



Microwave-assisted hydrothermal processing of pine nut shells for oligosaccharide production

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Abstract

Pine nut shells, a biomass residue from the Mediterranean *Pinus pinea* pine nut industrial processing, were treated by microwave-assisted autohydrolysis to produce xylo-oligosaccharides. Microwave-assisted processes provide alternative heating that may reduce energy input and increase overall process efficiency. The autohydrolysis treatments were performed under isothermal and non-isothermal operations within a wide range of operational conditions (temperature/reaction times) covering several severity regimes (as measured by the log R_0 severity factor). The composition of the autohydrolysis liquors was determined in terms of oligo- and monosaccharides, aliphatic acids and degradation compounds. The process was highly selective towards hemicelluloses hydrolysis and liquid streams containing a mixture of oligomeric compounds (mainly xylo-oligosaccharides) could be obtained under relatively mild operation conditions (190 °C, 30 min) with a maximal oligosaccharides' concentration of 18.48 g/L. The average polymerization degree of the obtained oligosaccharides was characterised by HPLC, showing that for the optimal conditions a mixture of oligomers with DPs from 2 to 6.

Keywords Biorefineries · Fractionation · Hemicelluloses · Microwave heating · *Pinus pinea* · Severity factor

Highlights

- Pine nut shell was subjected for the first time to microwave-assisted (MW) autohydrolysis.
- MW autohydrolysis process is highly selective for hemicellulose removal.
- Hydrolysates contained mixture of oligomeric compounds, mainly xylo-oligosaccharides.
- Mild operation conditions (190 °C, 30 min) produced 18.48 g/L of oligosaccharides.
- Oligosaccharide polymerization degree varied from 2 to 6.

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1 Introduction

Pinus pinea L., stone pine or umbrella pine, is a typical Mediterranean forest species, mostly relevant in Portugal and Spain (especially in their southern regions) where about 70% of the *P. pinea* pine nut world production concentrates. The pine nut seeds have a thick ligneous shell that encloses the pine kernel and represents about 77% of the weight of the pine nuts [1]. These shells are potential biomass sources that have the logistical advantage of being

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concentrated at the industrial processing facility [2]. However, they are currently only used as a low added-value energy source for direct combustion without any valorisation upgrade technologies.

Pine nut shells have a high content of lignin and polysaccharides, with low extractives and ash content, which suggest possible processing routes in the biorefinery framework, namely targeted to a polysaccharide-based conversion [1]. The hemicelluloses consist mainly of xylans with very low acetylation and uronic units and a low proportion of galactoglucomannans. Hemicelluloses may be converted into oligosaccharides, compounds that have been increasingly valued in food and pharmaceutical industries, and have received a large attention in recent research [3–5].

Conversion of lignocellulosic biomass to biofuels and value-added bio-products has been attracting much attention in the research and industrial sectors, although some constraints exist for the economic feasibility in biorefineries [3, 4]. The development of efficient, cost-effective and eco-friendly pretreatment processes is one challenge in biomass conversion [6]. Pretreatments can be physical (mechanical, radiation), chemical (e.g. alkali, acid, organosolv, ionic liquid-based) or physical-chemical (e.g. steam explosion, ammonia fibre explosion). Hydrothermal processes such as autohydrolysis present several technological advantages and are therefore one of the best biomass pretreatment options [7]. No chemicals other than water are used, and low equipment corrosion and by-product generation are expected to occur, but it requires high temperature, typically in the range of 150–250 °C. The present energy context advises the use of alternative heating techniques that may reduce the energy input and increase the total process efficiency.

In this framework, microwave processing has attracted attention both from the R&D and industrial sectors since it satisfies many requirements of green chemistry [8]. Microwaves are electromagnetic irradiation in the frequency range of 0.3 to 300 GHz, whose energy per photon is less than 10^{-3} eV, even at the highest frequency. This is too low to induce ionization and to break chemical bonds and has also lesser energy than the Brownian motion [9]. Microwaves apply an electromagnetic field directly to the molecular structure of the heated material leading to physical, chemical or biological reactions. Microwave-assisted extraction is one of the techniques that allows faster and larger extractions of several compounds, including phenolic compounds, with several advantages over other methods [10, 11]. Among other advantages, microwaves allow the rapid heating of aqueous mixtures with non-ionising electromagnetic radiation, lower solvent use, greater selectivity for targeted compounds, better efficiency and lower extraction times in comparison with other heating systems (e.g. 10 times less), thereby decreasing the energy consumption and allowing additional automation [12].

