Selective Acetone-Water Delignification of Eucalyptus urograndis: An Alternative Towards the Biorefinery Approach

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Abstract: Acetone-water (50:50 v/v) delignification of Eucalyptus urograndis (a hybrid of Eucalyptus urophylla and Eucalyptus grandis) was studied at temperatures ranging from 145°C to 195°C. Lignin and polysaccharide extraction kinetics were determined from pulp yield and residual lignin measurements. Arrhenius-type activation energy of initial, bulk and residual delignification phases were estimated respectively as 102.3, 117.7 and 89.0 kJ.mol⁻¹. For polysaccharide removal, the values were 119.4 and 177.9 kJ for slower and faster reactions, respectively.

The selectivity of the acetone-water delignification was approached through the interrelationship between delignification and polysaccharide removal. In contrast to kraft pulping, high selectivity is observed in the beginning of the pulping process and then decreases along the process. The first derivative of the lignin extraction curve presents two maxima, one in the beginning of the extraction and the second one between 65 % and 50 % of pulp yield and can speculating be related to the initial and bulk delignification phases. On the other hand, polysaccharides, probably hemicelluloses, are selectively extracted between 85 and 65 % of pulp yield. The process fits perfectly with the biomass biorefinery approach which consists in selectively separate and add-value to different fractions of wood.

Keywords: Biorefinery, delignification, organosolv, pulping, acetone-water, Eucalyptus urograndis, kinetics, activation energy, selectivity.

INTRODUCTION

The interest in developing ecologically adapted, and both economically and technically feasible wood pulping processes, has increased in the last decades. In this context, organosolv pulping was proposed as a promising alternative for chemical pulp production in the past decades. Several papers have reviewed solvent mixtures, pulp characteristics, as well as the advantages and problems of the processes for pulping purposes [1-7]. In order to point out the economical features of the organosolv processes, Curvelo and co-workers have studied the delignification of lignocellulosic raw materials using low cost solvent mixtures [8-11]. An important regain of interest on organosolv pulping has been observed in the recent years as part of the biorefinery concept due to better selectivity of extraction of the different wood components compared to the traditional pulping liquors [12-16].

The purpose of a delignification process is to remove as much lignin as possible while preserving the wood polysaccharides. However, it is well established that it is practically impossible to remove lignin without some carbohydrate degradation. According to Göring, the extraction of polysaccharides leads to the formation of pores in the cell wall matrix allowing the lignin fragments from the secondary wall and middle lamella to be extracted [17]. Thus, the kinetics of delignification and polysaccharide removal, and selectivity are important parameters for the determination of the best operation and control conditions of a pulping process.

The kraft process presents low selectivity at the beginning and end of cooking because of high polysaccharide dissolution [18]. On the other hand, very good selectivity is observed in the pulp yield range of 75 - 50%. Autocatalysed, organosolv pulping processes tend to be more selective than conventional processes [1], but external addition of acid leads to extended polysaccharide degradation, and consequently, low selectivity [19]. An interesting way of improving the extraction selectivity is by coupling an organic solvent and supercritical carbon dioxide [20,21].

Delignification processes have been reported by several authors to occur in three consecutive phases with different kinetic rates, called initial, bulk and residual phases, for which rate-constants are decreasing in the order mentioned. An excellent review and compilation of kinetics parameters data is given in the handbook by Sixta et al. [22]. The most common approach for the kinetics of isothermal delignification is to consider it occurring as a pseudo, first order process:

\[ \text{Ln}(L) = k \cdot t \]  \hfill (1)

\( L \) being the residual lignin on wood and \( k \) the rate constant.
A most accurate treatment involves a non-linear fitting of a sum of three exponentials related to each delignification phase taking place in parallel [22-24]:

\[
\frac{L(t)}{L_0} = A_x e^{-k_xt} + A_y e^{-k_yt} + A_z e^{-k_zt} \tag{2}
\]

where \(L(t)\) = residual lignin on delignified wood,

\(L_0\) = initial lignin content in wood,

\(A_x, A_y, A_z\) = weight fraction related to phase \(x, y, z\), and

\(k_x, k_y, k_z\) = rate constant of the phase \(x, y, z\).

