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Tailoring the Molecular and Thermo-Mechanical Properties of Kraft Lignin by Ultrafiltration

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ABSTRACT: This study has shown that ultrafiltration allows the selective extraction from industrial black liquors of lignin fraction with specific thermo-mechanical properties, which can be matched to the intended end uses. Ultrafiltration resulted in the efficient fractionation of kraft lignin according to its molecular weight, with an accumulation of sulfur-containing compounds in the low-molecular weight fractions. The obtained lignin samples had a varying quantities of functional groups, which correlated with their molecular weight with decreased molecular size, the lignin fractions had a higher amount of phenolic hydroxyl groups and fewer aliphatic hydroxyl groups. Depending on the molecular weight, glass-transition temperatures (T_g) between 70 and 170°C were obtained for lignin samples isolated from the same batch of black liquor, a tendency confirmed by two independent methods, DSC, and dynamic rheology (DMA). The Fox–Flory equation adequately described the relationship between the number average molecular masses (M_n) and T_g 's-irrespective of the method applied. DMA showed that low-molecular-weight lignin exhibits a good flow behavior as well as high-temperature crosslinking capability. Unfractionated and high molecular weight lignin ($M_w > 5$ kDa), on the other hand, do not soften sufficiently and may require additional modifications for use in thermal processings where melt-flow is required as the first step. © 2014 Wiley Periodicals, Inc. *J. Appl. Polym. Sci.* **2014**, *131*, 40799.

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INTRODUCTION

Biorefinery of forest-based products represents an emerging opportunity that may have a dramatic impact on future energy and material supplies, as well as on the forest industry itself.¹ The primary aim of modern integrated forest biorefineries is to produce high-value products, such as chemicals and biopolymers, while improving the efficiency and profitability of their core operations. The three major components of lignocellulosic materials are cellulose, hemicellulose, and lignin. Traditional pulp and paper processes, as well as most of the current biorefinery projects, concentrate on the utilization of cellulose and hemicellulose. Lignin, the second most abundant macromolecule in nature, is usually considered to be a low-value residue and is simply burnt for the production of process steam and energy; for an all-encompassing biorefinery

concept; however, the value-adding utilization of lignin is critical.²

A major concern in utilizing lignin for the production of materials with consistent properties lies in the structural inhomogeneity of lignin itself. Lignin structure and properties vary depending on the wood species, the industrial processes utilized to isolate it, etc.³ Kraft lignin is generated in an amount of ~50 million tons per year as a by-product in the sulfate (kraft) wood-pulping process.⁴ During pulping, around 90–95% of the lignin contained in the wood is degraded and dissolved in the aqueous solution of sodium hydroxide and sodium sulfide together with carbohydrate degradation products forming black liquor. Lignin is degraded into fragments of different molecular weights, which are soluble in alkaline solution.⁵

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Commercially, kraft lignin plays a vital role in pulp and paper manufacturing as a part of the chemical recovery process. The chemical recovery boiler of a kraft pulp mill is its largest investment; however, it is often the bottleneck in production. Lignin accounts for about 35% of the total dry solids of the black liquor and is also the main contributor to its heat value. Since the capacity of the recovery boiler is often limited by the heat load, the throughput of black liquor cannot be increased. In chemical pulp mills, where the recovery boiler is a bottleneck for increased production of fiber, lignin can, however, be extracted and used as external fuel or in more advanced polymeric applications.^{6–10}

Kraft lignin has several characteristic features that distinguish it from native lignin and other technical lignins: it contains a larger amount of phenolic groups due to the extensive cleavage of β -aryl bonds during kraft pulping, and some biphenyl units, as well as other condensed structures as a result of the severe cooking conditions.¹¹ The weight-average molecular weight of kraft lignin obtained from black liquor can vary from about 600 to 180,000 Da.¹² Consequently, the main limitation for the use of kraft lignin in value-added applications is its poor quality with respect to its non-homogeneous molecular weight distribution, the impurities originating from both the wood itself and process elements, and the varying amount of functional groups, resulting in reactivity differences.

Molecular mass is the key parameter affecting the reactivity and thermo-mechanical behavior of lignin, and fractionation is therefore a necessary step in order to obtain lignin with a narrow distribution of properties.^{13–15} Different methods have been suggested to fractionate lignin, including successive extraction with organic solvents,^{12,16} selective precipitation,^{17–19} and ultrafiltration.^{20–25} Toledano et al.²⁴ have shown in their comparative study that ultrafiltration gives the best results, as the lignin obtained is less contaminated with hemicelluloses. Ultrafiltration also allows a more efficient control of the molecular mass distribution of the lignin by the choice of suitable membrane cut-offs. These are important prerequisites with respect to the application of lignin in the polymeric field, and especially for the use of lignin as a precursor for carbon fiber production.^{26–28} It has been stated that ultrafiltration is economically feasible on an industrial scale.²⁹ A further major advantage of this method for the separation and fractionation of kraft lignin is the possibility to withdraw and use the black liquor at any position in the mill process without the need for pH or temperature adjustment.

By systematically characterizing the fractions obtained using the ultrafiltration approach, by investigation of lignin macromolecular properties, determination of key functional groups, as well as thermo-mechanical properties and reactivity, a technological platform for the targeted isolation of lignin fractions exhibiting the tailored properties necessary for specific applications can be developed. To the best of our knowledge, no systematic study has been performed so far to investigate the suitability of the ultrafiltration technique as a means for isolating fractions of technical kraft lignin with controlled thermo-mechanical properties.

In the present study, we have isolated lignin fractions with narrow molecular weight distributions by ultrafiltration of indus-

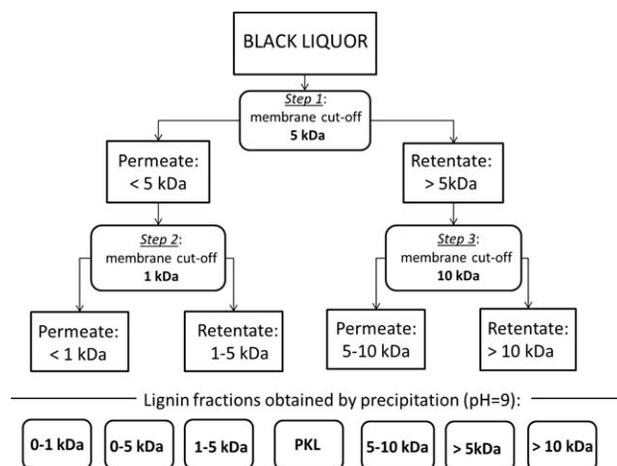


Figure 1. Experimental scheme for the fractionation and isolation of lignin fractions from industrial black liquor by ultrafiltration.

trial softwood weak black liquor, using membranes with molecular weight (MW) cut-offs of 1, 5, and 10 kDa. Detailed investigations of the composition, molecular weight, functional groups, as well as thermal and viscoelastic properties have been carried out for each lignin sample to investigate the efficiency of the ultrafiltration method for the fractionation of kraft lignin and to evaluate the potential of various lignin fractions for advanced polymeric applications, in particular for those where the melt-flow is required as the first step.