The efficient heating of materials by microwave dielectric heating effects is the basis for microwave-enhanced chemistry [13]. This is very attractive for many chemical applications and is now widely accepted as a non-conventional energy source for performing organic synthesis as shown by the increasing number of related publications in recent years and with the general availability of new and reliable microwave instrumentation [14]. Microwave heating has been successfully applied in biomass pretreatments where in addition to the heating process it causes swelling and fragmentation [15–17].

This work studies the use of microwave-assisted hydrothermal pretreatment of pine nut shells under various isothermal and non-isothermal conditions for the production of oligosaccharides and analyses their composition and polymerization degree.

2 Experimental

2.1 Feedstock

Pine nut shells were provided by a Portuguese pine nut processing industry. Upon reception, the shells were milled with a benchtop grinder and screened to obtain a homogenous size lot (1–3.55 mm). The material was stored in vacuum-sealed bags until required for processing or analysis. Granulometric characterization of the milled sample was carried out in a sieve shaker (Endecotts, England) with sieves from 1.00 to 4.00 mm. The material (approximately 100 g) was screened for 20 min, and the fraction retained on each sieve was weighed to quantify the respective mass fraction. These assays were carried out in duplicate.

2.2 Microwave-assisted autohydrolysis

The microwave-assisted autohydrolysis assays were performed in a Microwave Digestor (Ethos Easy, Milestone Srl, Italy) using a magnetron frequency of 2450 MHz and EasyWave software for monitoring and control. Pine nut shells were reacted with water in capped Teflon vessels, with a loading of 15 g of dry biomass and 45 g of water, corresponding to a liquid-to-solid ratio of 3:1 (mass ratio of H₂O/biomass). Several isothermal and non-isothermal conditions were tested, and the microwave power was continuously varied to maintain the prescribed temperature profile. The overall temperature profiles were recorded for all treatments (see below).

In non-isothermal conditions, the reactors were heated to a final temperature between 170 and 230 °C at a rate of 3.6 °C/min, as in a conventional non-isothermal autohydrolysis process [18]. In isothermal assays, the temperature was fixed at 190 °C which was attained also with a rate of 3.6 °C/min, and different isothermal periods between 0 and 60 min were

tested. After reaching the targeted temperature or reaction period time, the reactors were rapidly cooled by immersion in an ice bath. All assays were carried out in triplicate.

The separation of the liquid and solid streams was made by using filtering crucibles under vacuum. The liquid fraction was further filtered through quantitative filter paper (Whatman nr. 41) to remove any remaining solids. The solid fraction was washed with two volumes (90 g each) of water, filtered and frozen before chemical composition analysis. For comparison purposes, the severity of the treatments was estimated as summatory of heating, isothermal and cooling step calculating the log R_0 , based on the measured temperature profile data for each step [19] according to Eq. (1).

$$R_0 = \int_0^t \exp\left(\frac{T-100}{14.75}\right) dt \quad (1)$$

2.3 Chemical composition analysis

The initial pine nut shells and the solid residues obtained after microwave-assisted autohydrolysis were characterised following standard NREL protocols [20]. Shortly, this characterization consists firstly in dissolving 0.3g of material to analyse in 3 mL 72% (w/w) sulfuric acid at 30 °C for 1h. Afterward, the acid concentration is diluted to 4% (w/w) and then all mixture is subjected to a heating process in an autoclave for 1h at 121 °C.

The liquid fraction was analysed in relation to solubilised monosaccharides (glucose, xylose and arabinose), aliphatic acids (formic, acetic and levulinic) and furan derivatives (5-hydroxymethylfurfural and furfural) by high-pressure liquid chromatography using a Thermo Scientific system (USA), equipped with a refractive index detector (Refractomax521) controlled at 39 °C and a diode array detector (DAD 3000), using a Bio-Rad Aminex HPX-87H column (300 × 7.8 mm) (Hercules, CA). The mobile phase was H₂SO₄ 5 mM, the column temperature 50 °C, the flow rate 0.6 mL min⁻¹ and the injection volume was 5 μL. Mannose was quantified in the same chromatographic system using a column REZEX RPM-Monosaccharide (Pb²⁺, 8% cross-linked Pb²⁺; 300 × 7.8 mm, 9 μm; Phenomenex) at a column temperature of 85 °C and a flow rate of 0.6 mL/min. All samples were filtered with Millipore® (Cork, Ireland) 0.45 μm cellulose acetate membrane filters prior to analysis.

