Integrated processing of sugarcane bagasse: Arabinoxylan extraction integrated with ethanol production.

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Abstract

A proof-of-concept project compared extraction of arabinoxylans (AX) from sugarcane bagasse and wheat bran via alkaline hydrogen peroxide followed by enzyme-assisted extraction with combinations of feruloyl esterases and a xylanase. Bagasse contains comparable amounts of AX to wheat bran, but with a much lower arabinoxylan substitution on the xylan backbone (A:X ratio of around 0.2 compared with 0.6 for wheat bran), hence offering AX products with distinctive functionality and potential end uses. In the current work, bagasse released its AX more readily than wheat bran, and released a wider range of molecular weights. Use of feruloyl esterase and xylanase enzymes on their own or following alkaline peroxide extraction did not enhance AX release substantially; however, the xylanase appeared to be effective at reducing the size of AX molecules, and there is scope to optimise the effects of enzymes to produce specific AX product fractions. As bagasse frequently arises within the context of bioethanol production, integration of AX extraction with ethanol production could allow economic production of a portfolio of AX products, as has been demonstrated in principle for AX co-production in a wheat ethanol plant.

Keywords: sugarcane bagasse, wheat bran, arabinoxylans, bioethanol, biorefinery integration.

Abbreviations

AX Arabinoxylan
UoH University of Huddersfield
UoL University of Lincoln
UoY University of York
UoStA University of St Andrews
BDC Biorenewables Development Centre
LBNet Lignocellulosic Biorefinery Network
P2PNet Plants to Products Network
CE-High High cut-off fraction (from ultrafiltration over 10 kDa) following chemical extraction
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1. Introduction

For a reaction to take place, entities must come together and interact under the right conditions. The interactions leading to the current work arose from a workshop organized in September 2014 by the Lignocellulosic Biorefinery Network (LBNet, https://lb-net.net), a Biotechnology and Biological Sciences Research Council Network in Industrial Biotechnology and Bioenergy (BBSRC NIBB). At this workshop, one of those who eventually formed a consortium asked of another, “If I want to use a source of lignin for a chemical conversion process I have developed, what lignin would make a good source?”, to which the other advised “One that already arises naturally within existing biorefineries, such that transportation costs are negated and the infrastructure and integration opportunities are in place to enable the lignin processing to be undertaken economically, and in which the raw material might already be partially degraded through earlier processing, to give more ready access to the lignin.” At a later point in the workshop, the participants were asked to write on pieces of paper what we would like to research, and place them on the floor. One of us wrote “Arabinoxylans”, based on a long-standing interest in this subject as a promising coproduct of ethanol biorefineries, as the ethanol is used to precipitate the arabinoxylans, making the production of AX potentially economically viable in that context [1]. Meanwhile a representative of the company AB Sugar wrote “Something valuable from sugarcane bagasse (not furfural, we already do that)”. Seeing these notes together prompted the idea that arabinoxylans might similarly be extracted from bagasse in an integrated biorefinery producing ethanol from sugarcane, and that the residual bagasse following AX extraction might reveal a suitable source of lignin for further processing. A proof-of-concept project was constructed, to demonstrate the feasibility of AX extraction from sugarcane bagasse, and to examine the residual lignin, following partial deconstruction during the AX extraction process, for its suitability as a feedstock for conversion into a phenolic monomer using an established reaction sequence [2]. A parallel activity studying wheat bran was included, to extend the scope and make the findings more immediately applicable to the UK context (where bioethanol production is largely from wheat). Enzyme enhancement of the AX release, a scale-up component and a techno-economic analysis of AX production were also included in the project, to give a consortium comprising the Universities of Huddersfield, St Andrews, York, Lincoln and Nottingham along with the Biorenewables Development Centre in York (http://www.biorenewables.org/). Later a student project supported by the Plants to Products Network (another BBSRC NIBB, http://www.nibbp2p.org) extended the project by undertaking a bioethanol pinch analysis, following the approaches of Martinez et al. [3], to minimise ethanol usage while producing a range of AX products including arabinoxylan-oligosaccharides (AXOS) [4].

This paper describes the proof-of-concept work undertaken to demonstrate AX extraction from sugarcane bagasse and to compare it with extraction from wheat bran in terms of yield, composition

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and responsiveness of the extraction process to enzyme enhancement. A future paper will describe the investigation of the residual bagasse following AX extraction, in terms of the nature of its lignin and its suitability for further processing, including further integration opportunities with ethanol and butanol.

