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Effect of autohydrolysis and ionosolv treatments on eucalyptus fractionation and recovered lignin properties†

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Wood fractionation is key for the integral valorization of its three main components. In this sense, recovering the hemicellulosic fraction after the ionosolv treatment of lignocellulosic materials is one of the main drawbacks of this process. Thus, the incorporation of a previous autohydrolysis step to recover the hemicellulosic sugars before the ionosolv treatment is an interesting approach. The influence of both treatments, autohydrolysis and ionosolv, on the biomass fractions recovery yields was studied by a central composite design of experiments, varying the autohydrolysis temperature in a 175–195 °C range and ionosolv time between 1–5 h. Lignin recovery and cellulose purity were maximized at 184 °C and 3.5 h of autohydrolysis temperature and ionosolv time, respectively. In addition, lignin properties were incorporated to the statistical model, revealing lignin recondensation at severe conditions and a higher influence of the ionosolv treatment on lignin characteristics. These results remarked the importance of studying the effect of both treatments in the whole fractionation process and not each process separately and enhanced the understanding of the treatments combination in a complete fractionation biorefinery approach.

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1 Introduction

Over the last years, energy consumption has increased due to the technological advances in our world, causing an increment in fossil fuel utilization, decreasing fossil fuel reserves and increasing the level of greenhouse gas emissions, which has raised the Earth's average temperature by 1.9 °C above pre-industrial levels.^{1–3} To overcome this, biorefinery emerged as one of the most interesting alternatives, considering biomass utilization as substitute of fossil fuels.^{4,5} One of the most employed biomass sources is the lignocellulosic biomass due to its availability, abundance, and the fact that it does not conflict with the food sector. It is also cheaper than other types of agricultural raw materials, can be produced locally and has low nitrogen and sulfur contents.^{5,6} Its exploitation can be done through different approaches, being the biomass fractionation into its main components one of the most promising ones, and thus developing technically and economically viable processes for a biorefinery based on cellulose, hemicellulose, and lignin.⁷ A correct fractionation of lignocellulosic biomass is a key step in achieving economically competitive biorefineries, being able to

produce fuels, chemical products, and high value-added materials.⁸

Among the main biomass components, lignin has been underutilized and just burned for energy, leading to an ineffective economic efficiency.^{8,9} After cellulose, lignin is the second-most abundant biopolymer on Earth, making up 15–30% (w/w) of lignocellulosic biomass.¹⁰ In addition, it is the biggest biosource of aromatic compounds, which has led to growing interest in its economically competitive utilization.^{11,12} Lignin presents an heterogeneous structure and composition; thus, it can be employed in many different applications, for example, as a source of carbon fibers that can be mixed with biopolymers to enhance lithium-ion battery performance, dispersants, thermal reinforcements, coatings, adhesives, and as a source of platform chemicals (xylene, benzene, vanillin, *etc.*).^{13–18} It has also been used in the medical field, for example, in the formulation of hydrogels that can be employed as wound dressings, or as an additive in tablet formulation for drug release.^{19–22} Lately, lignin has attracted great attention in the formulation of lignin nanoparticles, overcoming the existing limitations for its employment (high heterogeneity, large particle size, among others).²³

Different processes have been proposed over the last years for lignin isolation and recovery, being one of them the ionosolv process. This process involves the (partial) solubilization of lignin and hemicellulose employing ionic liquids, while the cellulose fraction remains practically intact, making possible to recover the biomass fractions separately for valorization.^{24,25} Then, lignin can be

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precipitated, using water as an antisolvent.²⁶ The ionosolv process has been successfully employed with hardwoods, softwoods, and herbaceous woods, obtaining yields of up to 90% lignin recovery.^{27–33} However, the hemicellulosic fraction is normally difficult to recover after the ionosolv treatment, making the fractionation process less efficient. In this sense, the incorporation of a previous autohydrolysis step to recover hemicelluloses seems an interesting option.

Autohydrolysis is one of the most widely used physico-chemical treatments for the extraction of hemicelluloses from lignocellulosic biomass since it only uses water without any catalyst, which makes it attractive from an economic and environmental point of view.³⁴ The solubilization of hemicelluloses occurs by the action of hydronium ions from water and the release of acetic acid from the hemicelluloses in the biomass, which act as autocatalysts of the process.^{35,36} By means of autohydrolysis, a high recovery of hemicelluloses can be achieved in the liquid phase, being able to solubilize and recover between 50 and 90% of them.^{37,38}

In this work, the combination of autohydrolysis + ionosolv treatments to completely fractionate eucalyptus wood was studied, analyzing the influence of both processes on the recovered fractions. A central composite design of experiments was performed with autohydrolysis temperature, in the range of 175–195 °C, and ionosolv time, 1–5 h, as factors was performed. Hemicelluloses recovery, lignin and cellulose recoveries, and cellulose purity were set as the main responses. In addition, the recovered lignins were characterized in terms of thermal stability and molecular weight distribution. Although this is not the first time that autohydrolysis and ionosolv treatments are combined, it has always been done towards enhancing enzymatic digestibility, and not in a biomass fractionation concept. Thus, this work offers a first approach to a combined fractionation process and enhances the understanding of the influence of autohydrolysis and ionosolv treatments on both the fractionation efficiency and the recovered fraction characteristics.

2 Materials and methods

2.1 Materials

Eucalyptus globulus was used as feedstock and was provided by CIFOR-INIA (“Centro de Investigación Forestal-Instituto Nacional de Investigación y Tecnología Agraria y Alimentaria”, in Spain). The ionic liquid 1-methylimidazolium chloride ([Mim] [Cl], 98% purity) was supplied by Iolitec GmbH. Ethanol was used as antisolvent and supplied by Panreac (96% purity). *N,N*-Dimethylformamide (DMF, ≥99.9%, Panreac) and ReadyCal Kit PMMA standards (Polymer Standards Service GmbH) were used in GPC analysis.