The oligosaccharides were determined indirectly after quantitative acid hydrolysis according to NREL/TP-510-42623 [21]. Concentrated sulfuric acid (72%) was added to an aliquot of the liquor resulting from the autohydrolysis treatments, in order to attain a 4% H₂SO₄ concentration, and hydrolysed in an autoclave at 121 °C for 1 h. After completion, the hydrolysates were slowly cooled down to room temperature, a sample was collected, filtered using 0.45 μm

membranes (Millipore) and analysed by HPLC. The oligosaccharide concentration was calculated from the increase in sugar monomers in relation to their direct determination in the liquor. This procedure was performed in duplicate.

2.4 Polymerization degree of oligosaccharides

For characterization of the polymerization degree of oligosaccharides, liquors were injected in a REZEX RSO-Oligosaccharide (Phenomenex, Torrance, CA, USA) column filled with a sulphonated polystyrene-divinylbenzene resin in the Ag⁺-form (200 × 10.00 mm I.D., 4% cross-linking, particle size 8 μm), equipped with a guard column with the same filling, with the column at 75 °C. Ultrapure water at 0.3 mL/min was used as an eluent. Calibration for polymerization degree (PD) was carried out using a light corn syrup sample (mainly composed of malto-oligosaccharides) from Phenomenex. The calibration curve obtained is presented as supplementary material.

3 Results and discussion

3.1 Pine nut shell characterization

A granulometric characterization was performed after the size reduction of the pine nut shells (Fig. 1). Very few fines were produced in accordance with the brittleness of the pine shells. The 2.36 to 3.55 mm fraction accounted for more than 45% of the material and particles ranging from 1.00 to 3.55 mm corresponded to 75% of the total mass. This particle diameter range is within what is described in the literature as the most suitable for autohydrolysis processes [22].

The chemical characterization of the pine nut shells is detailed in Table 1, together with values reported in other

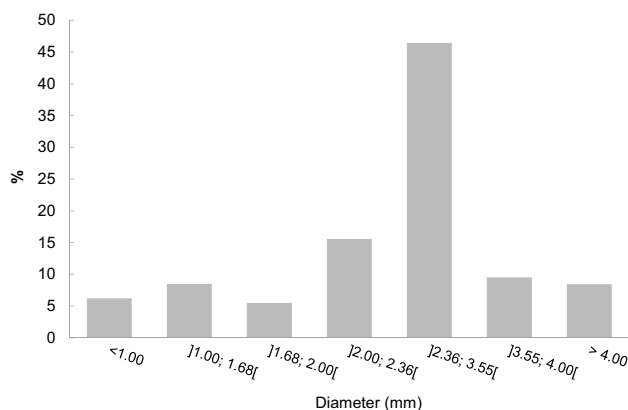


Fig. 1 Granulometric characterization of pine nut shell after milling using a laboratory benchtop grinder

Table 1 Summative chemical composition (% of total dry mass) of the *P. pinea* pine nut shells lot used in this work and comparison with reported literature values for *P. pinea* [1, 18, 23] and *P. koraiensis* [24]

| Component (%) | <i>P. pinea</i> | | | | <i>P. koraiensis</i> |
|----------------|-----------------|--------------|------|-----------|----------------------|
| | This work | [18] | [1] | [23] | [24] |
| Glucan | 25.90 ± 0.21 | 25.93 ± 0.06 | 29.1 | 29.5–32.4 | 36.13 |
| Hemicelluloses | 18.23 ± 0.15 | 20.98 ± 0.04 | 18.2 | 28.5–30.3 | 21.97 |
| Xylan | 11.52 ± 0.07 | 16.27 ± 0.05 | 12.7 | n.d | n.d |
| Mannan | 5.17 ± 0.05 | | 4.0 | n.d | n.d |
| Arabinan | 1.07 ± 0.24 | 2.72 ± 0.01 | 1.2 | n.d | n.d |
| Acetyl groups | 0.47 ± 0.03 | 1.99 ± 0.01 | 0.3 | n.d | n.d |
| Klason lignin | 44.48 ± 0.16 | 42.63 ± 0.24 | 39.9 | 37.6–40.1 | 39.59 |
| Ash content | 2.90 ± 0.10 | 3.04 ± 0.14 | 1.3 | n.d | n.d |
| Extractives | 5.10 ± 0.06 | 2.67 ± 0.13 | 4.5 | n.d | n.d |

[1] report 1.2% galactose and 0.2% uronic acids; in [18], the value of 16.27% for xylan includes xylose and mannose; *n.d.* not determined

studies for pine nut shells of *P. pinea* [1, 23] and *Pinus koraiensis* [24].