MATERIALS

Industrial softwood weak kraft black liquor was kindly supplied by the pulp and paper mill Billerud Gruvön, Sweden. The weak black liquor was taken prior to the evaporators. The pH was >13 and the solids content was ~17.6%. The liquor was used without pH adjustment.

METHODS

Ultrafiltration

Cross-flow filtration (or ultrafiltration, UF) was performed with pilot-scale membrane equipment. The system consists of a 30 L tank equipped with a stirrer, a heating element, a gear pump, and a membrane unit KerasepTM (Novasep, Pompay, France). The membranes used in the study were ceramic membranes made of TiO₂ and ZrO₂ with a surface area of 816 cm², suitable for the filtration of solutions with pH values between 0 and 14 at high pressures and high temperatures. Two membranes with 1 and 5 kDa cut-offs were used. All experiments were carried out with a total recirculation of the black liquor. The temperature range was 40–65°C, the trans-membrane pressure was ~3.5 bar and the pump velocity was 18.3 L/min. For the 10 kDa separation, a bench-scale solvent-resistant stirred cell with working volumes of 10–300 mL and a regenerated cellulose membrane with Mw cut-off 10 kDa from Millipore were used. The pressure was 0.35 MPa and the filtration was run at room temperature.

A starting volume of 14 L of softwood weak black liquor was processed according to the scheme shown in Figure 1. In the *first step*, the pilot system equipped with a ceramic membrane having a 5 kDa cut-off was used for ultra-filtration of the black liquor,

which resulted in a volume reduction (VR) of about 0.7 (i.e., permeate volume divided by initial feed volume) and a volume reduction factor (VRF) of 3.5 (i.e., initial feed volume divided by the retentate volume). In the *second step*, the permeate from the first step was further fractionated using a ceramic membrane with a 1 kDa cut-off, which resulted in a VR of about 0.6 corresponding to a VRF of 2.5. In the *third step*, the retentate from the first step was further fractionated in a stirred cell with a membrane having a 10 kDa cut-off, which resulted in a VR of about 0.7 and a VRF of 3.2. In total, six liquid samples were obtained in addition to the original weak black liquor. All the samples were stored at 4°C prior to lignin isolation.

Lignin Isolation and Purification

Lignin samples were precipitated by acidifying the corresponding liquors to pH 9 with 6 M sulfuric acid. The high molecular weight liquors (>5 kDa) were heated to 70°C. The low molecular weight fractions (<5 kDa) were precipitated following in general the procedure of Helander et al.²⁵ Prior to NMR analysis, lignin samples were further purified by consecutive extraction with toluene and pentane. In total, seven lignin samples were obtained, including a lignin directly precipitated from the original weak black liquor (PKL). The different fractions are referred to as PKL (unfractionated lignin), >10 kDa, >5 kDa, 5–10 kDa, 0–5 kDa, 1–5 kDa, and 0–1 kDa (Figure 1).

In ultrafiltration, the separation of solutes is defined by the molecular weight cut-off of the membrane used. The yield of specific fractions, thus, will depend on the type of the black liquors: on the amounts of high and low molecular weight kraft lignin present in it. Since the yield of the material separated by UF do not correlate with molecular weight cut-offs of membranes, the yields of lignin fractions were not recorded in this study. Instead, focus was placed on the correlation between the MW cut-offs of membranes and the molecular weight of lignin fractions, their composition, structure, and properties.^{7,8,21,23,24}

Chemical Composition of Lignin Samples

The solids content was determined using an infrared dryer balance PM460 (Mettler, Switzerland). The ash content was determined according to the ISO 1762. Elemental analysis (C, H, N, and S) was performed using 1–3 mg samples dried in a vacuum oven at 40°C for 48 h, using a Flash EA 1112, Thermo Finnigan, USA elemental analyzer (external service provided by the Elemental Analysis Unit of the Santiago de Compostela University, USC, Santiago de Compostela, Spain). Klason lignin and carbohydrate analyses were performed according to TAPPI 222 om-02 and SCAN-CM 71:09. The sugar content was determined by an ion exchange chromatography (IC) with a Dionex ICS3000, using a pulsed amperometric detector, a PA1 column (Dionex, Stockholm, Sweden), injection volume 10 µL and a flow rate of 1 mL/min. Carbohydrate standards used for calibration were arabinose, galactose, glucose, xylose, and mannose (Sigma-Aldrich, St. Louis, MO). The acid-soluble lignin content was determined according to Dence,³⁰ where the acid-soluble lignin was analyzed with Shimadzu dual beam UV-2550 UV/VIS spectrophotometer (Kyoto, Japan) using quartz cuvettes. Acid-soluble lignin concentrations were determined using a lignin extinction coefficient (absorptivity) of 128 L/g·cm.³¹

Determination of Molecular Weight by Gel-Permeation Chromatography

Molecular weight properties, such as the weight-average molecular weight (M_w), the number-average molecular weight (M_n), and the polydispersity index (PDI) of the isolated lignin fractions were determined by gel-permeation chromatography (GPC) according to Guerra et al.³² All samples were acetobrominated prior to analysis: a 5 mg sample was stirred for 2 h at room temperature in 1 mL of a 9/1 (v/v) mixture of glacial acetic acid and neat acetyl bromide before surplus reagents were removed rapidly *in vacuo*. The residue was dissolved in 1 mL of HPLC grade tetrahydrofuran (THF), and the resulting solution filtered through a 5 µm syringe filter to remove any eventually remaining solids. A 25 µL aliquot was injected into a 20 µL sample loop. GPC analysis was performed using a Shimadzu instrument consisting of a controller unit (CBM-20A), a pumping unit (LC 20AT), a degasser unit (DGPU-20A3), and a diode array detector (SPD-M20A); the instrumental set-up was controlled using the Shimadzu LC solution software package (Version 1.24 SP1). Two analytical GPC columns (each 7.5 × 30 mm) were connected in series for analyses: Agilent PL-gel 5 µm, 500 Å, followed by Agilent PL-gel 5 µm, 1000 Å. HPLC-grade THF was used as solvent for sample preparation and GPC analyses. The flow rate was kept constant at 0.5 mL/min during the analyses, and the UV signal was recorded for 60 min. Calibration was performed using polystyrene beads with molecular weights of 50,400, 21,000, 7200, 2960, 1270, and 580 Da. The final analysis of each sample was performed using the intensities of the UV signal at 280 nm employing a MS-Excel-based table calculation.