2.2 Design of experiments and statistical analysis

In order to study the influence of the different process variables in the combined fractionation process configuration of autohydrolysis + ionosolv, a 2² with 4 axial points and center point triplicates central composite design using response surface methodology was employed. The studied factors were the

autohydrolysis temperature (T_{AH}) and the ionosolv treatment time (t_{iono}). Hemicellulose recovery (only affected by the first step), lignin recovery, cellulose recovery, and cellulose purity were selected as responses. This selection also remarks the importance of studying the fate of the cellulosic fraction in a fractionation process. In addition, some lignin characterization responses, such as weight-average molecular weight, polydispersity, and degradation temperature at 10% weight loss, were also included.

All the data from the design were statistically analyzed to study the influence of each factor on the fractionation responses and model and optimize the autohydrolysis + ionosolv process. ANOVA was performed in the central composite design. All the statistical tests were done using the OriginPro software.

2.3 Autohydrolysis step

The autohydrolysis treatment was carried out in a 450 mL Parr reactor (Parr Instrument Company, Moline, IL, USA) with a turbine impeller stirrer. Previous autohydrolysis tests were performed to set the autohydrolysis temperature range for the design of experiments of the treatments' combination. Water and raw eucalyptus were mixed in a liquid to solid ratio of 5 kg of water per kg of wood. The mixture was heated up to 165, 175, 185, 195, and 205 °C during 50 min at a constant rate. Then, the set point temperature was maintained for 30 min and cooled at the maximum possible cooling rate (approximately 15 °C min⁻¹). The corresponding severity factors were 3.42, 3.72, 4.00, 4.26, and 4.57 for the selected temperatures. The detailed temperature profile can be found in Fig. S1 in the ESI.† After pretreatment, the solid and liquid phases were separated by vacuum filtration. The solid phase was washed with water, employing twice the volume used for the autohydrolysis treatment. The hemicellulose-free solid was freeze dried and characterized, following the NREL/TP-510-42618 procedure.³⁹ The composition of the resulting liquid phase was determined following the NREL/TP-510-42623 procedure.⁴⁰

2.4 Ionosolv treatment

The ionosolv treatment was performed in 25 mL glass reactors with anchor stirrer due to the high viscosity of the wood/protic ionic liquid (PIL) mixture. The PIL [Mim][Cl] was mixed with the autohydrolyzed wood at a 10% (w/w) biomass loading and heated up to 135 °C, according to our previous work, using a silicone bath and run during different treatment times (see Table 2).⁴¹ After the ionosolv treatment, 0.5 g of pure ethanol per g PIL were added as antisolvent to precipitate a cellulose rich fraction (CRF). The precipitated solid was washed with 2 g of ethanol per g of PIL for each washing step until no PIL was observed in the washing fractions (conductivity of washing fractions approximately 0.3 μS cm⁻¹). The CRF was vacuum dried at 70 °C and stored for further analysis. The ethanol of the PIL/ethanol resulting mixture was removed by vacuum distillation and 3 g of water per g PIL were added as antisolvent to precipitate a lignin fraction which was freeze dried and stored for further analysis. Both solid fractions were characterized following the NREL/TP-510-42618 procedure adapted to small amounts of sample.^{39,42}



2.5 Analytical methods

2.5.1 Liquid phases analysis. Sugar analysis of the different liquid phases was carried out using an Agilent 1260 Infinity HPLC (Agilent Technologies, Santa Clara, CA, USA). A CarboSep CHO-682 lead column, operating at 80 °C, was used for sugar separation before determination with a refractive index detector. Water was used as the mobile phase at 0.4 mL min⁻¹. All samples were neutralized with CaCO₃ before analysis, making necessary to use a Micro-Guard cartridge (BioRad Life Science Group, Hercules, CA, USA) to retain possible salty residues. The analysis of acetic acid and sugars' degradation products was performed using the same HPLC equipment operating with a Phenomenex Rezex-ROA column at 60 °C. In this case, 0.005 M H₂SO₄ at 0.6 mL min⁻¹ was used as mobile phase.

2.5.2 Gel permeation chromatography analysis of lignins. Weight-average (M_w) and number-average molecular weights (M_n), and polydispersity indexes (PDI) of lignins were measured by gel permeation chromatography (GPC) using an Agilent 1260 Infinity HPLC (Agilent Technologies, Santa Clara, CA, USA). Two PolarGel-M 300 × 7.5 mm columns with a PolarGel pre-column guard operating at 60 °C were used. The analysis was performed combining a concentration response (Agilent G1362A RID detector) and a viscosity response (Varian 390-LC viscosity detector), so the polymeric universal calibration was obtained and, therefore, the values can be considered as absolute values and not relative to the calibration.^{43,44} *N,N*-Dimethylformamide (DMF) + 1 g L⁻¹ LiBr was used as the mobile phase at 0.8 mL min⁻¹. ReadyCal Kit PMMA standards (M_w = 800–2 200 000 Da) from Polymer Standards Service GmbH were used for the equipment calibration. All samples were prepared at a concentration of 1 mg mL⁻¹ and filtered (PTFE, 0.22 μm filters) before analysis.

2.5.3 Thermogravimetric analysis of lignins. Thermal stability studies of lignins were performed in a Mettler-Toledo TGA/DSC 1e. Approximately 8 mg of sample were placed in a 70 μL alumina crucible and dry off by heating them from 30 up to 105 °C at a heating rate of 10 °C min⁻¹, followed by a 30 min isotherm at 105 °C. Then, samples were heated from 105 up to 950 °C at a heating rate of 10 °C min⁻¹. Nitrogen was used as reaction and protection gas at 20 mL min⁻¹. Degradation temperature at 10% mass loss ($T_{D_{10\%}}$) was established as an indicative of thermal stability of lignins.

2.5.4 Chemical structure of lignins. Lignins were analysed by HSQC-NMR to study their chemical structure. Approximately 30 mg of lignin were dissolved in 750 μL of DMSO-d₆ and analysed in a Bruker AVIII spectrometer at 700.17 MHz, according to our previous work.⁴⁶

3 Results and discussion

3.1 Autohydrolysis temperature (T_{AH}) and ionosolv time (t_{iono}) interval selection

A previous study of autohydrolysis with eucalyptus wood in the range of 165–205 °C was carried out to determine the temperature operating range in the design of experiments for the

autohydrolysis + ionosolv treatments combination. The composition of the obtained liquors is shown in Table 1.

