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Identification and structural analysis of cereal arabinoxylan-derived oligosaccharides by negative ionization HILIC-MS/MS

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ABSTRACT

Recent works provide evidence of the prebiotic potential of arabinoxylan-derived oligosaccharides (A)XOS. In this study, we developed a structural analysis for cereal-derived (A)XOS by negative ionization HILIC-MS/MS. Initially, we assessed twelve (A)XOS samples of known structures with different linkage positions and branching points by direct-infusion negative ESI-MSⁿ. We subsequently developed the negative ion HILIC-MS/MS with a post-column addition of ammonium chloride. The selected (A)XOS represented both linear (arabinofuranosyl residue linked to the non-reducing end of xylooligosaccharide) and branched structures. Each (A)XOS sample produced a specific spectrum in negative ion ESI-MSⁿ. By analyzing cross-ring fragment ions, we determined the linkage positions of linear (A)XOS. The presence or absence of diagnostic ions in the MS³ allowed us to detect different branches (*O*-2- or/and *O*-3-linked arabinofuranosyl with/or without *O*-4-linked xylopyranosyl at the non-reducing end). Furthermore, we could identify all analyzed samples by HILIC-MS/MS, based on the formed spectral library and chromatographic retention times.

1. Introduction

Arabinoxylan (AX) is a major polysaccharide found in the cell walls of cereal grain endosperm (Saulnier, Guillon, Sado, & Rouau, 2007). AX degrades to (arabino)xylooligosaccharides [(A)XOS] by enzymatic hydrolysis in the colon by means of intestinal microbiota or during food processing, such as brewing (Broekaert et al., 2011). Both AX and (A) XOS are considered as dietary fibers (Mendis & Simsek, 2014). (A)XOS include both xylooligosaccharides (XOS) and arabinoxylooligosaccharides (AXOS). Since recent studies have shown evidence of the prebiotic effects of AX and (A)XOS in the colons of humans and animals, interest in them has increased. AX and (A)XOS are associated with many health benefits, including immunomodulatory activity and attenuation of type II diabetes (Mendis & Simsek, 2014). (A)XOS have also been reported to attenuate serum cholesterol, triglyceride, and glucose levels (Broekaert et al., 2011).

Cereal grain AX consists of $\beta(1 \rightarrow 4)$ -linked xylopyranosyl residue (Xylp) backbone with arabinose mono- and di-substituents, as reviewed by Biliaderis and Izydorczyk (2007), Izydorczyk and Biliaderis (1995) and Saulnier et al. (2007). Arabinose substituents are $\alpha(1 \rightarrow 2)$ - or $\alpha(1 \rightarrow 3)$ -linked furanosyl residues (Araf) or both. Small amounts of β -D-Xylp-(1 $\rightarrow 2$)- α -L-Araf-(1 $\rightarrow 3$)-sidechains have also been found. Acidic AXs are present in the outer tissues of cereal grains. They carry $\alpha(1 \rightarrow 3)$ -

2)-linked glucopyranosyluronic acid or 4-*O*-methyl-glucopyranosyluronic acid substituents. Feruloyl residues linked to Araf residues are also found. The number of substituents is known to vary in different cereals, tissues, and even in tissue layers (Saulnier et al., 2007). Since structural features are related to physicochemical properties, such as water solubility, viscosity, and gelation (Saulnier, Sado, Branlard, Charmet, & Guillon, 2007), identification of the structure of AX is important if it is added to baking products or exploited in other industrial applications.

The structural analysis of large polysaccharide molecules is challenging; therefore, AXs are commonly hydrolyzed into (A)XOS prior to analysis. Nuclear magnetic resonance (NMR) spectroscopy (Duus, Gotfredsen, & Bock, 2000), high-performance anion-exchange chromatography with pulsed amperometric detection (HPAEC-PAD) (Pastell, Tuomainen, Virkki, & Tenkanen, 2008, Pastell, Virkki, Harju, Tuomainen, & Tenkanen, 2009), and gas chromatography-mass spectrometry (GC–MS) analysis of methylated (A)XOS (Ciucanu, 2006) have been popular techniques for the structural characterization of (A)XOS. The last two decades have seen an increased interest in using mass spectrometry (MS) and tandem mass spectrometry (MS/MS) for the structural analyses of carbohydrates (Mischnick, 2011). One specific benefit of MS and MS/MS analyses is that they require only small amounts of a sample to achieve a very sensitive analysis. Matrix-

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Fig. 1. The structural models and abbreviations of xylooligosaccharide and arabinoxylooligosaccharide samples (XOS and AXOS). Abbreviations of AXOS by Fauré et al. (2009).

assisted laser desorption ionization mass spectrometry (MALDI-MS) has been shown to be an efficient tool for determining the molecular weight distribution of (A)XOS. Reverse-phase high-performance liquid chromatography (RP-HPLC) combined with MS/MS has been used for the structural characterization of derivatized (A)XOS (Bowman, Dien, Vermillion, & Mertens, 2014). Recent and novel stationary phases, such as porous graphitized carbon (PGC) and hydrophilic interaction liquid chromatography (HILIC) phases, have enabled the HPLC-MS or HPLC-MS/MS analysis of oligosaccharides without derivatization (Everest-Dass, Kolarich, Campbell, & Packer, 2013, Everest-Dass et al., 2013, Hernandez-Hernandez, Calvillo, Lebron-Aguilar, Moreno, & Sanz, 2012, Leijdekkers, Sanders, Schols, & Gruppen, 2011, Leijdekkers et al., 2011, Liu, Kisonen, Willför, Xu, & Vilaplana, 2016, Liu et al., 2016, Pu, Zhao, Xiao, & Zhao, 2017).