Functional Group Analysis by Quantitative ³¹P NMR

Phosphorus-31 Nuclear Magnetic Resonance (³¹P NMR) analysis with 90° pulse angle, an inverse gated proton decoupling, and a delay time of 10 s was used for the identification and quantification of hydroxyl and carboxyl groups. Prior to analysis, a 20–25 mg sample of the respective lignin fraction was functionalized using 2-chloro-4,4,5,5-tetramethyl-1,3,2-dioxaphospholane in a 1/1.6 mixture (v/v) of CDCl₃ and pyridine for 2 h at room temperature.³³

Thermal Analysis and Dynamic Rheology

Thermo-gravimetric analysis (TGA) was performed using a TA Instruments Q500 TGA. The decomposition temperature (T_d) was determined by heating the samples from 25 to 500°C at a heating rate of 10°C/min under a dry nitrogen (N₂) atmosphere. The T_d of each sample was reported as the temperature at which there was a 5% weight loss. The glass transition temperature (T_g) was determined using a TA Instrument Q1000 DSC at a scan rate of 10°C/min over a temperature range from 0°C to 180–250°C (depending on the specimen). All measurements were made using 2–3 mg samples placed in aluminum hermetic pans under a nitrogen atmosphere. T_g was recorded at the midpoint temperature of the heat capacity transition of the second heating run. The results for each sample were reported as the average of 3 runs.

Viscoelastic responses of selected lignin samples were analyzed using a TA Instruments AR2000 dynamic torsional rheometer

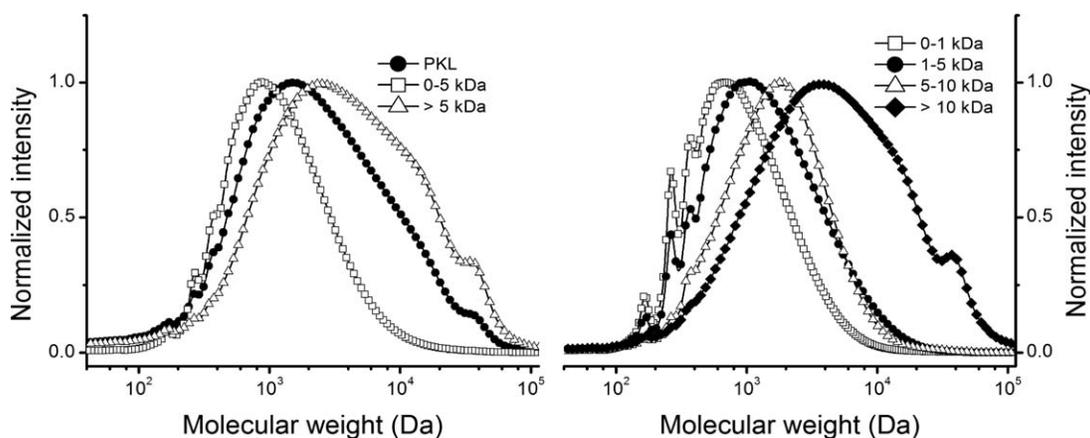


Figure 2. Molecular weight distribution curves obtained by GPC analysis for lignin fractions obtained by ultrafiltration of industrial black liquor. GPC measurements were performed using acetobrominated samples.

in the compression torsion mode. Details of this technique, and its advantages and limitations are described elsewhere.³⁴ About 0.1 g of dry lignin powder was loaded between two parallel plates of 25 mm in diameter and dynamically strained (0.1% strain, 1 Hz frequency) with increasing temperature at a rate of 3°C/min. Temperature-dependent viscoelastic responses were analyzed by sequential heat-cool-heat cycles as following: (1) heating from room temperature to 160–200°C (depending on the specimen), (2) cooling to room temperature, (3) heating to 250°C. All experiments were conducted in a dry N₂ atmosphere. The glass transition temperature was determined as the peak maximum of the loss modulus (G'') for both heating steps.

RESULTS AND DISCUSSIONS

Effect of Ultrafiltration on the Molecular Weight of Lignin Fractions

The efficiency of the ultrafiltration approach for fractionating lignin from the kraft black liquor and obtaining a material with specific characteristics was evaluated by studying the molecular weight properties of lignin fractions.

Over the years, several SEC/GPC procedures were examined and suggested for the determination of the molecular mass of lignin samples of various type and origins.^{35–39} In the recent Round Robin effort to standardize SEC analysis of technical lignins undertaken within the European thematic network “Euro lignin” it was shown that high-molecular-mass fractions of polydisperse lignins are the main source of intra- and interlaboratory variations.⁴⁰ Variations mainly resulted from the gel type used for separation, the eluent, the detection mode and the mode of analyzing. Incomplete solubility of lignin samples prior to GPC analysis was found to be a serious source of error. To reduce the variations, crosslinked polyvinyl styrene columns, THF as the mobile phase, and acetylation as the derivatization method were suggested, and it was recommended that the acetylation should be conducted over a period of 6 days (room temperature) to ensure complete dissolution.⁴⁰ In the present work, in order to investigate the efficiency of the fractionation of lignin samples obtained by ultrafiltration, we followed a GPC analysis using an acetobromination as derivatization procedure as suggested by Guerra et al.³² The main advantage of this method is

the comparably short reaction period, which minimizes any structural alterations, and yields lignin derivatives that are highly soluble in THF.^{41,42} It has been shown that molecular masses observed when the acetobromination derivatization is used, which are comparable to those obtained with acetic anhydride in pyridine derivatization.⁴²

The molecular weight distribution curve (Figure 2) for the 0–1 kD lignin sample had a number of peaks in the low molecular weight (MW) region, as shown in Figure 2b. The same peaks were found in the 1–5 kDa and 0–5 kDa samples, with lower relative intensities. The presence of these similar lignin fragments in both samples, albeit in different quantities, indicates that small amounts of low molecular weight fragments (mono- and dimers) are present in the low molecular weight lignin fractions. A similar phenomenon was observed in the high molecular weight region: the chromatograms for the reference sample PKL and the two high-molecular weight samples, >5 kDa and >10 kDa, all showed a peak at the same position in a very high MW range (at around 50–60 kDa). These peaks may correspond to LCC complexes of very high molecular weight present in the lignin samples. These samples had a higher carbohydrate content, as shown in Table II and discussed below.