Hemicelluloses concentration increased with treatment temperature up to a maximum at 185 °C, indicating a higher sugar solubilization and conversion to their dissolved oligomers and to monomeric sugars. This behavior can be most clearly observed in xylo-oligosaccharides and xylose, due to the composition of eucalyptus hemicelluloses (mostly xylan). A xylo-oligosaccharides concentration maximum of 26.10 g L⁻¹ can be found at 175 °C, then decreasing. The concentration of free xylose increased to 14.12 g L⁻¹ at 195 °C. This fact is due to a higher conversion of xylo-oligosaccharides to xylose with increasing temperature. However, the xylose concentration decreased significantly at 205 °C, dropping to 5.41 g L⁻¹. An increase in the furfural concentration was also observed since it is the main dehydration product of pentoses. As for glucose, an increase in the concentration of gluco-oligosaccharides and glucose was observed with temperature. However, the concentrations of both gluco-oligosaccharides and glucose are lower than 3 g L⁻¹, reflecting that the structural glucan was not solubilized.⁴⁵

Considering the hemicellulose recovery yields (Table 1), calculated as the total hemicellulosic sugar content in the autohydrolysis liquor compared to the hemicellulose content in the raw material, a maximum of 87.4% is observed at 185 °C. Recovery yields of 85.2 and 58.5% were obtained at 175 °C and 195 °C, respectively, indicating that the maximum recovery of hemicelluloses is in that range. Therefore, this range of autohydrolysis temperatures was selected for the design of experiments for the autohydrolysis + ionosolv treatments combination. This response was selected as the main one in the autohydrolysis step since no specific applications are considered in this work for the hemicellulosic fraction, so our goal was to maximize the total sugar recovery.

As for the ionosolv time, the interval was selected according to our previous work with the same ionic liquid and

Table 1 Autohydrolysis liquors composition in g L⁻¹ at different T_{AH}

Compound	165 °C	175 °C	185 °C	195 °C	205 °C
Gluco-oligosaccharides	0.74	0.96	1.37	1.28	1.84
Xylo-oligosaccharides	21.97	26.10	17.99	5.49	1.51
Galacto-oligosaccharides	3.16	2.85	1.86	0.70	0.62
Arabino-oligosaccharides	0.32	0.25	0.14	0.13	0.02
Manno-oligosaccharides	0.38	1.35	1.49	0.62	0.35
Glucose	0.17	0.26	0.46	1.25	2.57
Xylose	0.87	5.01	12.58	14.12	5.41
Galactose	1.18	2.13	2.78	2.62	1.39
Arabinose	0.86	0.87	0.76	0.57	0.25
Mannose	0.41	0.11	0.39	1.13	0.65
Formic acid	0.05	0.15	0.60	0.66	0.78
Acetic acid	0.95	2.09	4.20	6.74	8.70
5-Hydroxymethylfurfural	0.03	0.11	0.27	0.70	1.91
Furfural	0.06	0.36	1.23	3.13	5.12
Hemicelluloses recovery (%) ^a	58.98	85.21	87.45	58.49	23.96

^a Calculated regarding to total hemicelluloses present in eucalyptus wood.



autohydrolyzed poplar wood, a very similar hardwood, establishing an ionosolv treatment time interval of 1–5 h.⁴⁶ The levels of the selected factors of this work are shown in Table 2.

3.2 Influence of T_{AH} and t_{iono} on hemicelluloses, lignin, and cellulose recoveries and cellulose purity

The results of the hemicelluloses recovery (Rec_{HEM}), lignin recovery (Rec_{LIG}), CRF recovery (Rec_{CRF}) and lignin content in CRF (L_{CRF}) are shown in Table 2. It was possible to calculate statistical regressions for each of them. Details of the statistical regressions can be found in ANOVA table in Table S1 in the ESI.† All regressions passed the *F* test.

3.2.1 Hemicelluloses recovery. The first response of the treatments' combination study was the hemicellulose recovery yield (Rec_{HEM}), modeled by eqn (1), with a *R*-square of 0.976. Its regression is depicted in Fig. 1. In this case, it was only affected by the autohydrolysis temperature, being both autohydrolysis temperature and its interaction with itself significant for a 95% confidence interval. This parameter is closely related to autohydrolyzed wood recovery and its composition, which can be found in Table S2 in the ESI.†

$$Rec_{HEM} = 85.64 - 11.93 \cdot T_{AH} - 10.11 \cdot T_{AH}^2 \quad (1)$$

Hemicelluloses recovery yield followed a similar trend as described in the previous section for the recovery of sugars and oligosaccharides with temperature, reaching a maximum at 180 °C with a recovery yield of 89.07%, and then decreasing due to the degradation of sugars into furanic and acidic compounds.^{36,47} The trend observed here is similar to those described by other authors employing hardwoods as raw material at temperatures between 130–30 °C, increasing the concentration of hemicelluloses with the treatment temperature up to a maximum, subsequently decreasing due to their depolymerization and degradation to acids and furanic compounds.^{36,47–49}

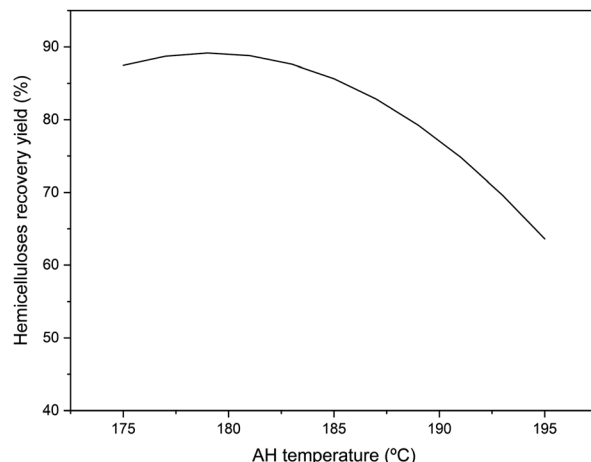


Fig. 1 Representation of statistical adjustment of hemicelluloses recovery as a function of autohydrolysis temperature.