Several authors have studied the kinetics of polysaccharide extraction during different pulping processes. In acidic media, the most common kinetic model involves two pseudo-first-order steps, one being much faster than the other [19, 26, 27]:

\[
\frac{P(t)}{P_0} = A_x e^{-k_xt} + A_y e^{-k_yt} + R \tag{3}
\]

where \(P(t)\) = residual polysaccharides on delignified wood,

\(P_0\) = initial polysaccharide content in wood,

\(A_x, A_y\) = weight fraction related to step \(x, y\), and

\(k_x, k_y\) = rate constant of the step \(x, y\).

\(R\) = Weight fraction of polysaccharides resistant to solubilization.

The faster step is related to hemicelluloses hydrolysis, particularly to xylanes. Two different explanations for the slower step are found in literature. Conner and Lorenz have attributed the existence of this step to the breakdown of lignin-carbohydrate complexes [26] while Aravamuthan et al. have related it to the hydrolysis of amorphous cellulose [27].

Another important effect taking place during the pulping is the topochemistry of delignification. At the beginning of kraft cooking, the lignin extracted comes almost exclusively from the secondary wall. Removal of the lignin present in the middle lamellae only becomes important when about 40% of the secondary wall lignin has been extracted [28]. In contrast, Pazar and Behera have shown that the extraction of lignin during an organosolv (methanol-water mixture) process occurs from secondary wall and middle lamellae, from the beginning of the process at approximately the same rate [29]. Acetone-water is a very good solvent for lignins, as shown by da Silva Perez et al. [11], Balogh et al. [30], Jimenez et al. [31], Araque et al. [32]. Therefore, great selectivity is expected to occur even from the beginning of the extraction process.

In this work, the kinetics of delignification and polysaccharide extractions, and the selectivity analysis for the organosolv, acetone-water cooking of a hybrid of *Eucalyptus urophylla* and *Eucalyptus grandis*, also called *Eucalyptus urograndis*, are reported. The kinetic parameters for delignification were obtained from both the procedures described by equations (1) and (2), while the polysaccharide extraction parameters were obtained by equation (3). Finally, the process selectivity, calculated as the ratio of extracted lignin over extracted polysaccharides, was studied as an indication for the separation of the different wood fractions in the frame of the biorefinery concept.

**EXPERIMENTAL**

*Eucalyptus urograndis* sawdust (20-60 mesh) was cooked in 120 mL stainless steel reactors heated by an oil bath, with 10:1 liquor - dry wood ratio at 145, 155, 165, 175, 185 and 195°C. Acetone-water mixture (1:1, v/v) was employed as cooking liquor, based on the conclusions of a previous study about the effect of the liquor composition on the selectivity of the process [11]. Once cooked for the given time, the reactor was immersed in a cold water bath to stop the pulping. The reactor was opened and the spent liquor was immediately separated from the pulp by filtering in order to avoid lignin reprecipitation. The pulp was then washed with an acetone-water solution (1:1 v/v) and then with water until neutral pH. Pulp yield and residual lignin content (Klason Lignin – TAPPi T230-om82 method) were determined. Carbohydrate contents were obtained by difference. Correction of the cooking times by H-factor was assumed unnecessary due to the efficient heating system. The chemical composition of the *Eucalyptus urograndis* wood is presented in Table 1.