Global sugarcane production is around 1.9 billion tonnes per annum (https://www.statista.com/statistics/249604/sugar-cane-production-worldwide/), resulting in around 570 million tonnes wet bagasse or half this amount if dried. Sugarcane bagasse thus represents a major waste stream arising from sugar and alcohol industries, typically containing around 40-50% cellulose, 25-35% hemicelluloses (predominantly xylans) and 20-30% lignin [5,6]. A focus of previous work has been to deploy feruloyl esterases and xylanases to assist the saccharification of bagasse to increase recovery of fermentable sugars, by removing the hemicelluloses that (to put it simply) link lignin and cellulose, thus increasing accessibility of the latter to cellulases and hence the release of glucose [7]. A difference in the current work is that the intention was not to hydrolyse arabinoxylan hemicelluloses to their constituent sugars, but rather to release and recover them as intact large AX molecules, in which form they have potential as functional food ingredients and non-food products including film forming, emulsifiers and stabilisers in the food, pharmaceutical and cosmetic industries [8,9]. The context of the ethanol biorefinery gives scope for AX co-production to be economic as a result of integration with ethanol production (used for precipitating the AX), as has been shown previously for AX production from wheat bran in a wheat ethanol biorefinery [1]. Wheat bran typically contains 20-30% AX [10,11], similar to bagasse, but with a much higher ratio of arabinose to xylose units; in wheat bran the A:X ratio is typically in the range 0.5-0.6 [12,13]), while in sugarcane bagasse it is much lower at around 0.2 typically [14]. This “cleaner” xylan backbone with fewer arabinose substitutions is likely to exhibit different functional properties compared with wheat bran AX, including reduced solubility and greater susceptibility to enzyme action, as well as effects on viscosity and gel formation, and performance in food products or animal feed.

2. Materials and Methods

Sugarcane bagasse (25 kg) and wheat bran (50 kg) were sourced commercially by AB Sugar and provided to the project. The bagasse and bran were milled at the Biorenewables Development Centre (BDC) using a Retsch Cutting Mill SM 300 (Retsch GmbH, Germany) with a 2.00 mm screen, and AX extraction studies undertaken at the University of Huddersfield (UoH). Proximate analysis and arabinoxylan (AX) content measurements were performed at the University of York (UoY). Lignin studies, to be presented in a future paper, were undertaken at the University of St Andrews (UoSfA).

Figure 1 describes the chemical extraction process using alkaline hydrogen peroxide applied to the milled wheat bran and sugarcane bagasse, based on the work of Hollmann and Lindhauer [15] and Du et al. [16]. Dilute alkaline peroxide reacts with lignin, breaking the association with polysaccharides and facilitating the release of arabinoxylans [10]. The residue material was then subjected to enzyme treatment to see if further release of AX resulted, using selected combinations of β-Xylanase (C. mixtus, PRO-E0051, referred to here as XYL) and two types of Feruloyl esterase (A. cellulolyticus, PRO-E0355 and PRO-E0356, referred to here as FE-E0355 and FE-E0356) from Prozomix UK. The hypothesis was that the feruloyl esterases, by breaking links between AX and
lignin, might enhance AX release, while the xylanase, which breaks the AX chains themselves, might similarly release AX fragments to enhance overall extraction, while also altering the size of the released molecules. Enzyme-assisted extraction on its own was also investigated, along with chemical extraction followed by further extraction with buffer solution, and buffer extraction on its own.

In total eight extractions were performed for each feedstock:
1. Direct buffer extraction (control for all trials, particularly Trials 7-8);
2. Chemical extraction followed by buffer extraction (control for Trials 3-6);
3. Chemical extraction followed by FE-E0355 and XYL;
4. Chemical extraction followed by FE-E0356 and XYL;
5. Chemical extraction followed by FE-E0355;
6. Chemical extraction followed by FE-E0356;
7. Direct enzyme extraction with FE-E0355 and XYL;
8. Direct enzyme extraction with FE-E0356 and XYL.

The supernatant following treatment, centrifugation and filtering was separated by ultrafiltration over a 10 kDa membrane (see below for details). The chemical extraction thus yielded High cut-off (CE-High) and Low cut-off (CE-Low) fractions and a pellet of residual material; enzyme extractions directly on the raw material or on the pellet following chemical extraction similarly yielded High cut-off (EE-High) and Low cut-off (EE-Low) fractions and a pellet. A single chemical extraction was performed to produce residual material for subsequent enzyme extraction in Trials 3-6; a second chemical extraction was performed for subsequent buffer extraction (Trial 2).

For the chemical extraction, 50 g bran or 11 g bagasse (because of the lower bulk density of the latter restricting the amount that could be processed in a bottle) was weighed into a 1 L Duran bottle. Foaming is a problem during AX extraction, so 15 drops of anti-foaming agent (Dimeticon SILFAR® SE 4, Wacker Chemie AG, Germany) were added, then 400 mL of 2% hydrogen peroxide (Fisher Scientific UK Limited, analytical grade) was added very slowly, with stirring with a magnetic stirrer. 5 more drops of anti-foaming agent were added, followed by the final 100 mL of hydrogen peroxide solution. The pH was adjusted to 11.5 with 50% NaOH. The bottle was placed in a 50°C water bath and the solution stirred for 4 h. The pH was controlled every hour and readjusted to 11.5 if needed. After cooling to room temperature the pH of the mixture was adjusted to 7 with concentrated sulphuric acid. The solution was centrifuged for 15 minutes at 4000 rpm using a Beckman GS-6S centrifuge (Beckman Coulter Life Sciences, USA). The supernatant was filtered and the remaining solids washed with 150 mL of water, then centrifuged, filtered, washed with 150 mL water and centrifuged again. The final supernatant was filtered and pooled with the previous two. The residual solid (the pellet) was placed in the oven to dry overnight at 50°C.