3.2.2 Lignin and cellulose rich fraction recoveries. Lignin recovery (Rec_{LIG}) and cellulose rich fraction recovery (Rec_{CRF}) are influenced by autohydrolysis and ionosolv treatments, making interesting to study the effect of both treatments combined. In this case, a confidence interval of 90% was selected since autohydrolysis temperature was close to be significant with a 95% interval. Thus, the autohydrolysis temperature (T_{AH}) and ionosolv time (t_{iono}) variables are significant with a *P* < 0.1, both with a positive effect on lignin recovery and a general negative effect on cellulose recovery. The statistical regressions are shown in eqn (2) and (3), with *R*-square coefficients of 0.941 and 0.838, respectively, and depicted in Fig. 2.

$$Rec_{LIG} = 5.77 + 2.19 \cdot T_{AH} + 9.18 \cdot t_{iono} \quad (2)$$

$$Rec_{CRF} = 56.79 + 0.85 \cdot T_{AH} - 3.93 \cdot t_{iono} + 3.03 \cdot t_{iono}^2 \quad (3)$$

Table 2 Experimental results of eucalyptus fractions recovery and lignin content in CRF

Factors	Responses					L_{CRF}^d	
	Run no.	T_{AH} (°C)	t_{iono} (h)	Rec_{HEM}^a (%)	Rec_{LIG}^b (%)	Rec_{CRF}^c (%)	
1	175	1		85.21	41.81	58.52	10.00
2	195	1		58.49	54.35	62.84	11.73
3	185	3		87.45	57.51	55.92	9.18
4	175	5		85.21	59.94	54.78	11.34
5	195	5		58.49	64.34	58.30	12.21
6	185	3		87.45	59.01	54.25	10.70
7	170.9	3		84.73	54.08	58.72	11.73
8	199.1	3		52.52	54.50	58.02	11.39
9	185	0.17		87.45	36.16	72.27	18.00
10	185	5.83		87.45	68.20	55.87	13.92
11	185	3		87.45	55.32	58.62	10.03

^a g recovered hemicelluloses/100 g of hemicelluloses in eucalyptus. ^b g lignin/100 g lignin in autohydrolyzed eucalyptus. ^c g CRF/100 g of autohydrolyzed eucalyptus. ^d g lignin/100 g CRF, calculated from NREL/TP-510-42618 procedure.



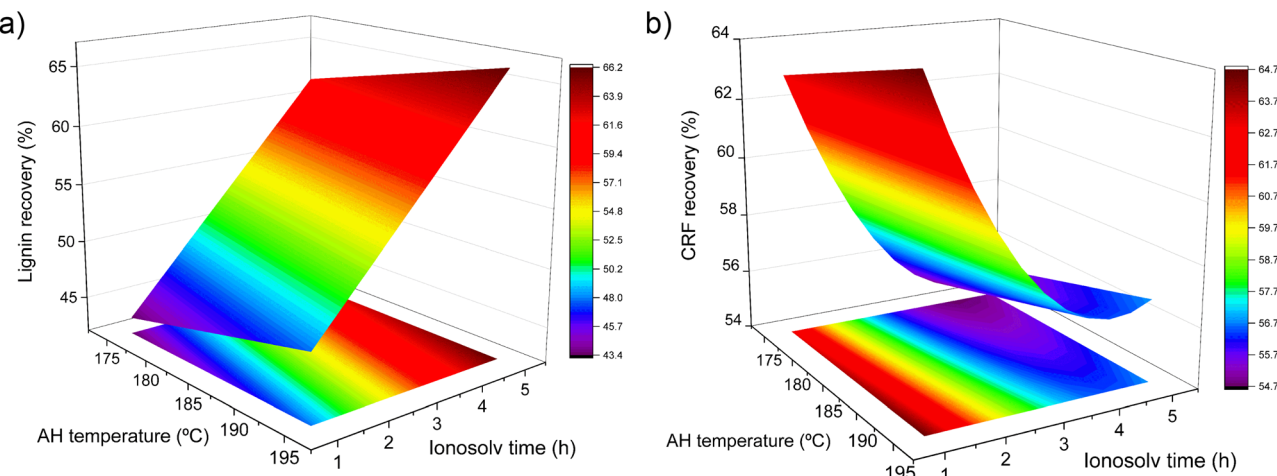


Fig. 2 Response surface plots of lignin (a) and CRF (b) recoveries.

In the regression eqn (3) and in Fig. 2a, it is observed that the ionosolv treatment time has a greater influence than the autohydrolysis temperature on lignin recovery. Thus, for an ionosolv time of 1 h, approximately 40% of lignin is recovered, compared to 5 h, where approximately 60% is recovered, for an autohydrolysis temperature of 175 °C, whereas approximately 45% and 65% of lignin is recovered at 1 and 5 h, respectively, for an autohydrolysis temperature of 195 °C. Lignin recovery always increases with increasing ionosolv time, without an observed maximum. We have observed in a previous work, with the same ionic liquid and eucalyptus as feedstock, that greatly increasing ionosolv time leads to a decrease in lignin recovery.⁴¹ This is caused by the formation of different degradation products that can act as inhibitors and possible degradation of the ionic liquid.⁵⁰ However, in this case, there is not a maximum in this response, which indicates that either the operating conditions are not severe enough to decrease lignin recovery or some of those degradation products are precipitating with the lignin fraction. This could not be verified by the compositional analysis since these degradation products are measured as Klason lignin by the NREL/TP-510-42618 procedure.

The autohydrolysis temperature presents a low influence on CRF recovery (Fig. 2b), although a slight slope is observed on that axis. With increasing ionosolv treatment time, CRF recovery decreased to a minimum at around 3 h, reaching a value of 54.25% and increasing thereafter. This is due to the precipitation of degradation and condensation products with the CRF under severe treatment conditions. This fact that was already observed by Tu *et al.*⁵¹ As a consequence, cellulose purity decreased from this point, making it more difficult to valorize it, as will be shown below.

