistance to degradation of hyaluronic acid gels is the low swelling ratio resulting from a 397 high concentration of polysaccharides and a high degree of crosslinking. Hence, the slow 398 removal of the arabinose side chains from MBAX-6 may have been due to its more 399 400 compact structure, which slowed down enzymes penetration.



#### 401

402 Fig. 5. Changes in Arabinose/Xylose ratio during *in vitro* fermentation of ungelified
403 MBAX and MBAX-4 and MBAX-6 gels by a mixture of *Bifidobacterium* strains (*B. longum*, *B. adolescentis*).

405

The results above are consistent with the production of short-chain fatty acids registered 406 during the in vitro fermentation by the Bifidobacterium mix (Fig. 6). All three samples 407 produced significantly higher concentrations of total SCFA in comparison to the control 408 cultures at different times (MBAX-free substrate). The SCFA production increased with the 409 time that the MBAX-4 and MBAX-6 gels were fermented. In contrast, the SCFA 410 production in the ungelified MBAX remained constant from 36 to 72 h. Ungelified MBAX 411 produced a significantly (p < 0.05) higher amount of SCFA in comparison to the MBAX 412 gels at all fermentation times. At 18 h, both MBAX gels showed no significant differences 413 in the amount of SCFA produced. However, at 36 and 72 h, MBAX-6 presented a lower 414 amount of SCFA than MBAX-4. The total SCFA decreased significantly when the degree 415 416 of cross-linking increased in the MBAX gels. Acetic acid was the main acid produced,

followed by propionic and butyric acids. The ungelified MBAX and MBAX-4 produced 417 more acetic and propionic acids than MBAX-6, while butyric acid production was not 418 significantly different among the three fermented samples. The SCFA profiles resulting 419 420 from the fermentation of the ungelified MBAX and MBAX gels were similar to the one reported for alkali-soluble arabinoxylans from maize (Rose et al., 2010). These results 421 indicate that a certain portion of the AX chains released during the degradation of MBAX 422 gels was fermentable and that carbohydrates were efficiently converted to SCFA by the 423 424 Bifidobacterium mix.



425

426 Fig. 6. Total short-chain fatty acid (SCFA) production during fermentation of ungelified 427 MBAX, MBAX-4 gel and MBAX-6 gel by a mixture of *Bifidobacterium* strains (*B.* 428 *longum*, *B. adolescentis*) after incubation for 18, 36 and 72 h. Different letters (a to e) 429 indicate significant differences between samples (p < 0.05).

#### 431 **4.** Conclusions

Covalently cross-linked arabinoxylan gels with different structures were obtained by 432 increasing the concentration of AX undergoing oxidative gelation by laccase. The presence 433 of a more compact structure with a higher crosslinking density allowed for a slower 434 degradation of the three-dimensional gel structure by fermentation with a mixture of 435 Bifidobacterium. The ability to modulate the structural parameters of the AX gels to change 436 their biodegradability suggest that covalently cross-linked arabinoxylan gels could be used 437 for the design of controlled delivery systems activated by the colon microbiota. The in vitro 438 and in vivo simulation of arabinoxylan gels degradation in the colon will be further 439 investigated. 440

441

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