Positive ion MS/MS can determine linkage positions and sequences of different oligosaccharides (Asam & Glish, 1997, Hofmeister, Zhou, & Leary, 1991, Mischnick, 2011). However, in MS/MS, positively charged oligosaccharide ions fragment from both ends, and, if the oligosaccharides consist of the same mass monomers, such as (A)XOS, identification of isobaric fragment ions is difficult and sometimes even impossible. Therefore, derivatization techniques, such as derivatization by per-O-methylation (Ciucanu, 2006, Matamoros Fernández, Obel, Scheller, & Roepstorff, 2003) and labeling the reducing end, for example, with ¹⁸O isotope (Hofmeister et al., 1991) or reagents (Suzuki, 2013), have commonly been used to differentiate the reducing end and non-reducing end fragment ions.

However, some studies have indicated that derivatization may not be needed if negative ionization is used. Negatively charged oligosaccharides have been reported to fragment mainly from the reducing end towards the non-reducing end and produce non-reducing end consisting ions (A-, B-, and C-ions) (Carroll, Willard, & Lebrilla, 1995, Chai, Piskarev, & Lawson, 2001, Harvey, 2005, Maina et al., 2013, Pfenninger, Karas, Finke, & Stahl, 2002a). Fragmentation in one direction can facilitate the analysis of cross-ring fragments from middle residues (Maina et al., 2013). By analyzing the cross-ring fragments, the linkages have been determined from negatively charged deprotonated (Harvey, 2005, Pfenninger et al., 2002a) or anion adduct (Guan & Cole, 2008, Zhu & Cole, 2001) disaccharides and oligosaccharides.

Although negative ionization MS/MS methods have been widely developed for the structural analysis of oligosaccharides, to our knowledge, only two MS/MS studies of AXOS using negative ionization have been published. Quemener, Ordaz-Ortiz, and Saulnier (2006) characterized the structure of neutral deprotonated AXOS by electrospray ionization quadrupole time-of-flight mass spectrometry (ESI-Q-TOFMS) and electrospray ionization ion trap mass spectrometry (ESI-ITMS). They showed that mono- or di-substituted AXOS (O-3substituted vs. O-3- and O-2-substituted) can be differentiated by the negative ion MS/MS method. Wang et al. (2009) developed the negative ion HPLC-MSⁿ method for structure elucidation of feruloylated AXOS. In both studies, the AX was hydrolyzed by glycoside hydrolase family 11 (GH11) endoxylanase. GH11 endoxylanases cleave the xylan backbone but not near the branching residues. Therefore the AXOS have unsubstituted +1 Xylp residues and +2 branched residues (Pollet, Delcour, & Courtin, 2010). Other common endoxylanase enzymes. which are also present endogenously in plants, hydrolyzing AX, are GH10 family enzymes. GH10 endoxylanases cleave the AX backbone near the branching residue (+1 being branched residue), and they form AXOS without the O-4 linked backbone Xvlp residue at the non-reducing end. We think that the development of an MS/MS method for structural analysis for the GH10 hydrolyzed AXOS is clearly needed. As AXOS are usually hydrolysate mixtures of AX, combining the negative MS/MS method with HILIC would enable high-throughput analysis of AXOS mixtures without sample derivatization.

In this study, we present a novel negative ionization HILIC-MS/MS method for the analysis of the linkage positions and branching points of AXOS. The present work achieves the following: 1) to elucidate the fragmentation behavior of chlorinated AXOS in negative electrospray ionization multiple-stage mass spectrometry (ESI-MSⁿ), 2) to determine how linkage positions influence the fragmentation of linear and branched AXOS by negative ion HILIC-MS/MS. Since our samples included AXOS from both GH10 and GH11 family hydrolysates, they comprise all possible linkage positions and branches that AXOS can form, including also the rare β -D-Xylp-(1 \rightarrow 2)- α -L-Araf-(1 \rightarrow 3)- β -D-Xylp-(1 \rightarrow 4)-Xylp and α -L-Araf-(1 \rightarrow 2)-mono-substituted samples.

2. Materials and methods

2.1. Samples and chemicals

Five linear XOS (X1, X2, X3, X4, and X5,) and three branched AXOS samples (XA³XX, XA²XX + XA³XX, and XA²⁺³XX, named GH11 AXOS) were purchased from Megazyme International (Bray, Ireland). Six AXOS samples (A²X, A³X, A²⁺³X, A²⁺³XX, A²XX, and D^{2,3}X, named GH10 AXOS) were prepared earlier by our group, as published elsewhere (Pastell et al., 2008, Pastell et al., 2009, Rantanen et al., 2007). The structures of the AXOS and XOS samples are shown in Fig. 1. Abbreviations are by Fauré et al. (2009). The $A^{3}X$, $A^{2+3}X$, $A^{2+3}XX$, $A^{2}X$, and A²XX were prepared from wheat arabinoxylan (Megazyme International, Bray, Ireland) and the D^{2,3}X from oat spelt arabinoxylan (Sigma Aldrich Chemie, Steinheim, Germany) by using different hydrolyzing enzyme treatments. In short, the treatments included enzymatic hydrolysis by GH family 10 endo-1,4-β-D-xylanase (Shearzyme; Novozymes, Bagsvaerd, Denmark), GH family 51 α -L-arabinofuranosidase (Megazyme International, Bray, Ireland), and GH family 43 α -L-arabinofuranosidase (Megazyme International, Bray, Ireland). Hydrolysates were fractionated by gel permeation chromatography (GPC) and identified by HPAEC-PAD and NMR spectroscopy. All chemicals were of analytical grade unless otherwise stated.