3.2.3 Cellulose rich fraction purity. No sugars were found in the lignin fraction, with purities always higher than 97%, so the study of this response is not interesting from a process optimization point of view. These high purities were already reported with the same ionic liquid.^{29,46} However, lignin content in the CRF (L_{CRF}) varied with the treatment operating conditions. In this sense, it is an important parameter to study in

order to valorize the cellulosic fraction. In this case, it was only affected by ionosolv time for a 90% confidence interval. Eqn (4) shows the statistical regression of the data in Table 2, with a R^2 square coefficient of 0.822. The representation of the adjustment is depicted in Fig. 3.

$$L_{\text{CRF}} = 10.17 - 0.17 \cdot t_{\text{iono}} + 2.31 \cdot t_{\text{iono}}^2 \quad (4)$$

A minimum lignin content was observed around 3 h of ionosolv treatment, a condition that maximized CRF purity, obtaining an experimental lignin content of 9.18 g lignin/100 g CRF. From this point on, the lignin content increases with ionosolv treatment time, probably due to pseudolignin precipitation with increasing treatment severity.⁵¹ This result justifies the aforementioned fact that CRF recovery increases at higher ionosolv treatment severities as pseudolignin precipitates with CRF, formed by degraded and recondensed products.

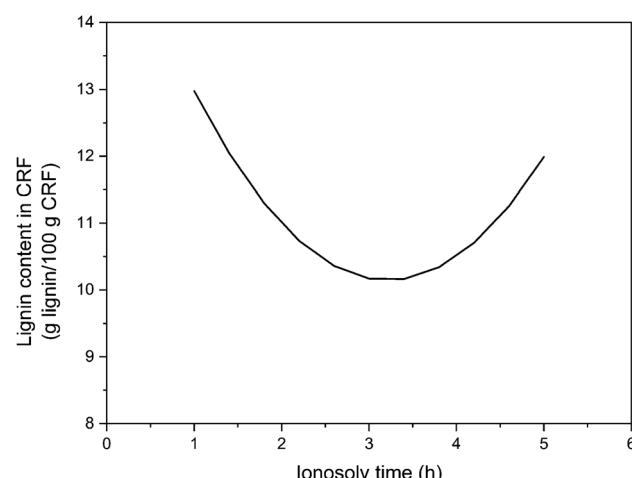


Fig. 3 Representation of statistical adjustment of lignin content in the cellulose rich fraction as a function of ionosolv time.



3.3 Optimization and validation

The process combination proposed in this work is an interesting alternative for wood fractionation, and specially for lignin isolation. Therefore, the operating conditions need to be optimized to maximize lignin recovery and minimize lignin content in CRF. From Fig. 1–3, it was found that lignin recovery could be maximized while maintaining a low lignin content in the CRF at a T_{AH} of 184 °C and t_{iono} of 3.5 h. Under these conditions, the statistical model predicted a 56.85% of lignin recovery, with a 10.19% of lignin in the CRF. In addition, a 55.91% of CRF and an 86.73% of hemicelluloses are predicted to be recovered after the ionosolv treatment and in the autohydrolysis liquid phase, respectively.

To validate the predicted operating conditions, an additional experiment was carried out at a T_{AH} of 184 °C and t_{iono} of 3.5 h. Thus, 89.22% of the hemicelluloses, 54.23% of lignin, and 56.05% of CRF were recovered, with a 10.38% lignin content in the CRF. All these results are within a 5% interval of the predicted values, indicating a good accuracy of the statistical model. The mass balance of the combined fractionation process under these conditions is depicted in Fig. 4.

The raw eucalyptus wood was fractionated into its three main components: hemicelluloses, cellulose, and lignin. The first autohydrolysis step made possible to recover 18.59 g of hemicelluloses/100 g of eucalyptus, with almost no glucan solubilization in this step and negligible lignin losses. 72.43 g of autohydrolyzed eucalyptus/100 g of wood were recovered, with a very low hemicelluloses content. Then, this solid was separated into lignin and a cellulose-rich fraction. 40.60 g of CRF/100 g of eucalyptus were recovered, of which only 4.21 g were lignin, indicating a high glucan purity. 13.68 g lignin/100 g eucalyptus were recovered in another fraction, with no sugars in it. It can be observed that there were still some unrecovered glucan and lignin after the ionosolv step, indicating that they were possibly degraded into furanic and phenolic compounds,

respectively, which could present a high added value. However, these could not be measured because of their low concentrations in the ionic liquid phase.

3.4 Effect of T_{AH} and t_{iono} on lignin properties

The recovered lignins after the combination of AH + ionosolv treatments were characterized to study the influence of autohydrolysis temperature and ionosolv treatment time on molecular weight distribution, thermal stability, and chemical structure. The thermograms and the molecular weight distributions are exhibited in Fig. 5. From them, the mass average molecular weight (M_w), polydispersity index (PDI), and degradation temperature at 10% mass loss ($T_{D_{10\%}}$) were calculated and are shown in Table 3. They were incorporated into the central composite design so the data could be statistically adjusted, and response surfaces represented. The ANOVA table can be found in Table S3 in the ESI.† All regressions passed the *F* test, and a 95% confidence interval was selected, so effects with a $P > 0.05$ were not considered.

3.4.1 Molecular weight and polydispersity index. Weight-average molecular weight (M_w) was influenced by both autohydrolysis temperature and ionosolv time, whereas polydispersity index (PDI) was only affected by ionosolv time in the studied range for a 95% confidence interval. The adjustment of experimental data for M_w and PDI, according to the regression eqn (5) and (6) (*R*-square coefficients of 0.963 and 0.863, respectively), are shown in Fig. 6.

$$M_w (\text{Da}) = 25\,734 + 15\,109 \cdot t_{iono} + 6469 \cdot T_{AH}^2 + 9183 \cdot t_{iono}^2 - 7110 \cdot T_{AH} \quad (5)$$

$$\text{PDI} = 27.92 + 14.90 \cdot t_{iono} + 8.47 \cdot t_{iono}^2 \quad (6)$$

From the molecular weight distributions (Fig. 5a and b) and the representation of M_w as a function of T_{AH} and t_{iono} (Fig. 6a), it can be observed how the mass average molecular weight increased with ionosolv time, whose effect is more remarkable than the autohydrolysis temperature (eqn (6)). The maximum M_w value of 60 000 Da was obtained at an autohydrolysis temperature of 175 °C and an ionosolv time of 5 h. The effect of autohydrolysis temperature is more significant at ionosolv treatment times shorter than 2 h, indicating a greater dominance of the latter over the M_w at longer times, as was already the case for lignin recovery (Fig. 2a). The increase in molecular weight with ionosolv time was already observed in our previous work with eucalyptus and the same ionic liquid.⁴¹ This fact is due to the recondensation of lignins in acidic medium with increasing treatment severity.⁵² On the other hand, the increase of the average molecular weight with autohydrolysis temperature at short ionosolv times is probably related to the recondensation of lignin on the surface of the autohydrolyzed material at severe autohydrolysis conditions, above 185 °C, so the lignin extracted by the ionic liquid was already recondensed before the ionic liquid acted.⁵³