The M_w , M_p , and PDI for the acetobrominated lignin samples are shown in Table I and illustrate a quite efficient fractionation of lignin based on the molecular size. There is a clear decrease in weight-average molecular weight with decreasing molecular weight cut-offs of the membranes used for the fractionation. The M_w value seems to be in a good agreement with cut-offs of the membranes. Polydispersities of lignin fractions were reduced as a result of fractionation, indicating that ultrafiltration results in more homogeneous materials with narrower molecular weight distributions.

It is difficult to compare our results with data available in the literature due to the nature of the lignin samples^{3,4} and the uncertainties related to the GPC analysis itself.⁴⁰ For unfractionated softwood kraft lignin, a M_w value of 39,000 was reported earlier.¹³ Toledano et al.²⁴ reported an efficient fractionation of lignin using an ultrafiltration technique with ceramic

Table I. Weight-Average (M_w), Number-Average (M_n) Molecular Weights and Polydispersity Index (PDI) of Different Fractions Obtained by Ultrafiltration^a

Samples	M_w	M_n	PDI
PKL	20,200	5000	4.1
>10	33,500	9500	3.5
>5	28,200	8000	3.5
5–10	4900	2300	2.2
1–5	4700	2000	2.3
0–5	4100	1700	2.4
0–1	2700	1200	2.1

^aThe molecular weights are relative values, related to polystyrene standards, and were obtained from GPC measurements of acetobrominated samples.

membranes of cut-offs 5, 10, and 15 kDa on black liquor from the pulping of *Miscanthus sinensis*. GPC data presented in that paper; however, report quite low M_w values for lignin fractions isolated with membranes having cut-offs of 10 and 15 kDa: 3500 and 6300 Da, respectively. This can be the result of a poor solubility of the lignin samples during the GPC analysis when acetylation was used as derivatization procedure.

The Composition and the Chemical Structure of the Lignin Fractions

Beside the molecular weight characteristics of lignin, its purity and structure, i.e., the presence of reactive functional groups, are key factors determining the utility of lignin for value-added applications. In the present study, we used traditional Klason lignin and carbohydrate analyses, in combination with elemental analysis, to evaluate the purity of the lignin fractions obtained by ultrafiltration and subsequent precipitation with acid. The number of the various functional groups was determined using quantitative ³¹P NMR spectroscopy.

The compositions with respect to carbohydrates, Klason lignin, acid-soluble lignin, and inorganic impurities of the reference (PKL) and the fractionated lignin samples are shown in Table II.

All the samples have a low ash content, which indicates an acceptable acid wash. In general, the lignin materials obtained by ultrafiltration have a high purity with respect to carbohydrates, which is in good agreement with a previous study.²⁴

Table II. Chemical Composition of PKL and Fractionated Lignin Samples

Samples	Carbohydrates (%)	Klason lignin (%)	Ash (%)	Acid-soluble lignin (%)
PKL	1.7	96.9	0.1	2.0
>10	5.2	94.4	0.2	1.0
>5	3.4	94.9	0.5	1.2
5–10	0.3	93.8	n/a	2.9
1–5	0.2	97.8	0.2	3.7
0–5	0.3	94.3	1.1	5.6
0–1	0.1	97.2	2.2	5.7

Only the high-molecular weight fractions, >10 kDa and >5 kDa, contain a noticeable amount of sugars, in the range of 3–5%. It is reasonable to assume that the carbohydrates detected are part of lignin-carbohydrate complexes (LCC). It has previously been reported that most of the lignin present in wood, and especially in the pulp, exists in form of LCC's with various types of hemicelluloses.^{43,44} During the kraft cooking, severely degraded low molecular weight fragments of the lignin macromolecule are dissolved in the pulping liquor.

As the ultra-filtration technique allows the fractionation of dissolved material based on its molecular weight, the fraction of molecular size >10 kDa (for the industrial liquor investigated in this study) contains a large amount of carbohydrates. The 5–10 kDa lignin sample contains only traces of sugars, which means that contamination in the >5 kDa sample was due to the lignin material with a molecular weight greater than 10 kDa (Table I). As expected, low molecular weight lignin fractions contain larger proportions of acid-soluble lignin. This is in the agreement with previous studies.²⁵

The Klason lignin analysis shows that the lignin fractions have high lignin content (Table II). Elemental analysis (Table III) shows; however, that the low MW lignin fractions 0–1 kDa and 0–5 kDa, respectively, contain 23–27% of sulfur, which coprecipitated with the lignin during acidic hydrolysis. It is not clear whether this sulfur comes from the cooking chemicals or was incorporated into the lignin molecules during the kraft cooking. Successive extraction with toluene and pentane removed most of the sulfur, giving lignin materials with sulfur contents in the range between 1.6 and 4.8%; a complete removal of sulfur; however, cannot be achieved in this way. Independent of the washing procedure, there is a trend towards high sulfur contents in lower molecular weight fractions (Table III).