2.2. Direct infusion ESI-MSⁿ

The positive and negative ion MSⁿ spectra of AXOS samples were produced by the Agilent XCT Plus model ion trap mass spectrometer (Agilent Technologies, Waldbronn, Germany) equipped with an electrospray ionization (ESI) source. The parameters were as reported previously (Maina et al., 2013), except the electrospray capillary voltage was set to (\pm) 3200 V. The samples were diluted in methanol/water/formic acid (50/49/1, $\nu/\nu/\nu$) to a final concentration of 8–12 µg/mL. The samples were directly infused into the ion source at a flow rate of 5 µL/min. Ammonium chloride (NH₄Cl, 40 µg/ml) was added in negative ionization mode for [M + Cl]⁻ adduct ion formation.

2.3. HILIC-MS/MS method development

2.3.1. Optimization of ion trap parameters

XOS standards (X_1-X_5) were analyzed using a Bruker Esquire quadrupole ion trap (Bruker, Bremen, Germany) equipped with an ESI source and using direct infusion (5 µL/min) and negative ionization. Samples of 10 µg/mL were prepared in 80% acetonitrile (ACN) with 0.1% NH₄OH. NH₄Cl (50 µg/mL) was added for chloride adduct formation. Ion trap parameters with the highest intensity level were optimized for each standard. The optimized fragmentation amplitudes were 0.65 V for X₂, 0.60 V for X₃, 0.70 V for X₄ and 0.55 V for X₅. For XA²⁺³XX the 0.5 V fragmentation amplitude was used.

2.3.2. HPLC setup

Optimized ion trap parameters were tested by HILIC-MS with a mixture of four XOS (X2-X5) and six AXOS samples. The chromatographic separation was performed with HPLC (Agilent 1100 Series HPLC, Agilent Technologies, Santa Clara, CA) on a 3.5 µm, $2.1 \times 150 \text{ mm}$ Xbridge BEH amide column (Waters, Milford, MA). The mobile phases were A) 80% ACN with 0.1% ammonium hydroxide (NH₄OH) and B) 20% ACN with 0.1% NH₄OH. The mobile phase elution profile was as follows: 80% ACN to 50% ACN in 31 min, isocratic (50% ACN) for 5 min, back to initial conditions in 1 min, and 15 min washing in initial conditions (based on Brokl, Hernández-Hernández, Soria and Sanz (2011)). Flow rate was 200 µL/min and oven temperature was set to 35 °C. Injection volume was 10 µL. The segment program was created for HILIC-MS. The segments were the following: 0-7 min with X₂ parameters, 7-12 min with X₃ parameters, 12-17 min with X₄ parameters, 17-23 min with X₅ parameters, and 23-31 min with X₅ parameters.

2.3.3. Optimization of post-column NH₄Cl addition parameters

The chloride adduct ions were formed by adding NH₄Cl (in 80% ACN with 0.1% NH₄OH) post-column to the mobile phase flow via a mixing tee with an HPLC pump (Waters 501 HPLC system; Millipore, Milford, MA). Post-column addition of NH₄Cl was tested with flow rates of 100 µL/min and 200 µL/min and NH₄Cl concentrations of 1 µg/mL, 2.5 µg/mL, 5 µg/mL, 7.5 µg/mL, and 10 µg/mL. Testing was performed with 5 µg/mL and 10 µg/mL XOS mixture (X₂–X₅) solutions. The aim was to determine the lowest concentration of NH₄Cl that produces the maximal [M + Cl]⁻ adduct ion signal.

2.3.4. HILIC-MS/MS of AXOS samples

The HILIC-MS/MS methods were set up for tri-, tetra-, penta- and hexasaccharides by optimized ion trap parameters and fragmentation amplitudes. The chloride adduct ions were formed by adding NH₄Cl post-column in an optimized concentration and flow rate ($10 \mu g/mL$ in 80% ACN with 0.1% NH₄OH, 200 μ L/min). The MS and MS² spectra were produced from all samples.

3. Results and discussion

3.1. Linkage determination of linear XOS and AXOS by direct infusion ESI- MS^n

The first part of the study was to find out how the XOS and AXOS linkage positions reflect on the product ion spectra. The negative ion MS^2 and MS^3 spectra of the linear XOS and AXOS are shown in Fig. 2. Product ions are named according to Domon and Costello (1988).