The high molecular weight material in the supernatant was concentrated by ultrafiltration using a Vivaflow™ 200 system (Sartorius Stedim Biotech GmbH, Germany) with polyethersulfone membranes with a molecular weight cut-off of 10 kDa. (Ultrafiltration would be done as part of a commercial process to reduce the amount of ethanol needed subsequently to precipitate the AX [1,15]; in the current work the fractions were not precipitated. The ultrafiltration served to separate the released AX into larger MW and smaller MW fractions.) Retentate was recycled until the volume was reduced to one fifth of the original. The retentate (High cut-off) and permeate (Low cut-off) fractions were freeze-dried using a Christ Freeze Dryer Alpha 1-4 LDplus, (Martin Christ
Gefriertrocknungsanlagen GmbH, Germany) at −47.8°C and 0.35 mbar. Samples of freeze-dried High
cut-off and Low cut-off material and oven-dried pellets were sent to UoY for AX analysis (see below).
Samples of the dried pellet material were also sent to the University of St Andrews (UoStA) for
assessment of its lignin, to be described in a future paper.

For enzyme-assisted extraction trials on raw wheat bran or sugarcane bagasse or on pellets after
chemical extraction, material (30 g for the wheat bran and 6 g for the sugarcane bagasse) was
weighed into a 1 L Duran bottle with a magnetic stirrer. 300 mL of a buffer (composed of 107.4 mL
of 0.2 M disodium phosphate, 42.6 mL of citric acid and 150 mL of water) at pH 6.5 was added, the
bottles placed in a 37°C water bath and the pH of the solution readjusted to 6.5 with citric acid when
the temperature reached 37°C. The different enzymes were added at a level of 10 µg/10 g for the
XYL and FE-E0355 and 5 µg/10 g for the FE-E0356, and the solution was stirred for 2 h. As above,
samples were centrifuged, filtered and washed, in this case using 100 mL of water for each washing,
and the residual solid pellet oven-dried overnight at 50°C. Again the supernatant was passed
through ultrafiltration over 10 kDa and the retentate (High cut-off, EE-H) and permeate (Low cut-off,
EE-L) freeze dried and sent to UoY for analysis along with the oven-dried pellet, with pellet samples
also sent to UoStA for lignin analysis.

Due to the limited scope of this small proof-of-concept project, replicate enzyme extractions were
not performed, the aim being to demonstrate broad effects in relation to AX yields from wheat bran
and bagasse and the potential effects of enzyme-assisted extraction. Five replicate chemical
extractions were performed, as enzyme extraction Trials 3-6 and buffer Trial 2 each required a
chemical extraction first, although High and Low cut-off fractions were subsequently produced for
only two of these (2 and 3), and only these two pellets were sent for lignin analysis. Subsequent
similar work with replication has confirmed the broad trends reported here and shown sufficient
reproducibility to have confidence in the trends, which are reported and discussed here within the
limits of the acknowledged lack of replication.

Proximate analysis of the wheat bran and sugar cane bagasse was undertaken as follows:

**Lignin determination: acetyl bromide method**

Biomass powder was weighed out (4 mg) into 2 mL tubes. The biomass was heated at 50°C for 3
hours after adding 250 µL of acetyl bromide solution (25% acetyl bromide and 75% glacial acetic acid
by volume) and vortexing every 15 minutes. After the samples were cooled to room temperature,
the contents were transferred into 5 mL volumetric flasks. A further 1 mL of NaOH (2 mol L⁻¹) was
used to rinse the tubes pouring the NaOH into the 5 mL flasks. 175 µL of hydroxylamine HCl (0.5 mol
L⁻¹) was added to the volumetric flasks and, after vortexing, the latter were filled up to 5 mL with
glacial acetic acid and mixed several times. Finally, in order to measure the 280 nm UV adsorption
by spectrophotometer, 100 µL of each sample was diluted in 900 µL of glacial acetic acid. The
amount of lignin was calculated using the following formula: [absorbance/(coefficient pathlength)] ·
[(total volume · 100%)/biomass weight], where coefficient = 15.69, pathlength = 1, total volume = 5,
biomass weight = 4.

**Non-cellulosic monosaccharide determination**

Following the method of Fry [17], biomass dry powder (4 mg) was partially hydrolysed by adding 0.5
mL of trifluoroacetic acid (TFA, 2 mol L⁻¹). Then, the vials were flushed with dry argon, mixed and
heated at 100°C for 4 hours, mixing periodically. The vials were then cooled to room temperature
and dried in centrifugal evaporator with fume extraction overnight. The pellets were washed twice with 500 µL of 2-propanol and vacuum dried. Finally, the samples were resuspended in 200 µL of deionised water, filtered with 0.45 µm PTFE filters, and analysed by HPAEC (see below).