³¹P NMR data (Table IV) show that the unfractionated lignin, PKL, has a high number of phenolic hydroxyl and aliphatic hydroxyl groups, which is a known feature of kraft lignins. A significant number of phenolic groups in technical kraft lignin provides high reactivity and thus makes the kraft lignin an attractive biopolymer for utilization in various applications. Ultrafiltration has shown; however, that kraft lignin in black liquor is nonhomogeneous not only with respect to its molecular size, but also with respect to the quantity of functional groups. With increasing molecular size, the different lignin fractions tend to exhibit an increasing number of aliphatic hydroxyl groups, and a decreasing amount of phenolic hydroxyl groups. The content of carboxylic groups was not affected by the fractionation, i.e., not dependent on the molecular size of the isolated lignin fractions. These observations are in accordance with the mechanism of lignin degradation during kraft cooking. The formation of low molecular weight fractions could be the result of a breakdown of the linkages during kraft cooking; more free phenolic groups—in relation to aliphatic hydroxyl groups—are present in these fractions. This could also be the reaction leading to higher amounts of sulfur with decreasing molecular weight of lignin. The high molecular weight lignin fractions, on the other hand, probably preserved more side chains and β -O-4 linkages, and hence exhibit less free phenolic and comparably

Table III. Elemental Composition of PKL and Fractionated Lignin Samples

Samples	Before extraction (%)				After extraction (%)			
	S	N	C	H	S	N	C	H
PKL	2.5	0.1	64.9	5.5	1.8	0.1	64.7	5.9
>10	1.7	0.1	62.3	5.1	1.6	0.1	62.1	5.9
>5	1.8	0.1	64.9	5.3	1.6	0.1	63.5	5.8
5–10	2.0	0.1	62.6	5.4	1.7	0.1	65.1	6.0
1–5	7.2	0.1	62.1	5.4	3.0	0.1	68.5	6.4
0–5	23.0	0.1	51.6	4.8	4.0	0.1	66.1	6.3
0–1	27.1	0.1	48.9	4.2	4.8	0.1	63.4	5.9

more aliphatic hydroxyls. The higher content of carbohydrates in the high MW fraction may also contribute to the observed increased content of aliphatic hydroxyls. These results are in a good agreement with previous findings based on quantitative HSQC NMR studies of kraft lignin samples obtained by a solubility-based fractionation.⁴⁵

Thermal Behavior of Kraft Lignin Fractions: Role of Molecular Weight and Structure

Thermal analyses such as TGA and differential scanning calorimetry, DSC, are useful methods for studying the physico-chemical properties of polymers. As vaporization, chemical reactions and thermal degradation are accompanied by a weight loss, TGA can be a useful tool for elucidating the physico-chemical properties of lignin. However, for isolated lignins, overall degradation is a very complex process involving the simultaneous formation/degradation of several interunit linkages, and this makes the interpretation of TGA spectra of lignins ambiguous. In the present study, we used the TGA method to compare the thermal stabilities of the lignin fractions. The thermal analysis by TGA reveals significant differences in decomposition temperatures (T_d), defined by 5% of weight-loss of dry samples, between the lignin fractions as shown in Figure 3 and Table V.

The decomposition temperature generally increased with increasing molecular weight, with exception of the 5–10 kDa sample. The decomposition temperature of this sample was 240°C, which is higher than that of the two samples with higher

Table IV. Content of Functional Groups in PKL and Fractionated Lignin Samples

Samples	Aliph-OH (mmol/g)	Ph-OH (mmol/g)	-COOH (mmol/g)
PKL	2.25	3.50	0.36
>10	2.62	2.81	0.52
>5	2.33	2.95	0.49
5–10	1.88	3.87	0.49
1–5	1.37	3.70	0.35
0–5	1.42	3.61	0.46
0–1	1.25	4.11	0.38

molecular weight, i.e., >10 and >5 kDa (Figure 3). The 5–10 kDa sample is the purest lignin sample as it contains less sulfur than the 0–1, 0–5, and 1–5 kDa fractions, and less carbohydrates than the high-molecular weight fraction (>5 kDa, >10 kDa) and the unfractionated lignin. This lignin fraction also has a low polydispersity (PDI 2.2), which is comparable to those of the lower MW fractions. All these factors probably resulted in the observed higher thermal stability.

The T_g of lignin is affected by factors such as molecular weight in the first place, but also by the thermal history, by the presence of low molecular weight contaminants (including water and solvents), and by crosslinking. In conventional DSC studies, the glass transition temperature (T_g) of a polymer is obtained from the second heating scan, while the initial scan (above its T_g) is usually used to eliminate the thermal history stored within the polymer's glassy state.^{46,47} Kraft lignin with its amorphous nature and its propensity for hydrogen bonding interactions usually exhibits a significant enthalpy relaxation resulting in a broad endothermic peak covering its glassy region during the initial heating period in the DSC scan (Figure 4, 1st heat).

To compare the lignin fractions in the present study, heating to 150°C was selected as the annealing temperature for all samples. This choice was based on the fact that this temperature is expected to be above the T_g for most of the lignin samples. The DSC heating curves between the 1st and 2nd heating scans are shown in Figure 4. Heat flow curves from the 1st heating run for most samples have a shape typical for lignin, except for the low MW fractions 0–1 kDa and 0–5 kDa. These two samples exhibit two obvious transition regions below the T_g of lignin as well as an endothermic peak at 117°C. The first transition is predominant and occurs between 50 and 60°C. This transition may be associated with the relaxation of lignin segments that are plasticized by trace amounts of nonbonded water. On the other hand, the smaller transition observed at around 100°C could be associated with changes in lignin molecules caused by evaporation of bonded water.⁴⁸ These two transitions are less obvious in other lignin fractions as they are covered by a broad endothermic peak resulting from the lignin enthalpy relaxation process. The transitions and peaks at 117°C were eliminated as a result of the 1st heating scan. The major difference between these two low MW fractions and the rest of the samples is their significantly higher sulfur content, which leads to the

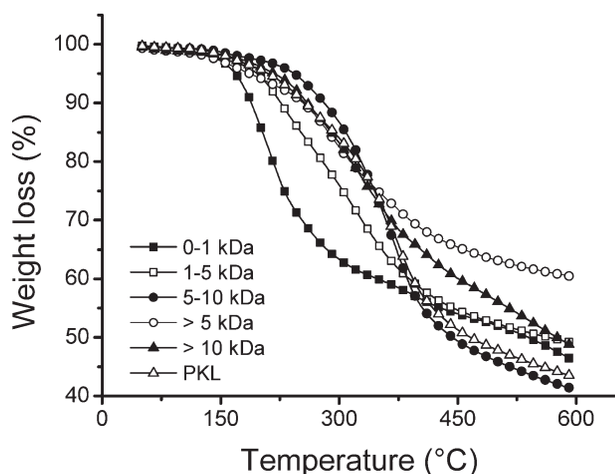


Figure 3. Result of thermo-gravimetric (TGA) analysis of lignin fractions. Decomposition temperature was estimated at 5% of weight loss.

conclusion that the endothermic peaks in the initial DSC scan are probably due to the presence of low molecular weight sulfur-containing compounds in addition to traces of water. The T_g 's of the lignin fractions were determined as the midpoint of the heat transition of the 2nd heating run, after the thermal history of sample had been eliminated (Figure 4 and Table V).