3.1.1. Determining the $(1 \rightarrow 4)$ -linkage

All the commercial XOS samples consisted of $\beta(1 \rightarrow 4)$ -linked Xylp residues. The MS² spectrum of X₄ shows that the reducing end Xylp residue formed ^{0,2}A and ^{0,2}A-H₂O cross-ring fragment ions at *m/z* 485 and *m/z* 467 (Fig. 2a). The (1 \rightarrow 4)-linked middle residues fragmented to ^{0,2}A₃-ions (*m/z* 353) and ^{0,2}A₃-H₂O-ions (*m/z* 335) as well. All the



Fig. 2. Negative ion MSⁿ spectra of linear (A)XOS samples and characterization of fragmentation. Precursor ions are $[M + Cl]^-$ and product ions are deprotonated. / = double cleavage. (a) X₄, MS² m/z 581, (b) X₄, MS³ m/z 581 \rightarrow 413, (c) X₄, MS³ m/z 449 \rightarrow 281, (d) A³X, MS² m/z 449, (e) A³X, MS³ m/z 449 \rightarrow 281, (f) A²XX, MS³ m/z 581 \rightarrow 413, (h) A²XX, MS³ m/z 581 \rightarrow 281, (i) D^{2,3}X, MS² m/z 581, (j) D^{2,3}X, MS³ m/z 581 \rightarrow 413, and (k) D^{2,3}X, MS³ m/z 581 \rightarrow 281. Product ions named according to Domon and Costello (1988).

linear GH10 AXOS samples were also found to produce ^{0,2}A and ^{0,2}A-H₂O cross-ring fragment ions from their $(1 \rightarrow 4)$ -linked reducing ends in MS² (A²XX: *m*/*z* 485 and *m*/*z* 467 in Fig. 2f; A³X: *m*/*z* 353 and *m*/*z* 335 in Fig. 2d, and $D^{2,3}X$: m/z 485 and m/z 467 in Fig. 2i). This was an expected result, since many studies of negatively charged oligosaccharides have reported the $(1 \rightarrow 4)$ -linked oligosaccharides fragmenting by ^{0,2}A and ^{0,2}A-H₂O cross-ring cleavages (Pfenninger, Karas, Finke, & Stahl, 2002b, Zhu & Cole, 2001), and this has also been observed in deprotonated (A)XOS (Quemener et al., 2006, Wang et al., 2009). The C₂-ions and C₃-ions were isolated and further fragmented in MS^3 . The C₃-ions and C₂-ions of commercial X₄, which contained (1 \rightarrow 4)-linkage at the reducing end, were again found to produce 0,2 A-ions (m/z 353 in Fig. 2b and m/z 221 in Fig. 2c), but the ^{0,2}A-H₂O cross-ring ions were not formed. The $(1 \rightarrow 4)$ -linked C₃-ions of A²XX fragmented to both 0,2 A- and 0,2 A-H₂O-ions in MS³ (m/z 335 and m/z 353 in Fig. 2g).

3.1.2. Determining the $(1 \rightarrow 2)$ -linkage

The C₃-ions and C₂-ions of A²XX produced cross-ring fragment ions with m/z 191 (C₂-90 Da) and m/z 233 (C₂-48 Da) in MS³ (Fig. 2g-h). We suggest that the m/z 191 is a ^{0,2}X₂-ion, since the $(1 \rightarrow 2)$ -linked hexose oligosaccharides have been found to fragment by ^{0,2}X-type ions due to the loss of 120 Da neutrals (Maina et al., 2013). The m/z 233 could be a ^{1,5}A₂-ion, formed by the loss of the reducing end of the C₂-ion. As expected, the same cross-ring ion peaks were seen in the spectra of D^{2,3}X, which also contained $(1 \rightarrow 2)$ -linked middle residue (m/z 191: ^{0,2}X and m/z 233: ^{1,4}A in Fig. 2j-k). It was also noted that the MS³ spectrum of the C₂-ion of D^{2,3}X (Fig. 2k) was nearly identical to the MS³ spectrum of the C₂-ion of A²XX (Fig. 2h). Therefore, we suggest that the M-90 Da (^{0,2}X) and M-48 Da (^{1,4}A if Araf and ^{1,5}A if Xylp) cross-ring cleavages are diagnostic for $(1 \rightarrow 2)$ -linkages in pentoses. Unfortunately, the method is not capable of differentiating β -p-Xylp- $(1 \rightarrow 2)$ - α -L-Araf from α -L-Araf- $(1 \rightarrow 2)$ - β -p-Xylp.

3.1.3. Determining the $(1 \rightarrow 3)$ -linkage

The A³X (Fig. 2d) and D^{2,3}X (Fig. 2i), which had $(1 \rightarrow 3)$ -linkages between their middle residues, did not form ^{0,2}A, or any other kind of cross-ring fragments from O-3-position linked residues. The finding that $(1 \rightarrow 3)$ -linked oligosaccharides do not fragment via cross-ring cleavages in negative mode is consistent with the results of deprotonated AXOS (Quemener et al., 2006). In MS³, the C₃-ion of D^{2,3}X, which had a $(1 \rightarrow 3)$ -linked reducing end residue, did not produce cross-ring fragments from the reducing end residue (Fig. 2j). However, in the C2-ion spectrum of $(1 \rightarrow 3)$ -linked A³X, a small number of the cross-ring ions were observed with m/z 221 (Fig. 2e). The spectrum was very similar to the C₂-ion spectrum of the $(1 \rightarrow 4)$ -linked X₄ (Fig. 2c), but did not have a loss of water peak (m/z 263). These cross-ring ions are most likely 0,3 X₁-ions, since we have experienced that deprotonated (1 \rightarrow 3)-linked hexose disaccharides form $^{0,3}X_1$ -ions by the loss of 90 Da units (Juvonen, 2011). On the other hand, the chlorinated $(1 \rightarrow 3)$ -linked hexose disaccharides were observed to lack the cross-ring fragmentation (Maina et al., 2013). These differences in fragmentation of $(1 \rightarrow 3)$ linked deprotonated and chlorinated ions should be kept in mind if interpreting spectra of unknown oligosaccharides.