It became evident that these hydrolysis conditions were inadequate to release all the AX in the original wheat bran (see below, where the mass balance indicates more AX in the extracted fractions than appeared to be present initially in the bran). The wheat bran was therefore hydrolysed under a range of conditions, to investigate the effects on AX measurement and to draw conclusions about the most appropriate conditions for AX analysis in wheat bran. The bran samples were hydrolysed in 4M TFA (2 mL per 4 mg bran) for 1, 2 and 4 hours, and in 2M (2 mL per 4 mg bran) for 4 and 6 hours at 120°C. In addition, samples were pretreated in 98% TFA for 1, 2, 4 and 6 hours at room temperature, then diluted to 4M and boiled for 1 hour at 120°C.

Crystalline cellulose
Biomass dry pellets after TFA hydrolysis were washed once with 1.5 mL of water, and twice using 1.5 mL of acetone. The dried pellets were left to air dry overnight before complete hydrolysis by adding 90 µL of 72% w/w sulphuric acid, incubating at room temperature for 4 hours. 1.89 mL of water was subsequently added and the sample was heated for 4 hours at 120°C. The glucose content of the supernatant was assessed using the colorimetric Anthrone assay, using a glucose standard curve.

Analysis of the sugar compositions of the wheat bran and bagasse samples and of fractions and residues following the various chemical and enzyme extractions was undertaken as follows:

Monosaccharide analysis was performed by high performance anion-exchange chromatography (HPAEC) (Dionex IC 2500) on a Dionex Carbopac PA-10 column with integrated amperometry detection [18]. The separated monosaccharides were quantified using external calibration with an equimolar mixture of nine monosaccharide standards (arabinose, fucose, galactose, galacturonic acid, glucose, glucuronic acid, mannose, rhamnose, and xylose), which were subjected to TFA hydrolysis in parallel with the samples.

3. Results and Discussion
Table 1 shows the compositions of the wheat bran and sugarcane bagasse used in the current work. The wheat bran had a significant starch content; therefore for the scale-up work undertaken at BDC, the wheat bran was washed with water to remove starch prior to extraction, in line with the recommendation of Du et al. [16]. The bagasse had 28% lignin and nearly 20% AX, broadly in line with literature [5,6,14], and suggesting it was a promising candidate for recovery of both materials. The analytical procedure used indicated 10% lignin in the wheat bran, although subsequent NMR work suggested a much lower lignin content, in line with other recent reports that wheat bran contains less lignin than previously thought. These results also suggest an AX content of only 8.64% in the wheat bran; this figure is lower than the 20-30% generally expected for wheat bran, and later proved to be incompatible with the mass balance for AX recovered in the various fractions and residues, which suggested an AX content in the original material of around 24% (see below). The A:X ratio for the wheat bran was 0.57, and for the bagasse 0.21, in line with typical values expected from the literature, and showing the much “cleaner” xylan backbone for the bagasse AX, with fewer arabinose substitutions compared with the wheat AX.
Tables 2 and 3 report the crude yields, AX concentrations and hence AX yields from 50 g wheat bran and 11 g sugarcane bagasse, respectively, in the starting materials and in the fractions following the various chemical, enzyme and buffer treatments. The second column in each table is for chemical extraction only, showing the data from two replicates, with good agreement. Considering the wheat bran results first, chemical extraction of around 50.6 g of bran (moisture content 9.5%) yielded a residual dry pellet weighing 29.94 g (averaged from Trials 2 and 3), a High cut-off (CE-High) of 8.21 g, and a Low cut-off (CE-Low) of 4.40 g (total 42.55 g, c.f. about 45 g solids in the original material).

The AX concentration in the pellet was 35.81% and in the High and Low cut-off fractions was 16.32% and 1.25%, respectively. This implies a total amount of AX in the pellet and two fractions of 29.94×35.81% + 8.21×16.32% + 4.40×1.25% = 10.72 + 1.34 + 0.06 = 12.12 g. The mass balance therefore implies an AX concentration of 12.12/50.6 = 24% in the original wheat bran, higher than the 8.64% reported in Table 1, and more in line with the expected AX content of wheat bran, suggesting that the 8.64% figure is erroneous.

It is well known that the appropriate hydrolysis conditions for this sort of analysis depend on the nature of the sample, with different samples requiring different combinations of time, temperature and acidity to get an optimum balance between release and degradation of monosaccharides [19,20]. The bran sample was therefore reanalysed under a range of hydrolysis conditions as described above. Figure 2 shows the AX contents calculated for each of the hydrolysis regimes. Under similar conditions to the original analysis (2M for 4 hours, but at a higher temperature, 120°C), the measurement was similar at 8.79%, while 2M for 6 hours released more AX to give 14.04%. 4M for 1 or 2 hours released even more (18.58-19.71%), but 4M for 4 hours appears to give substantial degradation and a final measurement of only 5.50%. Pretreatment in concentrated acid appeared effective at releasing more AX for effective hydrolysis, giving measurements of around 26.7% after 1 or 2 hours of pretreatment, decreasing to 22.8 and 20.6% after 4 and 6 hours, suggesting degradation at the high acid concentration despite the low temperature. Overall, these results demonstrate that an AX content of around 24%, as implied by the mass balance, is plausible, but that measuring the AX content of raw wheat bran requires different hydrolysis conditions compared to extracts or the residual pellet following extraction.