Table V summarizes the decomposition and transition temperature data obtained by the TGA, DSC, and dynamic rheology (DMA) for unfractionated lignin and lignin fractions obtained by the ultrafiltration of kraft black liquor. Three different T_g measurement methods are demonstrated: as the midpoint of the heat capacity transition of the 2nd DSC heating scan, as a peak maximum of G' and as a peak maximum of $\tan \delta$ for both 1st and 2nd heating scans. The glass transition temperatures, T_g , obtained for the fractionated samples lie between 70°C and 170°C (Table V). The reference sample, PKL, had a T_g of 144°C, while the >10 kDa sample had a T_g of 170°C and the 0–1 kDa sample has a T_g of 70°C. The dependence of T_g on the molecular weight has been reported earlier for lignin samples obtained by the solvent fractionation

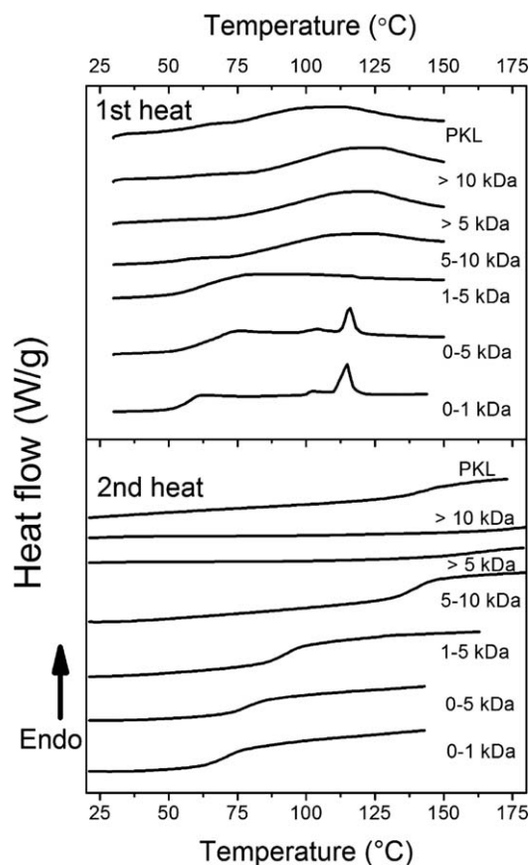


Figure 4. Heat flow curves on the 1st and 2nd heating scans of lignin fractions and PKL lignin measured by differential scanning calorimetry (DSC).

procedure,¹³ also indicating that T_g is highly influenced by the morphology of the polymer. As molecular weight increases, the entanglement, and consequently the rigidity of the polymer increases, resulting in the increase of T_g . This trend can be explained by the well-known Fox-Flory equation,^{49,50} as will be described later in the article. This variation in T_g of kraft lignins isolated from the same black liquor illustrate the potential of ultrafiltration for isolating a specific

Table V. Decomposition and Transition Temperature Data Obtained by the TGA, DSC, and Dynamic Rheology for Unfractionated Lignin and Lignin Fractions Obtained by the Ultrafiltration of Kraft Black Liquor

Samples	TGA T_d (°C)	DSC T_g (°C)	Dynamic rheology			
			Peak maximum loss modulus (G'')		Peak maximum $\tan \delta$	
			T_{g1} , (°C) (1st scan)	T_{g2} (°C) (2nd scan)	T_{g1} , (°C) (1st scan)	T_{g2} (°C) (2nd scan)
PKL	215	144	149	165	183	192
>10	217	170	n/a	n/a	n/a	n/a
>5	190	159	158	185	199	221
5–10	240	140	140	155	194	210
1–5	193	94	93	127	147	179
0–5	175	82	96	114	150	165
0–1	163	70	84	97	135	150

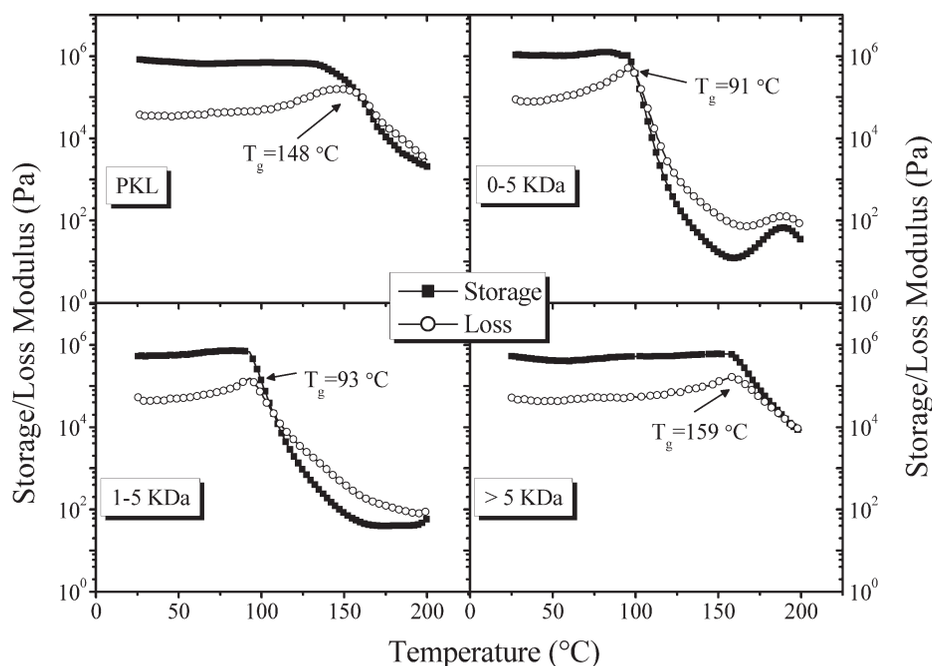


Figure 5. Dynamic rheological response of lignin powder of different molecular weight. First heat responses for unfractionated (PKL) and fractionated lignin (on the basis of MW) are demonstrated.

lignin fraction with the properties desirable for a certain application.

Although there is a clear tendency for T_g to increase with molecular weight, the exact T_g value for each fraction is quite ambiguous. This is due to the fact that nonreversible thermal events, such as condensation reactions, may take place in lignin molecules upon heating above the T_g as a result of breakage of intermolecular hydrogen bonds and increased chain mobility.^{48,51} In their detailed study on the molecular understanding of the way in which the polymeric properties of kraft lignin are affected by various thermal treatments, Cui et al.⁵¹ have shown

that a significant crosslinking through a radical coupling mechanism may take place in lignin molecules when they are heated to temperatures close to their T_g s during the initial DSC scan. This may lead to an increase in molecular weight and polydispersity, resulting in a higher T_g observed in the second DSC scan.