3.2. Structural analysis of branched AXOS by direct infusion ESI-MSⁿ

The analysis continued by examining how branching points of AXOS influence on fragmentation. The negative ion MS^2 spectra of branched AXOS are shown in Fig. 3. As XA^2XX was available only as part of the mixture, the MS^2 spectrum of XA^2XX is from HILIC-MS/MS.

3.2.1. Determining the $(1 \rightarrow 4)$ -linkages from linear parts of branched AXOS

In MS^2 , as observed previously with linear (A)XOS, the branched GH11 AXOS samples also formed ^{0,2}A and ^{0,2}A-H₂O type cross-ring

fragment ions from their $(1 \rightarrow 4)$ -linked linear reducing end residue (m/z 749 and m/z 731 in Fig. 3a; m/z 617 and m/z 599 in Fig. 3b-c). Furthermore, ^{0,2}A and ^{0,2}A-H₂O cross-ring fragment ions were formed from $(1 \rightarrow 4)$ -linked middle residues (m/z 617 and m/z 599 in Fig. 3a; m/z 485 and m/z 467 in Fig. 3b-c). Also, as expected, the branched GH10 AXOS, A²⁺³XX, and A²⁺³X fragmented from their $(1 \rightarrow 4)$ -linked xylan backbone by ^{0,2}A and ^{0,2}A-H₂O type cross-ring cleavages (m/z 617, m/z 599, m/z 485, and m/z 467 in Fig. 3d-e).

3.2.2. Branching point determination of AXOS by MS^2

The hypothesis was that, if the branched AXOS followed the same fragmentation behavior as linear AXOS and started fragmenting from the reducing end, the branching points could be determined by a lack of cross-ring ions in the spectrum. However, it was found that the lack of ions was observed only in the spectra of the Araf mono-substituted GH11 AXOS samples (XA²XX in Fig. 3b; XA³XX in Fig. 3c) between m/z281 and m/z 395. The di-substituted XA²⁺³XX showed a peak at m/z467 (Fig. 3a), even though the spectrum should have been lacking ions between m/z 395 and m/z 527, if it has been fragmented only from the reducing end. The ions at m/z 467 were characterized as ${}^{0,2}A_4/Y_{3x}$ -ions formed by co-fragmentation of the reducing and the non-reducing ends. Interestingly, the branched GH10 AXOS, A²⁺³XX, and A²⁺³X were also found to co-fragment from both ends, similarly to the Araf di-substituted XA²⁺³XX; therefore, no lack of ions indicating branching residue were shown in their spectra either (m/z 323 and m/z 335 in)Fig. 3d-e). The peak was characterized at m/z 323 as a ${}^{2,4}A_3/Y_{3x}$ -ion and at m/z 335 as a 0,2 A-H₂O/Y_{3x}-ion. Di-substituted samples fragmenting from both ends implies that di-substitution enhances the Ytype fragmentation of Araf substituents.

3.2.3. Characterization of branched AXOS by MS^2

The Araf mono-substituted pentasaccharides XA³XX and XA²XX were easily differentiated from each other by the peak at m/z 299 in the spectrum of XA³XX (Fig. 3c). It was reported earlier that *O*-3-branched AXOS produce these unique types of ions by double C- and Y-type cleavages combined with a water molecule addition in the negative mode (Quemener et al., 2006). Therefore, the m/z 299 was characterized as C₂/Y_{3x} + H₂O-ions. Since these fragment ions were absent also in the MS² spectrum of pentasaccharide A²⁺³XX (Fig. 3d), the peak at m/z 299 is an excellent indicator of XA³XX.

The Araf di-substituted sample $XA^{2+3}XX$ (Fig. 3a) was found to produce more abundant m/z 587 (C-90 Da) compared to the monosubstituted samples XA^2XX and XA^3XX (Fig. 3b-c). Since the cross-ring fragmentation most likely happens from the reducing end toward the non-reducing end by retro-aldol reaction, it is likely ^{2,4}A-ion. The same ^{2,4}A-type ions were found from spectra of branched GH10 AXOS samples: m/z 587 and m/z 455 in $A^{2+3}XX$ (Fig. 3d) and m/z 455 and m/z323 in $A^{2+3}X$ (Fig. 3e). It should be noted that the linear samples Xylp backbone did not fragment by these cross-ring cleavages.