Thus, of a total of around 12 g AX in the original 50 sample, 1.4 g or 11.5% was released by the alkaline hydrogen peroxide extraction process. This is much lower than the recoveries of 46-50% reported by Hollmann and Lindhauer [15] and Du et al. [16] for alkali-extracted AX from wheat bran using similar conditions. In those studies the wheat bran was boiled in 70% ethanol at 80°C for 4 hours prior to alkaline \( H_2O_2 \) treatment, which Hollmann and Lindhauer [15] advised was necessary to achieve high yields; this step was omitted in the current work (for safety and cost reasons in relation to the planned scale-up work, and because related unpublished work from our labs on extraction from maize meal had found that this step was not needed, as also confirmed by work from Doner and Hicks [21] on AX extraction from maize fibre). The omission of this ethanol boiling step is possibly the reason for the lower yields than in this previously reported work.

Considering the sugarcane bagasse results, chemical extraction of around 11.5 g of bagasse (moisture content 9.3%) yielded a residual dry pellet weighing 5.43 g (averaged from Trials 2 and 3), a High cut-off (CE-High) of 3.11 g, and a Low cut-off (CE-Low) of 8.70 g. The total appears to be 17.34 g, c.f. about 10.5 g solids in the original material; the mass balance does not give good agreement in this case. This is probably because the Low cut-off was extrapolated from the solids
left after freeze-drying dilute samples (and the freeze-dried samples may not have been completely dry), such that the 8.70 g figure is not accurate, while overall the mass balance from just 11 g of bagasse is inherently less accurate than that from 50 g wheat bran, and the contribution from salts formed on neutralisation relatively greater.

The AX concentration in the pellet was 14.02% (averaged from Trials 2 and 3, although higher in Trials 4-6) and in the High and Low cut-off fractions was 23.64% and 7.23%, respectively. This implies a total amount of AX in the pellet and two fractions of 5.53×14.02% + 3.11×23.64% + 8.70×7.23% = 0.775 + 0.735 + 0.629 = 2.139 g, compared with 11.54×19.58% = 2.260 g of AX in the original sample. This mass balance appears reasonable, although the uncertainty over the Low cut-off contribution is acknowledged. In this case it appears that the AX left in the pellet (assuming this to be a more accurate figure) is about one-third of the AX in the original material (0.775/2.25 = 34%), implying nearly two-thirds has been released, compared with only 11.5% for the wheat bran. This suggests sugarcane bagasse may be very amenable to AX extraction, as it appears to yield its AX more readily than wheat bran (although the higher water:solids ratio used for the bagasse extraction is noted). The readiness of the bagasse to release its AX compared with wheat bran suggests the expensive and hazardous ethanol boiling step advised by Hollmann and Lindhauer [15] for wheat bran may not be needed in a bagasse-based AX extraction process. As noted above, Doner and Hicks [21] advised that dewaxing with toluene-ethanol was unnecessary for AX extraction from maize fibre.

Figure 3 shows the crude yields of High and Low cut-off material following the various wheat bran extractions, the AX concentrations in the fractions and hence the absolute yields of AX in each fraction. Trial 1 shows the recovery of High and Low fractions from just extraction with Buffer. Clearly, while quite a lot of small molecular weight material (<10 kDa) was recovered in the Low cut-off fraction, it contained very little AX, while some high MW AX was extracted just with the use of Buffer. Chemical extraction is shown in Trials 2 and 3, with good agreement, showing roughly twice as much High cut-off material as Low was recovered, but that the latter contained very little AX, such that the majority of the recovered AX was in the High cut-off fraction, giving a yield of around 2.6% compared with only 0.62% with Buffer. Thus chemical extraction using alkaline hydrogen peroxide was somewhat successful at releasing high MW AX from wheat bran. Further extraction with Buffer (Trial 2) recovered a little more high MW AX.

Turning to Trials 7 and 8, use of FEA enzymes in combination with XYL appeared to enhance release of AX a little compared with just Buffer, but not substantially. Trials 3-7 taken together indicate that enzyme treatment following chemical extraction was able to release a little more AX, with the balance changing towards low MW material when the xylanase was included. This makes sense, although the absence of any detectable AX in the fractions from Trial 6 is unexpected. However, in general the enzymes did not dramatically enhance the further extraction of AX, and it is not possible to infer any meaningful differences in the performance of the different enzyme combinations.

Figure 4 shows the equivalent results for sugarcane bagasse. Clearly the patterns are overall quite contrasting to those for the wheat bran. Most obviously, from Figure 4(c), the recovery of AX by chemical extraction was much greater than for wheat bran for both High and Low cut-off material, with yields of around 6-7% for high MW AX and 5.5% for low MW AX, compared with 2.6% and 0.1%, respectively, for wheat bran. Thus the bagasse released its AX much more readily following chemical extraction, and released a more balanced profile of AX between large and small MW molecules; this
is also apparent in Figure 4(a) which shows much greater crude yields of Low cut-off material than High, in contrast to the yields from wheat bran.