The pine nut shells used in this study have a high lignin content (44.5%), and a significant content of polysaccharides, including cellulose (25.9%, estimated as glucan), and hemicelluloses (18.2%, as the sum of xylan, mannan, arabinan and acetyl groups). This composition is similar to values found in the literature for pine nut shells of *P. pinea* [25] and *Pinus koraiensis* [24, 25].

The pine nut shell hemicelluloses comprise mostly xylans with low proportions of arabinose units and acetyl substitutions, corresponding to about 13.1% of the material. This is not typical for softwood species for which galactoglucomannans are the main wood hemicelluloses. Mannan is however still significant, being almost 30% of the hemicellulosic sugars. Galactose was not detected and non-cellulosic glucose was not determined. Although Queirós et al. [1] obtained 1.2% of galactose, the methodology used in the present work (NREL vs TAPPI) did not allow its detection. Also, it must be kept in mind that there could be a variability in the composition of the pine nut shell. On the other hand, the pine nut shell may not have the same hemicellulosic composition as softwood species despite galactoglucomannans being its major component [26].

The chemical characterization of biomass materials is determined for evaluation of their potential for the biochemical/chemical platform of biorefineries. A significant content in hemicelluloses allows considering it for the production of oligosaccharides, as it occurs in the present study.

3.2 Microwave-assisted autohydrolysis under non-isothermal conditions

The microwave-assisted autohydrolysis process was studied using non-isothermal and isothermal conditions. The heating profiles for the various experiments are linear and overlap, showing a high degree of reproducibility of the equipment

operating conditions, as shown in the Supplementary material (S1).

The pine nut shells were subjected to microwave-assisted autohydrolysis treatments under non-isothermal conditions, i.e. the reaction was stopped when the desired final temperature was attained. Final temperatures between 170 and 230 °C were tested, corresponding to the following severities: log R_0 2.82 (170 °C), log R_0 3.45 (190 °C), log R_0 3.78 (200 °C), log R_0 4.10 (210 °C), log R_0 4.66 (230 °C), log R_0 5.17 (250 °C). The heating profile was similar to that used in a previous study using standard autohydrolysis of pine nut shells [18] carried out in a Parr reactor with electric heating through an external mantle. Figure 2 shows the composition of the autohydrolysis liquor as a function of the severity factor.

Oligosaccharides (OS) were the most relevant products, reaching a total concentration of 6.4 g/L for log R_0 4.66, which corresponds to a yield of 1.59% (g XOS/100g pine nut shells). The hemicellulose-derived oligosaccharides contain xylose, arabinose and mannose as main sugars, and acetyl groups, reflecting the composition of the pine nut shells hemicelluloses. They are here referred to as xylo-oligosaccharides as xylose accounts for the majority of the hemicellulosic sugars. Gluco-oligosaccharides are present with a low concentration corresponding to 5% of the total oligosaccharides.

The concentration of xylo-oligosaccharides increased from 0.16 to 5.96 g/L and decreased after that. The severity condition leading to the highest production of oligosaccharides (log R_0 4.66) is higher than for other lignocellulosic materials that typically present values close to 4 [27, 28]. Pine nut shells are more recalcitrant to hydrolysis and will require more severe operational conditions at the industrial level.

For the most severe condition tested (log R_0 5.17), the xylo-oligosaccharide concentration decreased abruptly to 0.89 g/L, as a result of their depolymerization to