The PDI value (Fig. 6b) increased with ionosolv treatment time. This fact corroborates the recondensation of lignin with

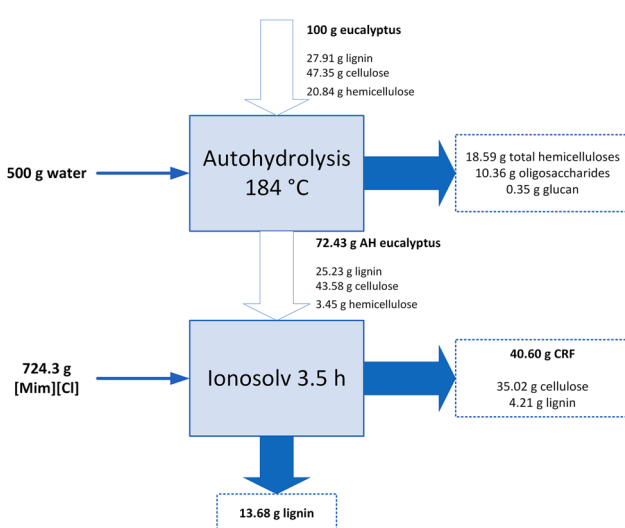


Fig. 4 Mass balance per 100 g of eucalyptus wood at the optimum combined process conditions.



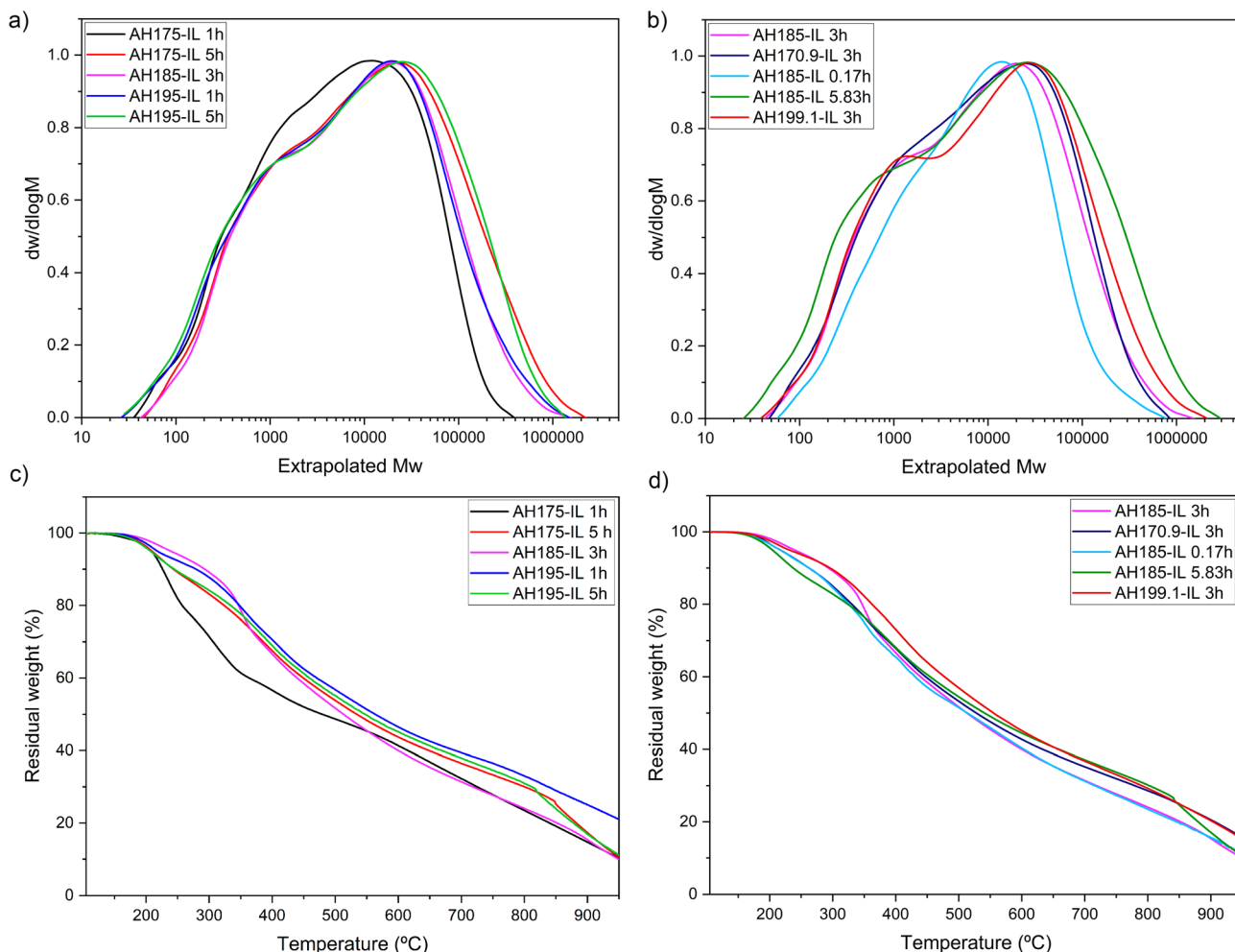


Fig. 5 Representation of (a) and (b) molecular weight distributions (c) and (d) thermograms of the recovered lignins.