Viscoelastic Behavior and Crosslinking Tendency of Lignin Fractions

DMA analysis is known to be a more adept technique for polymer glass transition analysis. The viscoelastic responses of the dry lignin powders were thus studied using parallel-plate

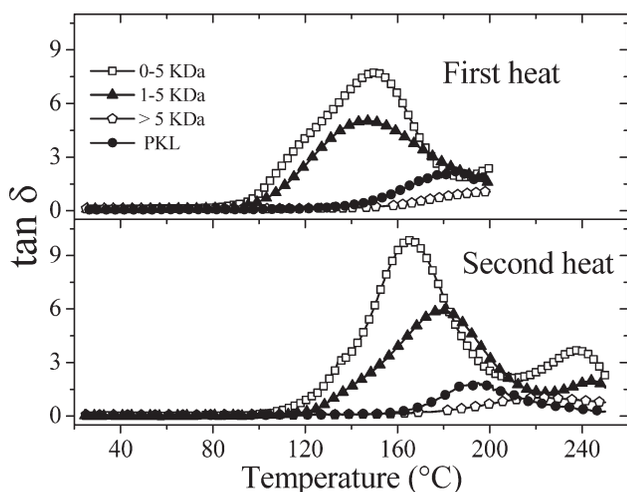


Figure 6. Dynamic rheological response of membrane separated lignin. First and second heat $\tan \delta$ for PKL (reference lignin) and lignin samples with varying molecular weights are shown.

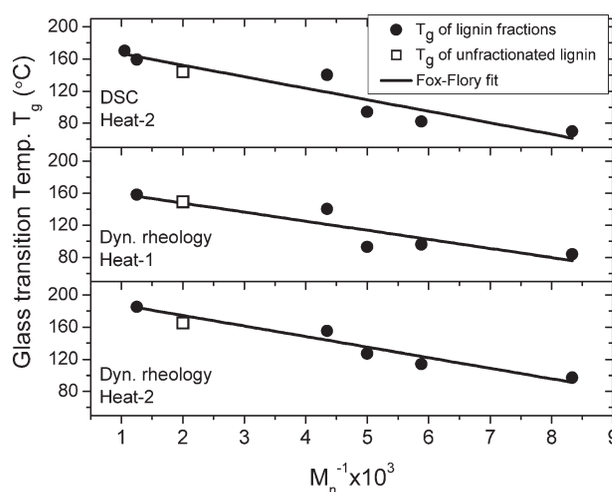


Figure 7. Fox-Flory plot for unfractionated and fractionated lignin. Glass transition temperature versus reciprocal of number average molecular weight are plotted.

Table VI. Estimated Fox–Flory Parameters, Obtained from the Experimental T_g from DSC and Dynamic Rheology

Method	Heating cycle	$T_{g\infty}$ (°C)	k_g
DSC	2nd	181	1.4×10^5
Dynamic rheology	1st	170	1.1×10^5
Dynamic rheology	2nd	201	1.3×10^5

dynamic torsional rheology (Figures 5 and 6). The temperature-dependent evolutions of the storage (G') and the loss (G'') moduli were observed. The molecular weight had a distinct effect on the viscoelasticity of lignin, especially the T_g (identified from the peak of G'') and the degree of softening (identified as the change in G' across the glass transition). With increasing molecular weight, T_g increased—indicating a trend similar to that shown by the DSC analysis (Figure 4 and Table V). Moreover, ^{31}P NMR data (Table IV) show that the higher molecular weight fractions had higher proportions of aliphatic —OH groups are compared to the lower MW fractions: aliphatic —OH groups are more capable of intermolecular hydrogen bonding and generate a greater mobility restriction.^{52,53}

The lignin glass-rubber transition—the change in G' from the glassy to the rubbery phase—was associated with significant softening. Differences in the molecular weight clearly affected the degree of softening also. Unfractionated lignin (PKL) showed a drop in G' of one decade across the glass transition; lower molecular weight fractions softened more (up to a G' drop of 5 decades for 0–5 kDa and 1–5 kDa) and the high molecular weight fraction hardly softened (a G' drop of less than 2 decades for the >5 kDa fraction) and never reached the rubbery plateau within the experimental temperature range. The absence of rubbery plateau for PKL and high MW fractions indicates that the lignin in high MW fractions are more condensed (have a higher effective crosslink density) than in the low MW fraction. This response is, again, rationalized on the basis of more efficient intermolecular hydrogen bonding in the high MW fractions because of the presence of more aliphatic hydroxyl groups in the lignin molecules as well as due to the presence of larger quantity of carbohydrates in them.

A rubbery plateau region is indicative of the effective crosslink density in the polymer.⁵⁴ After the drop that the storage modulus took in the glass transition region, it became almost constant for 1–5 kDa sample within a temperature region between 150 and 200°C. For the 0–5 kDa lignin fraction an additional high-temperature crosslinking reaction was observed during the rubbery plateau phase indicated as a peak in G' at around 180°C. This fraction has a higher amount of sulfur in it than high MW lignin and the 1–5 kDa fraction, which could have some effect on increased reactivity. The loss modulus dominated in the rubbery plateau phase for these samples.

A lower crosslink density, as in case of the low MW fractions 0–5 kDa and 1–5 kDa, leads to a well-defined lower rubbery plateau modulus and a higher flow at high temperature.⁵⁴ This is extremely important in the thermal processing of lignin, where

melt-flow is required as the first step, e.g., fiber spinning. Higher flow is required for optimum melt extrusion and spinning.

Figure 6 shows the $\tan \delta$ of the first and second heat scans of the lignin samples. Although, T_g 's can be obtained from the peak of G'' (Figure 5), the peak of $\tan \delta$ also presents a popular way to determine T_g . The peak $\tan \delta$ -temperature showed a prominent shift to a higher temperatures from the 1st to the 2nd heating scan, presumably due to the temperature-induced crosslinking of the lignin by aryl ether linkage formation between two phenylpropane units.⁴⁸ It is also evident in the $\tan \delta$ profile of 2nd heating scan of the 0–5 kDa sample that there is, again, another minor transition at around 240°C. This is due to additional high-temperature crosslinking reaction. This transition is also weakly visible in the 1–5 kDa specimen. Intriguingly, this high-temperature transition was not prominent in unfractionated and >5 kDa lignin fractions. This perhaps indicates a higher reactivity of the low MW lignin fractions, containing also higher amount of sulfur, than of the higher MW fraction. In unfractionated lignin, the higher molecular weight fraction dominated the viscoelastic responses and the additional high-temperature crosslinking was not visible.