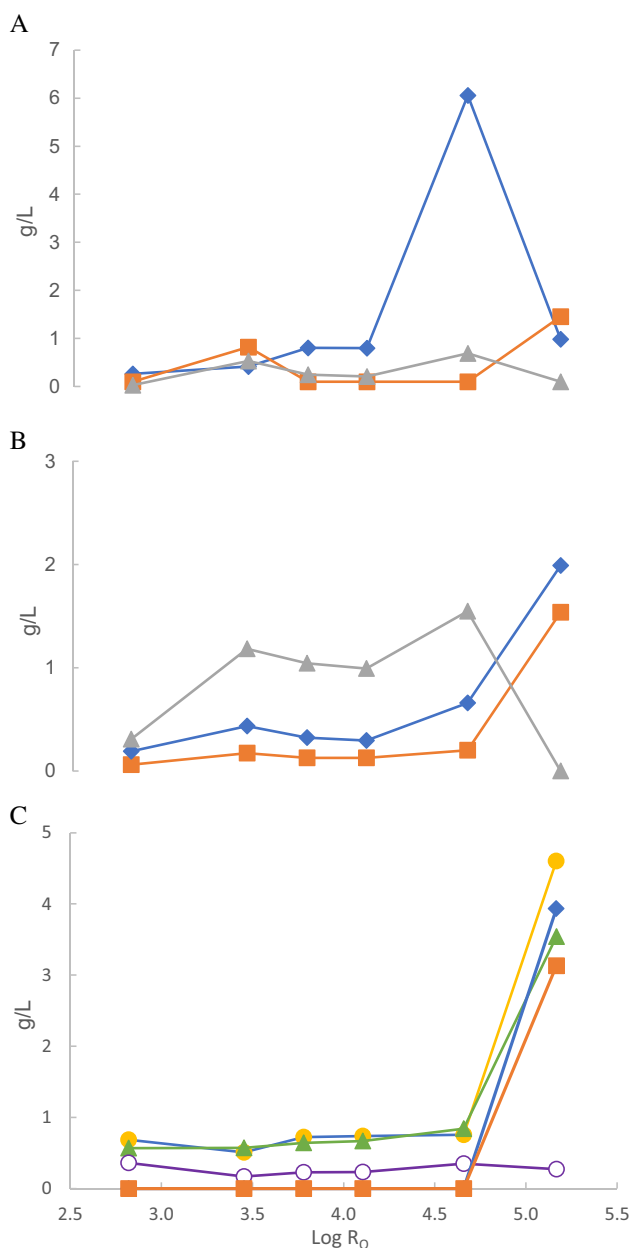


Fig. 2 Composition (g/L) of the liquid fractions obtained after non-isothermal microwave-assisted autohydrolysis of pine nut shells as a function of $\log R_0$: **A** filled diamond, xylo- and mannan-oligosaccharides; filled triangle, arabino-oligosaccharides; filled square, gluco-oligosaccharides; **B** filled square, glucose; filled diamond, xylose and mannose; filled triangle, arabinose; **C** filled circle, acetic acid; filled triangle, formic acid; empty circle, levulinic acid; filled diamond, furfural; filled square, HMF. Lines are used for eye guide only

monosaccharides and the degradation of monosaccharides to aliphatic acids and furans, which increased significantly. Such carbohydrate behaviour in autohydrolysis has been reported for several biomass materials and is typically observed for severities above 4.01 [29–31]. This is also consistent with studies on wood and other agricultural and

agro-industrial residues reporting the autohydrolysis selectivity towards hemicellulose depolymerisation [32, 33].

The results show that autohydrolysis has a higher selectivity towards xylans, i.e. higher yields were obtained for xylo-oligosaccharides than for gluco-oligosaccharides in a proportion much above their original ratio in the pine nut shells hemicelluloses. Chemical bonds between xylose units in the xylan backbone as well as chemical bonds to arabinose were more easily cleaved than glucose and mannose bonds in glucomannans, as it is well known from the overall hydrolysis reactivity of the different hemicelluloses [34, 35].

The microwave-assisted autohydrolysis was less effective when compared to standard autohydrolysis, for which an equivalent maximal oligosaccharides yield was obtained for lower severity ($\log R_0$ 4.01, 16.11 g/L) [18]. This suggests that autohydrolysis, either microwave-assisted or conventional, requires for optimal oligosaccharides production process severities between 4 and 4.5.

3.3 Microwave-assisted autohydrolysis under isothermal conditions

The microwave-assisted autohydrolysis was evaluated in more detail for process severities between 4 and 4.5, using an isothermal process at 190 °C with different durations of the isothermal periods from 0 to 60 min. The results are shown in Fig. 3. The best condition, with a xylo-oligosaccharides recovery of 34.64 g/100 g of the xylan in pine nut shells (concentration of 16.49 g/L), was obtained at $\log R_0$ 4.2. This corresponds to a higher yield as compared to that obtained with a standard autohydrolysis of 28.74 g/100 g xylan [18], which was obtained with a $\log R_0$ 4.01.

For non-isothermal processes at comparable severity, microwave-assisted autohydrolysis seemed to show a lower performance than standard autohydrolysis, which could be explained by the fact that the ideal conditions have been exceeded. Precisely the isothermal conditions showed that in fact microwave treatment was more efficient than non-isothermal with standard methods.

Oligosaccharide hydrolysis to monosaccharides started simultaneously and slightly ahead of hemicellulose depolymerization to oligosaccharides. Xylose and mannose increased strongly in the autohydrolysis liquor from $\log R_0$ 4.0 to 4.15 and remained constant subsequently until their decrease starting at $\log R_0$ 4.47 (Fig. 3B). Arabinose content increased up to $\log R_0$ 4.2 and decreased subsequently to zero at $\log R_0$ 4.47. Glucose contents were rather stable with the different severities.