The branched, Araf di-substituted $A^{2+3}XX$ (Fig. 3d) and $A^{2+3}X$ (Fig. 3e) both produced peaks at m/z 263 in MS². Since the m/z 263 was not found in the MS² spectra of the branched Araf mono-substituted pentasaccharides XA²XX or XA³XX, the $A^{2+3}XX$ can be distinguished by it. The peak is probably formed by p-type glycosidic bond fragment ions (C₂/Z_{3x}-ions). The p-ions have been reported to fragment from some *O*-3-branched oligosaccharides next to the *O*-3-branching unit by double C- and Z-cleavages in negative mode (Chai et al., 2001, Chai, Lawson, & Piskarev, 2002, Everest-Dass et al., 2013, Harvey, 2005). The assumed p-ions were also found in the MS² spectra of di-substituted XA²⁺³XX (m/z 395, Fig. 3a). Since the p-ions are also isobaric to B-ions formed from linear AXOS, they cannot, however, act as an indicator of di-substituted or branched structures.

3.2.4. Characterization of branched AXOS by MS^3

The C_{2} - and C_{3} -ions of branched samples were isolated and fragmented in MS^{3} . The precursor ions might have consisted of some minor



Fig. 3. Negative ion MS² spectra of the branched AXOS and characterization of fragmentation. Precursors are $[M + Cl]^-$ adduct ions. Product ions are deprotonated. / = double cleavage, $x = \alpha$, β , or γ ; branching unit = the *m/z* area of branching unit if fragmentation from reducing end. (a) XA²⁺³XX, *m/z* 845, (b) XA²XX (HILIC-MS²), *m/z* 713, (c) XA³XX, *m/z* 713, (d) A²⁺³XX, *m/z* 713, and (e) A²⁺³X, *m/z* 581. Product ions named according to Domon and Costello (1988).

proportion of Y-ions, since non-reducing end fragmentations were detected in MS^2 . Each MS^3 spectrum of branched AXOS was observed to be different from the others (Fig. 4). This means that all these branched tri- or tetrasaccharide ions could be identified by the MS^3 spectrum. For example, the C₂-ion of $XA^{2+3}XX$ (Fig. 4b) showed peaks at m/z 431 and m/z 395. The spectra of the C₂- and C₃-ions of XA^3XX had abundant peaks at m/z 299 (Fig. 4d-e) which were absent in other spectra. The C₃-ion spectrum of $A^{2+3}XX$ showed ^{2,4}A-ions at m/z 455 but was lacking

the B₂-ions at m/z 395. The C₂-ion spectrum had a strong peak at m/z 263, but showed no peak at m/z 299. The C₃-ion spectrum of XA²XX (Fig. 4c) was lacking both the m/z 299 and m/z 431, but had an abundant B₂-ion peak at m/z 395.

The ions with m/z 299, which were earlier identified as C₂/ Y_{3x} + H₂O in the MS² of XA³XX, were found also in the MS³ spectra of XA³XX (Fig. 4d-e). A surprising finding was the peak at m/z 431, which was found in the C₂-ion spectrum of XA²⁺³XX (Fig. 4b) and in the C₃-



Fig. 4. Negative ion MS^3 spectra of the branched AXOS' C₃- and C₂-ions and characterization of fragmentation. Precursors are $[M + Cl]^-$ adduct ions, and product ions are deprotonated. (a) $XA^{2+3}XX$, m/z 845 \rightarrow 677, (b) $XA^{2+3}XX$, m/z 845 \rightarrow 545, (c) XA^2XX , m/z 713 \rightarrow 545, (d) XA^3XX , m/z 713 \rightarrow 545, (e) XA^3XX , m/z 713 \rightarrow 413, (f) $A^{2+3}XX$, m/z 713 \rightarrow 545, and (g) $A^{2+3}XX$, m/z 713 \rightarrow 413. Product ions named according to Domon and Costello (1988).

ion spectrum of XA³XX (Fig. 4d). These fragment ions are one pentose unit larger than the ions at m/z 299, and, therefore, they are also assumed to be $C/Y_{3x} + H_2O$ ions. Both of these ions (m/z 299 and m/z 431) were only found in the spectra of O-4- and O-3-branched samples. We suggest that a water molecule has been added to the C-ion after the Y-type cleavage of (1 \rightarrow 3)-linked Araf or (1 \rightarrow 4)-linked Xylp residues from the non-reducing end, as then the ring has the potential to open with the formation of an aldehyde, which is known to react with water to form a hydrate.

The fragment ions, which we earlier suggested to be p-ions (C_2/Z_{3x} ions), were observed in all MS³ spectra of the O-3-branched samples. We assume them to be p-ions, not Z-ions, because Quemener et al. (2006) reported the same m/z at the spectrum of deprotonated XA³X, when the reducing end was differentiated from the non-reducing end by ¹⁸O-labelling. The p-ions were found in the MS³ of XA²⁺³XX (Fig. 4a-b, m/z 395), XA³XX (Fig. 4d-e, m/z 263), and A²⁺³XX (Fig. 4f-g, m/z 263). Unfortunately, the m/z 263 was also found from the spectrum of O-2branched XA²XX (Fig. 4c), and, therefore, it cannot be a diagnostic peak of the O-3-linked branch. As previously commented, the p-ions are also isobaric to B-ions formed from linear AXOS; thus, they cannot act as an indicator of the branched structure either.