A consequence of the ready release of AX under chemical extraction is that there is therefore less material for the enzymes subsequently to work on, hence the subsequent enzyme treatments yield very little extra AX, although again there is evidence that the presence of the xylanase shifts the balance towards smaller molecules, as expected; this is clearer in Figure 5 which presents an expanded view of the Absolute yields, to allow the effects of the enzyme treatments, such as they are, to be seen more clearly. Even on their own, however, without prior chemical extraction, the enzymes release little more than Buffer alone (Trials 7-8 c.f. Trial 1), with the extra being entirely small MW material. It is recognised, however, that this small study did not explore a wider range of enzyme dosages and incubation conditions; it is likely that the effects of the enzymes could be enhanced under optimised conditions.

Figure 6 shows the A:X ratios in the various wheat bran and bagasse extracts. In line with the starting material and as expected from literature reports, the wheat bran extracts have much higher A:X ratios than those from bagasse. Given the limitations of the work, not much more can be read into the fine detail of Figure 6, beyond noting that for the bagasse extracts, the smaller molecular weight material (Low cut-off fractions) consistently had lower A:X ratios than the corresponding larger MW fractions, whereas the wheat-derived AX presents a more mixed picture. It is well established that in general different parts of the biomass structure contain AX with different molecular weights and A:X ratios, reflective of different botanical functions of AX in different parts of the plant [22]. For the bagasse extracts, it appears to be consistently the case that material initially released by whichever means (chemical extraction, enzymes or buffer) has higher A:X ratios than material released subsequently via further extraction with buffer or enzymes, again reflecting differences in the nature of AX material given up easily compared with that released on further processing.

The picture is less consistent for the wheat bran extracts, reflecting that the initial release was less extensive from the wheat bran than from the bagasse, such that comparisons are less dominated by that initial release; for the bagasse, so much was released initially that the remaining AX material is understandably quite different, whereas for the wheat bran, so little was released at all that what was released at any point was similar in structure. The A:X ratios greater than 1 for some of the Low cut-off wheat bran fractions (Trials 1 and 2) are probably erroneous, arising from errors in measuring very low concentrations of A and X in these samples (see Figure 2(b)), although some components of wheat outer layers (cross-cells and pericarp) can have A:X ratios great than 1 [22].

As noted already, the cleaner xylan chains of bagasse AX would offer somewhat different properties compared to those of wheat AX. This is an important consideration in developing commercial products; the challenge is to understand the functional performance and potential uses of AX fractions as affected by molecular weight and A:X ratio, and hence to understand which feedstocks and extraction processes are suitable for producing specific fractions. Even then, the likely scenario is not that specific fractions would be targeted for exclusive production, but rather that processing would co-produce a range of AX fractions, each suitable for different end-use applications, including small AXOS fractions with prebiotic functionality in food and in animal feed, alongside mid-range and large molecular weight fractions offering gradations of product functionality in relation to viscosity, gel formation and interaction with other food components [4]. In this respect, commercialisation of
AX-based products is likely to follow the fractionation paradigm of crude oil cracking, to produce a range of products and to find markets for each. The use of enzymes would form part of the approach for creating specific fractions with targeted end-uses.

The above observations and comments regarding the results from the current work are made in full recognition of the limits of replication and accuracy of the study; nevertheless, the overall patterns are clear, relative to the objectives of the work and the wider commercial context, and lead confidently to the following conclusions:

1. Arabinoxylans can be extracted from sugarcane bagasse via similar protocols previously used for wheat bran, yielding AX with a lower A:X ratio than for wheat bran, hence offering different functional properties for end-use applications.

2. Bagasse released its AX more readily than wheat bran, and released a wider range of AX molecules, with a greater proportion of small MW (<10 kDa) molecules. In the current work, around two-thirds of the AX in the bagasse was released by chemical extraction, more or less equally divided between High and Low cut-off material, compared with just 11% of mostly large MW AX initially in the wheat bran.

3. Boiling wheat bran in ethanol prior to extraction appears to be necessary to obtain good yields of AX.

4. Within the conditions used, feruloyl esterase and xylanase enzymes had small effects on releasing AX from raw wheat bran or sugarcane bagasse or from residues following chemical extraction, with the xylanase tending to shift the balance from large to small MW molecules.

Following these results, larger scale extractions of AX from bran and bagasse were performed at the Biorenewables Development Centre (BDC) using alkaline hydrogen peroxide extraction. Bran was washed with water prior to extraction to remove the starch. 25 kg of bran and bagasse, in 5 kg batches, were subject to chemical extraction, centrifugation, ultrafiltration, ethanol precipitation, recovery and drying. A total of 1250 g of wheat bran extract (5% yield) of 54% purity and 848 g of bagasse extract (17% yield) at 52% purity were produced, with much of the rest being analysed as glucose, either from residual starch or from cellulose.