The Fox–Flory equation is one of the most widely used relations to explain polymer glass transition dependence on molecular weight:

$$T_g = T_{g\infty} - \frac{k_g}{M_n}$$

where T_g is the glass transition temperature of any fraction, M_n : the number average molecular weight of that fraction; $T_{g\infty}$: T_g of the material at infinite molecular weight; and k_g : parameter explaining the molecular weight dependence of T_g for the given material.^{50,55,56}

Figure 7 shows the fit for the T_g values obtained from DSC and DMA according to the Fox–Flory equation; delineated Fox–Flory parameters are compiled in Table VI.

Experimental T_g data show a reasonable agreement with the Fox–Flory prediction (Figure 7), exhibiting coefficients of determination of 0.944 for DSC, 0.887 for DMA heat-1, and 0.966 for DMA heat-2. The observed scatter in the data is due to the differences in the chemical nature of the fractions that is not explained by the Fox–Flory relation, which only predicts the effects of molecular weight change on the T_g . The T_g of unfractionated PKL sample lies within the regression in each case. $T_{g\infty}$ and k_g provide an indication of the stiffness of the polymer chains and relates to the effective crosslink density; materials with higher $T_{g\infty}$ provide higher k_g , indicating a stiffer polymeric system. Furthermore, k_g relates to the polarity and the abundance of chain ends; the lower the number of chain ends (i.e., higher crosslink density), the lower is the free volume, resulting in higher polymer stiffness.⁵⁰ The DMA data show this trend. The 2nd heat $T_{g\infty}$, as well as k_g increase as compared to the corresponding 1st heat parameters, reaffirming the increased crosslink density in lignin samples due to exposure to high temperatures.

The thermal analyses by DSC and DMA reveal significant differences in the glass-transition temperatures (T_g) between the lignin

fractions. At the same time, the tendency for T_g increase with increased molecular weight is obvious for each applied method. The shift in T_g to higher values from the 1st to the 2nd heating run, measured from both, the G-peak and the $\tan \delta$ peak, were similar, indicating the validity of the applied detection techniques.

DMA is an efficient tool for evaluating the behavior of lignin samples during various thermal processing reactions, e.g., for screening the spinnability of a lignin sample and predict its spinning temperature, as will be described elsewhere (Chowdhury et al., unpublished results).⁵⁷ It is evident based on the data presented in here, that the 0–5 kDa and 1–5 kDa molecular weight fractions, having good high-temperature flows and cross-linking capabilities, are most probably spinnable and should produce continuous fibers. Unfractionated, PKL, and >5 kDa molecular weight fractions, on the other hand, do not soften sufficiently for continuous spinning, and the addition of plasticizers or a chemical modification of the lignin may be required prior to thermal processing.

CONCLUSIONS

1. Ultrafiltration of industrial black liquor enables the efficient fractionation of kraft lignin in accordance with molecular weight cut-offs of membranes. Only higher MW fractions, e.g., >5 kDa, contain a noticeable amount (3–5%) of sugars. The lower MW fractions have significantly higher contents of sulfur than the high MW fractions.
2. The lignin fractions had different amounts of functional groups, which correlated with their MW—with decreasing molecular size, the lignin fractions tend to contain more phenolic hydroxyl groups, and fewer aliphatic hydroxyl groups. The content of carboxylic groups was not affected by fractionation.
3. The thermo-mechanical properties of lignin fractions showed a good correlation with their molecular weight. By fractionation it may be possible to tailor the T_g of lignin— T_g 's between 70°C and 170°C were obtained. This was confirmed by analyses using both, the DSC and DMA method.
4. The T_g 's obtained from DSC and DMA followed the Fox–Flory equation; the Fox–Flory parameters showed that high temperature exposure increased crosslink density in the lignin samples.
5. DMA method showed that low-molecular weight lignin fractions exhibit good flow behaviors as well as high-temperature crosslinking capability. Experimental data suggest that these fractions are suitable for e.g., melt spinning and should produce continuous fibers as a single component. Unfractionated and high molecular weight fractions (e.g., >5 kDa), on the other hand, do not soften sufficiently for continuous spinning.

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AUTHOR CONTRIBUTIONS

Dr. Sevastyanova O.: Planned the research, performed molecular weight (at URTV, Italy) and thermal analyses TGA, DSC (at UBC, Canada), coordinated the communication between coauthors in Sweden, Canada and Italy and wrote the manuscript. Ms. Helander M.: Helped with ultrafiltration experiment, performed chemical composition and elemental analysis, contributed to the writing of manuscript. Dr. Chowdhury S.: Performed the Dynamic rheology (DMA) analysis, wrote the part of the manuscript dealing with DMA analysis and the application of the Fox–Flory theory to the lignin fractions, contributed to the discussion related to the thermal properties of lignin. Dr. Lange H.: Helped with the molecular weight analysis, contributed to the discussions related to the 31P NMR analysis, participated in the writing of the manuscript. Dr. Wedin H.: Performed the ultrafiltration experiment and the extraction of lignin fractions. Dr. Chang L.: Performed the 31P NMR analysis. Prof. Ek M.: Encouraged to undertake of the present study, participated in the evaluation of results and discussions. Prof. Kadla J.F.: Advised on the studies of the thermal properties of lignin and lignin fractions, participated in the evaluation of results and discussions (hosted Dr. Sevastyanova at his Laboratory of Advanced Biobased Materials at UBC, Vancouver, Canada during July–August 2012. This visit was financially supported by the The Gunnar Sundblad Research Foundation). Prof. Crestini C.: Advised on the studies of the molecular weight and chemical structure of lignin and lignin fractions, participated in the evaluation of the results and discussions (hosted Dr. Sevastyanova at her laboratory at University ‘Tor Vergata’, Rome, Italy during the February 2013. This visit was financially supported by the Cost Action FP1105 Wood CellNet). Prof. Lindström M.E.: Supervised the research, advised on the design of the ultrafiltration experiment and the isolation of lignin fractions, participated in the evaluation of the results and discussions.

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