Interestingly, the MS^3 spectra were observed to also contain other kinds of double cleavage fragment ions. These ions were found as minor peaks at m/z 245 and were formed only from *O*-2-branched samples $(XA^{2+3}XX \text{ in Fig. 4a-b}, XA^2XX \text{ in Fig. 4c}, and A2^{+3}XX \text{ in Fig. 4g})$. These could be E- or F-type elimination ions as previously reported (Maslen, Goubet, Adam, Dupree, & Stephens, 2007, Spina et al., 2004) and formed by double B- and Z-cleavages (E-type) or two Z-cleavages (Ftype). Since linear AXOS did not form these ions in MS³, the ions at m/z245 are the potential diagnostic peaks of the branched structure. However, more research is needed to determine if the more complicated branched samples also fragment similarly.

3.3. Analysis of (A)XOS by positive $ESI-MS^n$

To show the difference between positive and negative ionization, the positive MS² and MS³ spectra of three lithium cationized tetrasaccharides (X₄, $D^{2,3}X$, and $A^{2,3}X$) are presented in Online Resource 1 (OR. 1). The branched $A^{2,3}X$ was observed to produce primarily the same peaks in MS^2 as the linear equivalents X_4 and $D^{2,3}X$. The only exception was a peak at m/z 253 in the A^{2,3}X spectrum (OR. 1e). Otherwise, only differences in the ion intensity occurred. All three samples were found to produce $^{0,2}A$ cross-ring fragments (*m*/*z* 493, neutral loss of 60 Da) from their $(1 \rightarrow 4)$ -linked reducing end residues. A small peak of the ^{2,4}A-ion (m/z 463, neutral loss of 90 Da) was also observed. The same ^{0,2}A- and ^{2,4}A-type of cross-ring fragment ions were formed in MS³ (*m*/*z* 361, *m*/*z* 331, and *m*/*z* 229) from the Y- and C-ions (OR. 1b, 1d, and 1f). Since the three MS³ spectra were very similar, the isolated ions for MS³ had to be mainly Y-ions formed by losing a nonreducing end. It was clear that the positive ion ESI-MSⁿ was not as effective as the negative ion ESI-MSⁿ to characterize the structural differences of (A)XOS samples which were located near the non-reducing end.

3.4. HILIC-MS/MS

3.4.1. Setting up HILIC-MS in negative mode

Based on the promising results in negative ionization, implementation of the chromatographic method with an HILIC column was initiated by using negative ionization. In the first phase, the ion trap parameters were optimized for every XOS standard (X_1-X_5) by using direct infusion. Then, the optimized parameters were tested by HILIC-MS with XOS and AXOS samples. The chromatograms of the XOS mixture (X_2-X_5) analyzed with different parameters can be found in OR. 2. It was observed that larger oligosaccharides formed smaller peaks when optimized parameters for X_2 and X_3 were used, and smaller



Fig. 5. Optimization of post-column addition of NH₄Cl to HILIC-MS with X₂-X₅ samples. Segment program: (a) Base peak chromatogram of X₂-X₅ mixture, without NH₄Cl addition, (b) Base peak chromatogram of X₂-X₅ mixture, 10 μ g/l NH₄Cl added with flow rate 200 μ L/min, (c-f) HILIC-MS spectrum of X₂, X₃, X₄ and X₅, respectively, without NH₄Cl addition, (g-j) HILIC-MS mass spectrum of X₂, X₃, X₄, and X₅, respectively, 10 μ g/L NH₄Cl added with flow rate of 200 μ L/min.

oligosaccharides formed smaller peaks on X_3 - X_5 optimized parameters. As a result, the segment program was applied for MS. The segment program changes the ion trap parameters during the run, to obtain the highest intensity levels (Fig. 5a).

The MS spectra of samples showed that various adduct ions were formed in the negative ionization mode (Fig. 5c-f). The intensity levels of base peaks were approximately 2×10^5 – 3×10^5 . NH₄Cl was added to form exclusively chloride adducts and to enhance the intensity level. The post-column addition of NH₄Cl was chosen to enable the use of the same mobile phase in the positive mode. Post-column addition was tested with two flow rates and five concentrations of NH₄Cl. The smallest concentration to form acceptable peaks was found to be $10 \,\mu\text{g/mL}$ with a flow rate of 200 $\mu\text{L/min}$ (Fig. 5b). The [M + Cl]⁻ adduct ions were observed to be the dominant peaks in the MS spectra of XOS (*m*/*z* 317, *m*/*z* 449, *m*/*z* 581, and *m*/*z* 713 in Fig. 5g-j). Furthermore, the addition of NH₄Cl increased the intensity level of base peaks to 4×10^5 – 6×10^6 .

3.4.2. HILIC-MS/MS of (A)XOS

In the last stage of the study, the (A)XOS samples were analyzed by



HILIC-MS and HILIC-MS/MS under optimized conditions. An HILIC-MS/MS method was created for each size of oligosaccharide. The $\rm MS^2$ spectra of HILIC-MS/MS (Fig. 6) and direct infusion MS/MS were

Fig. 6. Extracted ion chromatograms of (A)XOS analyzed by negative ion HILIC-MS and MS² spectra of HILIC-MS/MS. Precursors are [M + Cl] – adduct ions and product ions are deprotonated. (a) X₃, MS² m/z 449, (b) X₄, MS² m/z 581, (c) X₅, MS² m/z 713, (d) A²X, MS² m/z 449, (e) A³X, MS² m/z 449, (f) D²⁺³X, MS² m/z 581, (g) A²⁺³X, MS² m/z 581, (h) A²XX, MS² m/z 581, (i) A²⁺³XX, MS² m/z 713, (j) XA³XX, MS² m/z 713, (k) XA²XX, MS² m/z 713, and (l) XA²⁺³XX, MS² m/z 845. Product ions named according to Domon and Costello (1988).

compared to each other. Some differences in fragmentation pattern and ion intensity were found. This might be due to the different pH of the eluent in HILIC-MS/MS and direct infusion. HILIC-MS/MS was run at pH 10 (to prevent anomeric peaks from splitting) and directly infused ESI-MSⁿ was run at pH 5.