**4. Conclusions**

The hypotheses that arabinoxylans could be extracted from sugarcane bagasse using similar protocols used for wheat bran, and that enzyme treatment might enhance the extraction, were investigated in a small proof-of-concept project. Bagasse was shown to be a promising source of AX in terms of its content (around 20%) and structure (with a low A:X ratio) and the readiness with which it yielded its AX to give a balanced release of both large (>10 kDa) and small (<10 kDa) molecules. Use of feruloyl esterase and xylanase enzymes on their own or following alkaline peroxide extraction was not particularly effective at enhancing AX release; however, there was evidence that the xylanase was effective at reducing the size of AX molecules, and there is scope to optimise the action of the enzymes through a more comprehensive study of dosage and incubation effects.
Thus, the metaphorical reactions that arose from the LBNet workshop were successful in demonstrating the proof of concept, but the literal reactions in relation to optimising AX extraction from wheat bran and sugarcane bagasse retain some scope for further enhancement.

As bagasse frequently arises within the context of bioethanol production, integration of AX extraction with ethanol production could allow economic production of AX products, as has been demonstrated in principle for AX co-production in a wheat ethanol plant [1]. Further processing of the now lignin-rich residue could give even further opportunities within the biorefinery, both for co-production of additional products and for further integration (particularly if the lignin processing also involves ethanol). The nature and additional processing of the lignin in the residual fractions from the current work were therefore studied further, and will be the topic of a future paper.

Acknowledgements

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References


Table 1. Compositions of wheat bran and sugarcane bagasse (%w/w dry basis).

<table>
<thead>
<tr>
<th>Component</th>
<th>Wheat bran</th>
<th>Sugarcane bagasse</th>
</tr>
</thead>
<tbody>
<tr>
<td>Starch</td>
<td>29.8</td>
<td>ND</td>
</tr>
<tr>
<td>Hemicellulose</td>
<td>16.6</td>
<td>22.1</td>
</tr>
<tr>
<td>Cellulose</td>
<td>21.8</td>
<td>28.3</td>
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<tr>
<td>Lignin</td>
<td>10.4</td>
<td>22.0</td>
</tr>
<tr>
<td>Other</td>
<td>21.2</td>
<td>27.6</td>
</tr>
<tr>
<td>AX (=0.88×(A+X))</td>
<td>8.64</td>
<td>19.58</td>
</tr>
<tr>
<td>A:X ratio</td>
<td>0.57</td>
<td>0.21</td>
</tr>
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</table>
Table 2. Crude yields, AX concentrations and AX yields from wheat bran and its fractions following extraction under different treatments.

| Treatment                        | Initial Bran (g) | AX Concentration (%) | AX Amount (g) | CE Pellet (g) | AX Concentration (%) | AX Amount in CE Pellet (g) | CE High (g) | AX Concentration (%) | AX Amount in CE High (g) | Absolute Yield (%) | CE Low (g) | AX Concentration (%) | AX Amount in CE Low (g) | Absolute Yield (%) | EE Pellet (g) | AX Concentration (%) | AX Amount in EE Pellet (g) | Absolute Yield (%) | EE High (g) | AX Concentration (%) | AX Amount in EE High (g) | Absolute Yield (%) | EE Low (g) | AX Concentration (%) | AX Amount in EE Low (g) | Absolute Yield (%) |
|----------------------------------|------------------|----------------------|---------------|---------------|----------------------|---------------------------|---------------|----------------------|---------------------------|-------------------|---------------|----------------------|---------------------------|-------------------|------------------|----------------------|---------------------------|-------------------|------------------|----------------------|---------------------------|-------------------|------------------|----------------------|---------------------------|
| 1. Buffer (Control)              | 30.34            | 8.64                 | 2.62          | 30.29         | 34.05                | 10.31                     | 8.11          | 16.43                | 1.33                      | 2.63              | 3.84          | 1.75                 | 0.067                     | 0.13              | 14.63          | 31.00                | 4.54                      | 14.95             | 1.81          | 10.42                | 0.19                      | 0.62              | 4.78            | 1.15                | 0.05                      | 0.18              |
| 2. Chemical + Buffer (Control)   | 50.73            | 8.64                 | 4.38          | 29.58         | 38.50                | 11.39                     | 8.31          | 16.21                | 1.35                      | 2.68              | 4.97          | 0.74                 | 0.037                     | 0.07              | 16.02          | 35.82                | 5.47                      | 12.06             | 2.00          | 1.21                 | 0.24                      | 0.48              | 0.97            | 0.29                | 0.00                      | 0.18              |
| 3. Chemical + Enzymes FEA E0355 + XYL E0051 | 50.20 | 8.64 | 4.34 | 29.21 | 33.67 | 9.84 | 8.11 | 16.21 | 1.35 | 2.68 | 4.97 | 0.74 | 0.037 | 0.07 | 15.04 | 35.74 | 5.04 | 3.41 | 5.73 | 5.00 |
| 4. Chemical + Enzymes FEA E0356 + XYL E0051 | 50.47 | 8.64 | 4.36 | 29.54 | 36.92 | 10.92 | 8.31 | 16.21 | 1.35 | 2.68 | 4.97 | 0.74 | 0.037 | 0.07 | 14.10 | 22.61 | 5.04 | 3.41 | 5.73 | 5.00 |
| 5. Chemical + Enzyme FEA E0355 | 50.47 | 8.64 | 4.36 | 29.54 | 36.92 | 10.92 | 8.31 | 16.21 | 1.35 | 2.68 | 4.97 | 0.74 | 0.037 | 0.07 | 15.06 | 36.07 | 5.04 | 3.41 | 5.73 | 5.00 |
| 6. Chemical + Enzyme FEA E0356 | 50.47 | 8.64 | 4.36 | 29.54 | 36.92 | 10.92 | 8.31 | 16.21 | 1.35 | 2.68 | 4.97 | 0.74 | 0.037 | 0.07 | 14.10 | 22.61 | 5.04 | 3.41 | 5.73 | 5.00 |
| 7. Enzymes FEA E0355 + XYL E0051 | 30.61 | 8.64 | 2.64 | 30.29 | 35.82 | 10.92 | 8.11 | 16.21 | 1.35 | 2.68 | 4.97 | 0.74 | 0.037 | 0.07 | 30.47 | 35.87 | 5.04 | 3.41 | 5.73 | 5.00 |
| 8. Enzymes FEA E0356 + XYL E0051 | 30.47 | 8.64 | 2.63 | 30.47 | 35.87 | 10.86 | 8.11 | 16.21 | 1.35 | 2.68 | 4.97 | 0.74 | 0.037 | 0.07 | 30.61 | 35.87 | 5.04 | 3.41 | 5.73 | 5.00 |
a. The data reported for AX concentration and amount in the bran appear to be erroneous, as they indicate less AX in the raw material than in the residual pellet; a starting concentration of 24% is more in line with the mass balance and with the expected AX content of wheat bran.