Monosaccharide degradation accompanied their presence in the autohydrolysis liquor, with a considerable increase of furan and aliphatic acids, therefore starting before the maximum oligosaccharides yield. Acetic acid started to accumulate from $\log R_0$ 3.96 onwards, reaching

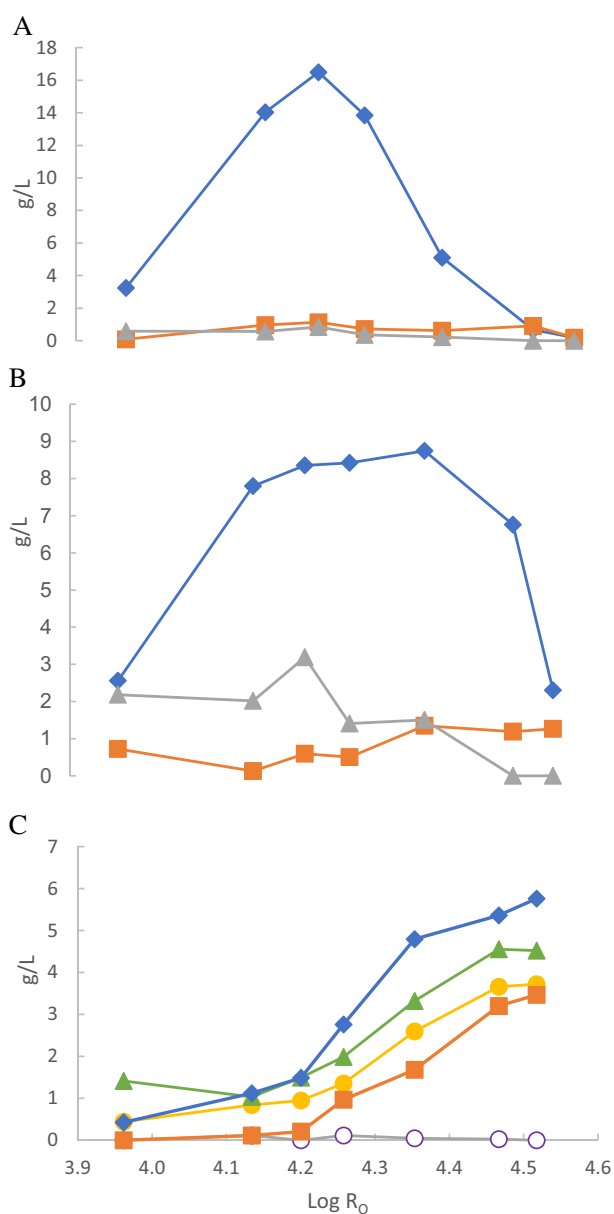


Fig. 3 Composition (g/L) of the liquid fractions obtained after isothermal microwave-assisted autohydrolysis, as a function of $\log R_0$: **A** filled diamond, xylo- and mannan-oligosaccharides; filled triangle, arabino-oligosaccharides; filled square, gluco-oligosaccharides; **B** filled square, glucose; filled diamond, xylose and mannose; filled triangle, arabinose; **C** filled circle, acetic acid; filled triangle, formic acid; empty circle, levulinic acid; filled diamond, furfural; filled square, 5-hydroxymethylfurfural. Lines are used for eye guidance only

3.72 g/L. Furfural and HMF were found in low concentrations, higher for furfural (1.49 g/L) given the higher amounts of pentoses in the pine nutshell hemicelluloses and their higher dehydration susceptibility. Other sugar degradation by-products such as levulinic and formic acids were also present at low concentrations [36].

The total aliphatic acid concentration (including acetic acid) for the optimal xylo-oligosaccharide production condition was below 3 g/L, thereby not precluding, as potential microbial inhibitors, the possible future use of this hydrolysate for fermentation processes [36].

Overall, microwave-assisted autohydrolysis of pine nut shells allowed the production of oligosaccharides in values similar to those obtained with conventional autohydrolysis of the same material. This result was slightly different from studies performed by obtained by del Río et al. [37] that compared conventional and microwave-assisted autohydrolysis for the obtention of oligosaccharides from Paulownia obtaining the higher values of OS, with milder conditions with microwave in comparison to conventional water treatment. Taking into account the raw material composition and structure differences in operational and energy conditions of the autohydrolysis is a key factor for evaluating both processes.

3.4 Degree of polymerization of oligosaccharides

The characterization of the polymerization degree (DP) of the oligosaccharides obtained for the different severities of the isothermal autohydrolysis of pine nut shells was performed. The REZEX™ 195 RCM-Monosaccharide column and a malto-oligosaccharide standard including maltose, maltotriose, maltopentose and maltohexose with DPs of 2 to 6, respectively, were used to estimate the DP of the obtained pine nut shells oligosaccharides. With this column, the oligomers with higher DP have shorter retention time [38].