Table 3 Experimental results of lignin properties

Factors	Responses					
	Run no.	T _{AH} (°C)	t _{iono} (h)	M _w (Da)	PDI ^a	T _{D10%} (°C)
1	175	1	17 853	18.57	226	
2	195	1	36 055	37.73	280	
3	185	3	22 720	24.90	293	
4	175	5	56 714	42.16	243	
5	195	5	46 473	46.65	245	
6	185	3	28 748	26.75	294	
7	170.9	3	31 854	25.79	262	
8	199.1	3	49 719	34.46	296	
9	185	0.17	20 902	14.33	262	
10	185	5.83	71 527	75.62	237	
11	185	3	26 058	26.32	293	

^a M_w/M_n.

increasing ionosolv treatment time, which would cause the lignins recovered under these severe conditions (times longer than 3 h) to present a heterogeneous structure, probably making their valorization difficult. The recondensation of lignin

in acidic media is associated to the breaking of C–O bonds and the formation of C–C bonds, resulting in more thermally and chemically stable lignins.^{52,54} It is striking that PDI does not depend on the autohydrolysis temperature when M_w does. This fact reflects that the autohydrolysis treatment modifies the M_n value in the same way it modifies the M_w and thus not having an influence on PDI.

Comparing the molecular weights and polydispersity indexes obtained by combining the autohydrolysis + ionosolv treatments with eucalyptus wood with those obtained with the ionosolv treatment alone using the same ionic liquid and eucalyptus wood, it can be observed that the inclusion of a previous autohydrolysis stage produces lignins with similar average molecular weights, between 10 000 and 100 000 Da in both cases, but higher PDI values (2–18 without autohydrolysis step vs. 18–75 with it).⁴¹ This has also been observed with different ionic liquids and raw materials. For example, the employment of protic ionic liquids with different chain lengths with anions derived from organic acids and sugarcane bagasse lead to the recovery of lignins with similar molecular weights (5000–20 000 Da) but with much lower PDI (7.1 as maximum).⁵⁵ Similar results were observed with an hydrogensulfate-based



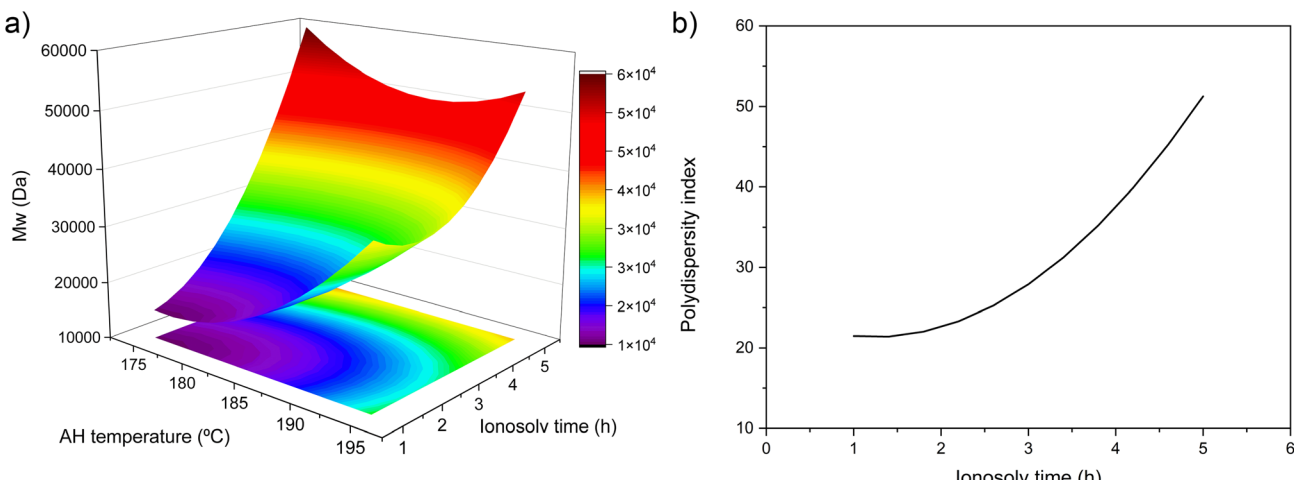


Fig. 6 (a) Response surface plot for M_w and (b) representation of statistical adjustment polydispersity index as a function of ionosolv time.

ionic liquids and miscanthus, recovering lignins with similar M_w but lower PDI (maximum of 4).³⁰ This indicates that the autohydrolysis stage is favoring the deposition of recondensed and heterogeneous lignin particles on the surface of eucalyptus wood, leading to the recovery of lignins with very high PDI, even in lignins with low molecular weights (around 10 000–30 000 Da), hindering their utilization as a base chemical compound and directing their utilization to the reinforcement of materials, or dispersant.^{15,16,18,56}

3.4.2 Thermal stability. Degradation temperature at 10% mass loss ($T_{D_{10\%}}$) from the thermograms (Fig. 5c and d) was selected to represent thermal stability of the extracted lignins. Both autohydrolysis temperature and ionosolv time had a significant influence on the $T_{D_{10\%}}$ for a 95% confidence interval. Its response surface is depicted in Fig. 7, according to eqn (7), with a R -square coefficient of 0.975.

$$T_{D_{10\%}} (\text{°C}) = 293 + 13.01 \cdot T_{\text{AH}} - 6.67 \cdot t_{\text{iono}} - 10.94 \cdot T_{\text{AH}}^2 - 25.69 \cdot t_{\text{iono}}^2 - 13 \cdot T_{\text{AH}} \cdot t_{\text{iono}} \quad (7)$$