b. The results for the pellet and high- and low-cutoff fractions following just buffer extraction are reported in these EE rows, but these are not to be understood as having undergone enzyme treatment.

c. Initial weights averaged from Trials 2 and 3.
Table 3. Crude yields, AX concentrations and AX yields from sugarcane bagasse and its fractions following extraction under different treatments.

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<td>11.76</td>
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<td>11.54</td>
<td>11.54</td>
<td>6.29</td>
<td>6.15</td>
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<td>19.54</td>
<td>19.54</td>
<td>19.54</td>
<td>19.54</td>
<td>19.54</td>
<td>19.54</td>
<td>19.54</td>
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<td>2.30</td>
<td>2.25</td>
<td>2.25</td>
<td>2.25</td>
<td>1.23</td>
<td>1.20</td>
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<td>CE Pellet (g)</td>
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<tr>
<td>AX concentration (%)</td>
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<td>1.10</td>
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<td>Absolute yield (%)</td>
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<td>CE Low (g)</td>
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<td>AX concentration (%)</td>
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<td>0.96</td>
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<td>0.06</td>
<td>0.04</td>
<td>0.03</td>
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<tr>
<td>Absolute yield (%)</td>
<td>0.61</td>
<td>0.52</td>
<td>0.37</td>
<td>0.24</td>
<td>0.38</td>
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<td>2.97</td>
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<tr>
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<td>0.01</td>
<td>0.03</td>
<td>0.03</td>
<td>0.01</td>
<td>0.02</td>
<td>0.06</td>
<td>0.05</td>
</tr>
</tbody>
</table>
### Absolute yield (%)

|          | 0.68 | 0.12 | 0.23 | 0.29 | 0.12 | 0.19 | 1.00 | 0.89 |

- a. The results for the pellet and high- and low-cutoff fractions following just buffer extraction are reported in these EE rows, but these are not to be understood as having undergone enzyme treatment.
- b. Initial weights averaged from Trials 2 and 3.
Figure 1. Procedure for chemical extraction of AX from wheat bran or sugarcane bagasse, yielding High cut-off (CE-High) and Low cut-off (CE-Low) fractions and a residual pellet; and procedure for further enzyme treatment of the pellet to yield further High (EE-High) and Low (EE-Low) cut-off fractions.
Figure 2. Measurement of arabinoxylan content in wheat bran following different hydrolysis conditions.
Figure 3. (a) Yield, (b) Concentration and (c) Absolute Yield in fractions following arabinoxylan extraction from wheat bran.
Figure 4. (a) Yield, (b) Concentration and (c) Absolute Yield in fractions following arabinoxylan extraction from sugarcane bagasse.
Figure 5. Expanded view of Absolute Yields in fractions following arabinoxylan extraction from sugarcane bagasse under various chemical and enzyme treatments.
Figure 6. Arabinose:Xylose ratios in fractions following arabinoxylan extraction from (a) wheat bran and (b) sugarcane bagasse.