Figure 4 shows the chromatographic profiles of seven pine nut shell hydrolysates obtained at the different severity conditions as well as of the malto-oligosaccharide standard (standard OS). The standard profile clearly shows the separation of all five oligosaccharide standards from DP2 to DP6. The oligosaccharides of the seven hydrolysates included disaccharides (DP2), trisaccharides (DP3), tetrasaccharides (DP4), pentasaccharides (DP5) and hexasaccharides (DP6). The peak for the column retention time of approximately 45 min corresponds to xylose.

Depending on the autohydrolysis severity, different molecular weight distributions were found for the produced oligosaccharides. For the mildest autohydrolysis conditions, the obtained oligosaccharides were oligomers with 6 and above monomers. With increased severity, the oligosaccharide molecular weight progressively decreased, leading preferably to the accumulation of dimers and trimers.

Figure 5 shows the concentration profile of the various oligosaccharides with the increase in autohydrolysis severity. At the conditions leading to the highest concentration ($\log R_0$ 4.2), the oligosaccharides were a mixture of oligomers with 2 to 5 units, with the highest proportion being dimers with a concentration of 3 g/L. Oligomers with DP4

Fig. 4 Molecular weight distribution of the soluble oligosaccharides obtained by isothermal microwave-assisted autohydrolysis of pine nut shells at 190 °C and different severities

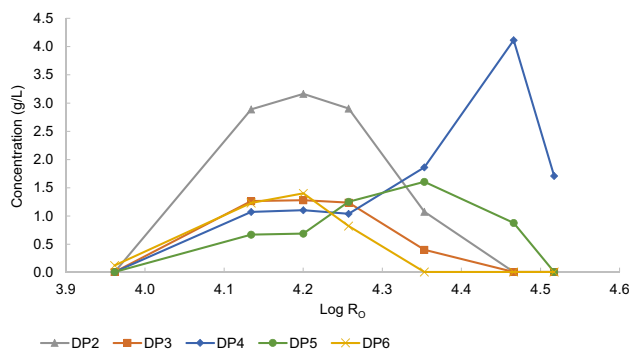
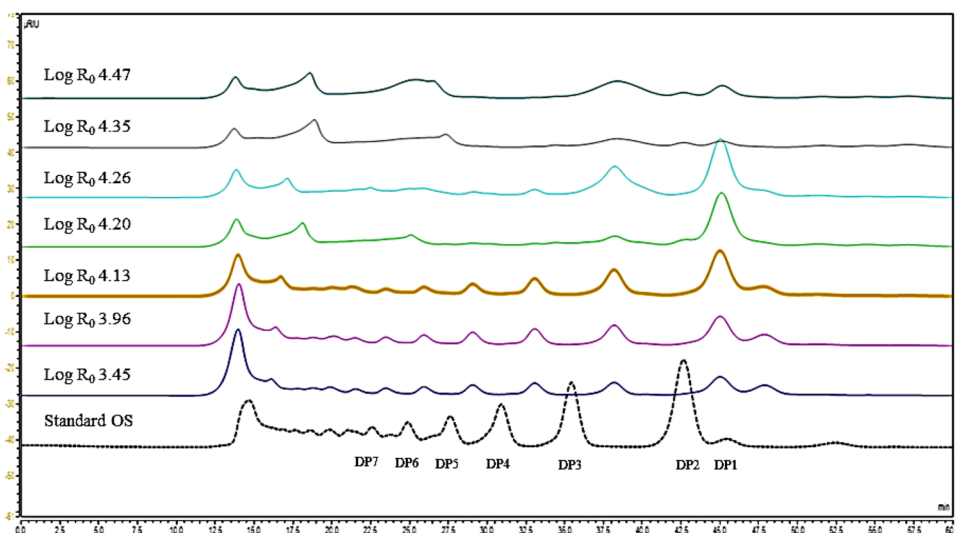


Fig. 5 Concentration of oligosaccharides (g/L) with different degrees of polymerization (DP2 to DP6) obtained from pine nut shells by isothermal microwave-assisted autohydrolysis at 190 °C with different severities

start to increase after $\log R_0$ 4.35, and are in the highest quantity at $\log R_0$ 4.47, reaching a concentration of 4.1 g/L. DP6 oligomers reach the maximum value (1.4 g/L) at $\log R_0$ 4.2. Monosaccharides measured in the RSO column (not shown in Fig. 5) follow the same trend as those measured in the Aminex 87H column, as previously discussed [39, 40].