All samples were found to form a specific MS² spectrum, although the spectra of XA²XX (Fig. 6k) and X₅ (Fig. 6c) had very similar ion distributions. It was noticed that trisaccharides and tetrasaccharides can be identified by their base peaks (table of diagnostic peaks in OR 3). X_3 had m/z 263 (Fig. 6a), A^2X had m/z 335 (Fig. 6d)), and A^3X had m/z281 (Fig. 6e). X₄ had m/z 395 (Fig. 6b), D^{2,3}X had m/z 413 (Fig. 6f), $A^{2+3}X$ had m/z 467 (Fig. 6g), and A^2XX had m/z 395 (Fig. 6h) as their base peaks in the MS² spectrum. Even though A²XX had the same base peak as X_4 , it was easily differentiated from X_4 by the peak at m/z 335. Pentasaccharides had only base peaks at m/z 599 or m/z 617. Therefore, the presence or absence of other peaks was needed to define them. X₅ had *m/z* 281, *m/z* 467, *m/z* 599, and *m/z* 617 peaks (Fig. 6c). $A^{2+3}XX$ had m/z 263, m/z 467, and m/z 599 peaks (Fig. 6i). XA³XX had *m*/*z* 263, *m*/*z* 299, *m*/*z* 599, and *m*/*z* 617 peaks (Fig. 6j). XA²XX had *m*/ z 467, m/z 599, and m/z 617 peaks (Fig. 6k). The following retention times were observed in HILIC in the current study: A³X 10.3 min, A²X 12.1 min, A²⁺³X 13.5 min, X₃ 15.3 min, D^{2,3}X 15.9 min, A²XX 17.6 min, A²⁺³XX 18.5 min, X₄ 20.7 min, XA³XX 21.3 min, XA²XX 22.4 min, XA²⁺³XX 24.3 min, and X₅ 25.6 min. It was noted that the O-3-Arafsubstituted samples eluted before the O-2-Araf-substituted samples and the di-substituted samples were last in the order. Samples with two Xylp residue backbones eluted in the order of X_2 , A^3X , A^2X , and $A^{2+3}X$, and samples with three Xylp residue backbones in the order of X_{3} , $A^{2}XX$, and $A^{2+3}XX$. The elution order of four Xylp residue backbone samples was X₄, XA³XX, XA²XX, and XA²⁺³XX. In addition, the lengthening of the Xylp backbone was found to increase the retention, and Araf substitution decreased the retention, since the A³X, A²X, and A²⁺³X samples eluted faster than the linear X₃. Furthermore, the retention times were all different, which will confirm their identification in future studies.

4. Conclusions

The feasibility of negative ionization direct infusion ESI-MSⁿ and HILIC-MS/MS to analyze and identify the structures of cereal-derived (A)XOS was studied. Initially, the influence of linkage positions on fragmentation of (A)XOS was examined. It was found that negative ionization ESI-MSⁿ can characterize each linkage of linear (A)XOS. Each linkage produced diagnostic cross-ring fragments, the presence or absence of which can determine the linkage positions in (A)XOS. The $(1 \rightarrow 4)$ -linked Xylp backbone produces $^{0,2}A$ - and $^{0,2}A$ -H₂O-ions. The $(1 \rightarrow 2)$ -linked residues produce ^{0,2}X₂-ions from both Xylp and Araf residues. The $(1 \rightarrow 3)$ -linked residues lack the cross-ring fragment ions, except for the $(1 \rightarrow 3)$ -linked C₂-ion of A³X. Also, the branched samples were found to fragment next to the branched residue by double glycosidic bond cleavages. The branched samples were determined by the presence or absence of diagnostic ions in the MS³ spectra. It was also found that diagnostic ions (m/z 245 or m/z 299) in MS³ can distinguish the studied branched samples from linear samples. Ion m/z 299 is also a diagnostic ion of O-3 and O-4 branched residues. Subsequently, HILIC-MS/MS was implemented for the identification of (A)XOS. The results showed that the negative ionization HILIC-MS/MS method was efficient for identification of (A)XOS samples by using a now formed spectral library and chromatographic retention times. The implemented method will be most useful for the structural analysis and profiling of AX and mixtures of (A)XOS from different plant materials. If the AXOS sample has many branching points, the method can be improved with ¹⁸O-labeling of the reducing end or separation with ion mobility mass spectrometry.

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Conflict of interest

The authors declare that they have no conflicts of interest regarding this work.

Appendix A. Supplementary data

Supplementary data to this article can be found online at https://doi.org/10.1016/j.foodchem.2018.09.074.

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