The degradation temperature at 10% mass loss of lignins (Fig. 7) increased with autohydrolysis temperature and ionosolv time up to a maximum of 297 °C at an autohydrolysis temperature of 190 °C and an ionosolv time of 2 h. $T_{D_{10\%}}$ does not decrease at autohydrolysis temperatures higher than 190 °C, which indicates that the autohydrolysis step is causing the deposition of recondensed lignins in the material surface, as it was stated before, a fact that is associated with a decrease in the relative content of C–O bonds and an increase in C–C bonds, resulting in more thermally stable lignins.⁵⁷ Lignin recondensation was also observed with ionosolv time, causing the aforementioned increase on thermal stability. However, at ionosolv times longer than 3 h, $T_{D_{10\%}}$ decreased for every T_{AH} , a fact that is possibly related to the recovery of pseudolignin and degradation products at long ionosolv times, as commented in Section 3.2.2. It is also important to remark that due to the negligible sugar content in the recovered lignins, the observed differences are due to the lignin structure. The lowest $T_{D_{10\%}}$ obtained was 226 °C at 175 °C and 1 h. This value is higher than that obtained for lignins from Kraft and organosolv processes (around 220 °C) with poplar wood, indicating that the combination of autohydrolysis + ionosolv treatments produces, at least, equally thermally stable lignins in the worst case scenario.⁵⁸ Comparing these lignins to those obtained without a previous autohydrolysis stage and the same ionic liquid with eucalyptus, it is clear that the inclusion of an autohydrolysis step produces lignins with higher thermal stabilities ($T_{D_{10\%}} = 225\text{--}265$ °C vs. 226–293 °C), a fact that remarks their possible utilization as a reinforcement in biomaterials.^{56,59,60}

3.4.3 Chemical structure. The two main interunit linkages content and S/G ratio were selected as the main responses of the HSQC-NMR characterization. In this case, it was not possible to perform statistical regressions of the experimental data since the adjustment was very poor (R -square values below 0.5). Thus, the experimental data are shown in Fig. 8. In addition, the spectra can be found in Fig. S2 in the ESI.†

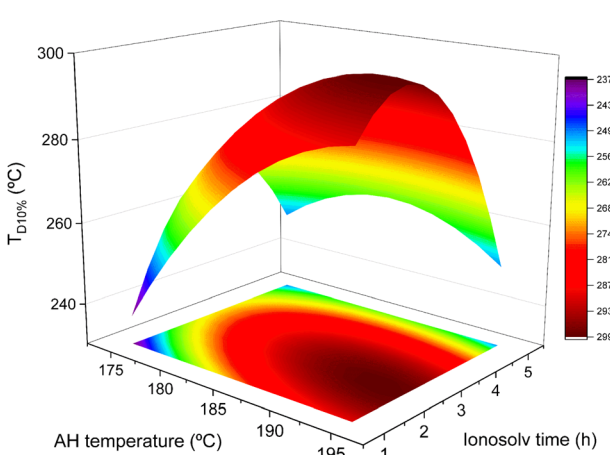


Fig. 7 Response surface plot of $T_{D_{10\%}}$.



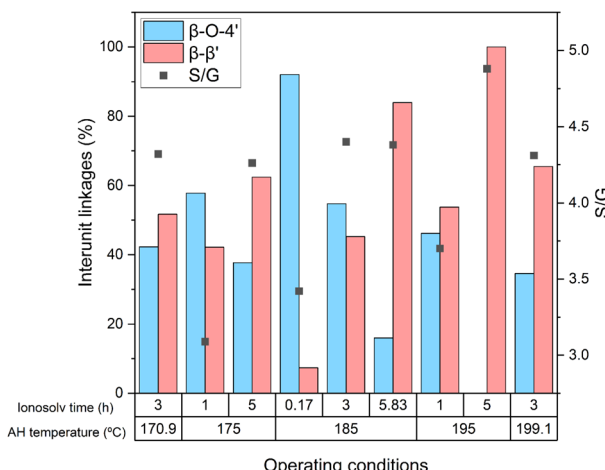


Fig. 8 β -O-4' and β - β' linkages content and S/G ratio of the recovered lignins.

Comparing the results obtained for different ionosolv times at the same autohydrolysis temperature, it can be seen that increasing treatment time produced a β -O-4' linkages degradation with an enrichment in β - β' ones. This increase in resinol substructures supports the previously mentioned lignin recondensation and, consequently, an increase in the molecular weight of lignin and thermal stability. Similar findings have been reported using different imidazolium-based ionic liquids to depolymerize lignin from poplar, eucalyptus, and model compounds.^{41,61,62} Autohydrolysis temperature presented a similar effect at temperatures above 185 °C, recovering lignins with higher β - β' content at higher autohydrolysis temperatures. This effect supports the higher thermal stability of lignins recovered at increasing autohydrolysis temperatures and the augment of M_w due to lignin recondensation in the autohydrolysis step that was anticipated before in Section 3.4.1.

Attending to the relative proportion of S and G units, S units were predominant as it is the main structural unit in hardwoods.^{63,64} The calculated S/G ratios were similar to that of the MWL lignin of the same eucalyptus species (3.7).⁴¹ S/G ratio increased from 1 to 3 h and then, excepting the lignin recovered at an autohydrolysis temperature of 195 °C and ionosolv time of 3 h, it remained practically constant at ionosolv times equal or longer than 3 h for every autohydrolysis temperature. Hence, the autohydrolysis step had no influence on the relative proportion of S and G units in the studied temperature interval. These results suggest a non-preferential action of the ionic liquids towards any of the lignin substructures. However, the degradation of G units from 1 to 3 h can be attributed to the preferential degradation of these units by the action of ionic liquids.^{65,66}

4 Conclusions

The combination of autohydrolysis and ionosolv treatments for eucalyptus fractionation made possible to recover hemi-celluloses, cellulose, and lignin separately. The design of

experiments approach was successful in statistically modeling the fractionation process, which allowed the maximization of lignin recovery (56.85%) and minimization of lignin content in the cellulosic fraction (10.19%) at an autohydrolysis temperature of 184 °C and an ionosolv time of 3.5 h. The prediction validation by an experiment under those operating conditions revealed that the statistical model was successful to describe the fractionation process. Lignin properties were also measured and varied with both treatments operating conditions, having the ionosolv treatment a higher influence on lignin characteristics. It was revealed that increasing treatment severity, especially ionosolv time, caused lignin recondensation and produced more thermally stable lignins. This study enhances the understanding of the effect of autohydrolysis and ionosolv treatments on the overall fractionation process and on lignin characteristics in a treatment combinations approach.

Author contributions

A. O.: investigation, writing-original draft, formal analysis. V. R.: investigation, conceptualization, methodology. J. C. D.: conceptualization, validation, methodology. M. V. A.: writing-review and editing, supervision, project administration. M. O.: writing-review and editing, supervision, project administration. F. R.: project administration, funding acquisition.

Conflicts of interest

There are no conflicts to declare.

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