The biological activity of xylo-oligosaccharides depends directly on their DP that typically ranges from 2 to 12. Xylo-oligosaccharides with $DP \leq 4$ have prebiotic applications because they promote the propagation of beneficial bacteria in the intestinal tract, such as *bifidobacteria*, which inhibit the growth of pathogenic bacteria [41, 42]. Uronic acid-containing xylo-oligosaccharides are known to present antioxidant and antiallergic properties [42].

The pine nut shell-derived oligosaccharides by autohydrolysis have previously been tested and proved chemically stable in time, and for temperature and pH under relevant conditions for stomachal digestive conditions, thereby

supporting their potential nutritional/bioactive properties [18]. The ability of these oligosaccharides to be processed and to pass the stomach undigested deems them as potential non-digestible oligosaccharides, a trait that should be further characterised and explored.

3.5 Composition of the solids

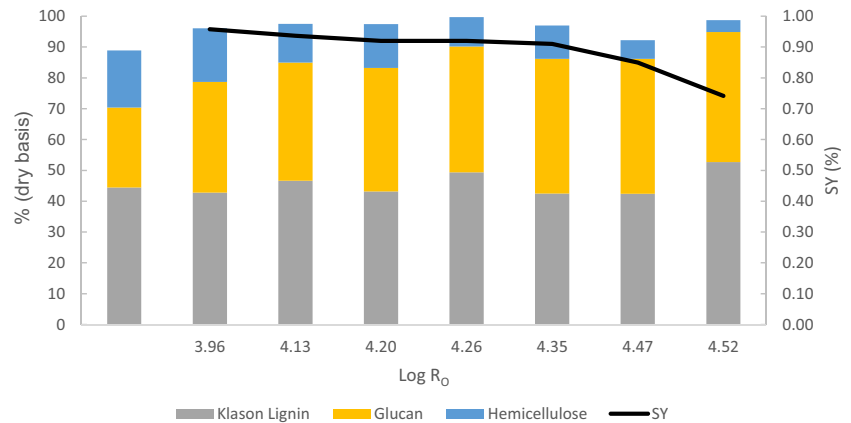
Within a biorefinery framework, it is necessary to envisage the full resource use, and therefore the biomass remaining after the microwave-assisted autohydrolysis has to be evaluated and valorised. Figure 6 shows the composition of the pine nut shells and of the solids obtained by autohydrolysis with the different hydrolysis severities and solid recovery. The values are given in mass proportion of each solid material.

The higher solid recovery of solids denotes the recalcitrance of the raw material related to the treatment. Only at severe conditions, the recovery decreased.

After autohydrolysis, the processed solids increase their glucan contents, due to the decrease of hemicelluloses that were solubilised to the liquor. The microwave-assisted process is selective for hemicellulose removal that increases with the severity of the treatment, and therefore the glucan proportion increased in the solids. This is consistent with previous results of biomass autohydrolysis, which were characterised to have a low effect on the cellulose and lignin fractions [30, 43].

The composition of the spent pine nut shells points out that they may be valorised by two pathways, either a biochemical route targeted to cellulose or a thermochemical route towards energy materials. Cellulose enzymatic hydrolysis was already tested with poor saccharification yields [18]. More severe chemical fractionation procedures are possibly necessary given that the lignin in pine nutshell is of the G type and therefore has low reactivity [1]. Current uses,

Fig. 6 Solid recovery and composition of the pine nut shells of solids in relation to structural components (% of the sum of lignin, glucan and hemicelluloses) obtained after microwave-assisted autohydrolysis of pine nut shells under isothermal conditions at 190 °C, with different severity factors



namely as solid biofuel for combustion, are certainly possible, but other upgrading processes in the thermochemical platform of the biorefinery (e.g. biochar or bio-oil production) are of interest given that the biomass has a high heating value (HHV) favoured by the high lignin and low hemicelluloses content [43, 44]. In fact, most works of research carried out so far on pine nut shell valorisation have considered pyrolysis and biochar production [24, 45].

4 Conclusions

Pine nut shells are a good source for the production of oligosaccharides, namely xylo-oligosaccharides, which can be easily produced under environmentally friendly conditions using microwave-assisted autohydrolysis that allows a better performance by using milder operation conditions. The OS degree of polymerization varied from 6 to 2, being enriched with the highest DP and the lowest inhibitors at the best conditions. The spent solids are enriched in cellulose and can be further directed towards the biochemical or thermochemical platforms within the biorefinery framework.

Supplementary Information The online version contains supplementary material available at <https://doi.org/10.1007/s13399-023-05244-z>.

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Declarations

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