**Table 1. Composition of Eucalyptus urograndis Wood**

<table>
<thead>
<tr>
<th>Component</th>
<th>%</th>
</tr>
</thead>
<tbody>
<tr>
<td>Extractives (cyclohexane - ethanol)</td>
<td>2.6</td>
</tr>
<tr>
<td>Extractives (water)</td>
<td>1.4</td>
</tr>
<tr>
<td>Klason Lignin</td>
<td>32.1*</td>
</tr>
<tr>
<td>Holocellulose</td>
<td>67.5*</td>
</tr>
<tr>
<td>Ash</td>
<td>0.34*</td>
</tr>
</tbody>
</table>

*extractive-free basis

**RESULTS AND DISCUSSIONS**

1. Delignification

The effect of temperature on the pulp yield is presented in Fig. (1). The pulp yield after 120 min. at 195°C was about 47%, which is comparable to the traditional chemical delignification processes of hardwood species presenting similar
lignin content. Since the pulping process involves both delignification and polysaccharide hydrolysis reactions, which are considerably different, we decided not to perform the kinetics studies (rate constants and activation energy) for the whole pulping process.

Lignin removal percentages are referred here to the initial lignin content. Delignification between 12.1 and 94.6% was observed under our experimental conditions. The delignification at 145°C does not exceed 40% even after 6.5 h of cooking, whereas at 195°C, 80% of the lignin is removed after 1 h and only about 5% of lignin remains in the pulp after 2 h.

The pseudo, first order kinetics behaviour of delignification process (eq. 1) is shown in Fig. (2a) and the values of rate constants obtained from the slope of the straight lines are presented in Table 2. We can identify the existence of bulk phase for all the temperatures studied while the residual phase is observed for temperatures above 165°C. The transition between bulk and residual phases is strongly affected by the cooking temperature. Two concurrent phenomena probably contribute to this behaviour: i) at higher temperatures the breakdown of lignin and lignin-carbohydrates complexes takes place easily; ii) the polysaccharides extraction rate is also heavily affected by the temperature takes leading to the creation of higher pores in the cell wall structure according to the ‘sieve effect’ theory.

The fitting of equation 2 by non-linear regression was also performed and the curves obtained are shown in Fig. (2b). The rate constants obtained from the previous treatment were used as input parameters for the iterative calculation process. Since the initial delignification phase was not identified by the linear treatment, hypothetical values were adopted for the rate constants in order to start the iterative calculation. By using this mathematical treatment, besides the bulk and the residual phases previously identified, the initial delignification phase for temperatures between 145 and 165°C, was also observed. The rate constants and the extracted lignin fractions obtained from the non-linear fitting of equation (2) are presented in Table 3. Excepting for the residual phase, the rate constant values were found to be slightly higher than those found by the linear fitting model.

We observe that most lignin (80 to 90%) is extracted during the bulk phase. During both initial and residual delignification phases the percentage of lignin extraction is no higher than 15%. The lignin extraction is therefore quite different from kraft cooking. Sixta et al. [22] collected literature data on the lignin extracted in each one of the three phases for different wood species. The values show that, for kraft cooking, the bulk phase represents 64 to 78% of the extracted lignin. Although important amount of lignin is extracted during initial phase of kraft cooking (18 to 31%), this fraction is a mixture of lignin and degraded polysaccharides. Indeed, large amount of polysaccharides-like materials

Table 2. Rate Constants and Delignification Ranges for the Pseudo, First Order Kinetics Model (eq. 1) of Acetone-Water Pulping of Eucalyptus urograndis Wood

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>k_b x 10^3 (min^-1)</th>
<th>k_r x 10^3 (min^-1)</th>
<th>Bulk-Residual Phases Transition (% of Delignification)</th>
</tr>
</thead>
<tbody>
<tr>
<td>145</td>
<td>0.97</td>
<td></td>
<td></td>
</tr>
<tr>
<td>155</td>
<td>2.30</td>
<td></td>
<td></td>
</tr>
<tr>
<td>165</td>
<td>5.41</td>
<td>2.68</td>
<td>64.4</td>
</tr>
<tr>
<td>175</td>
<td>11.7</td>
<td>4.56</td>
<td>78.8</td>
</tr>
<tr>
<td>185</td>
<td>21.7</td>
<td>6.90</td>
<td>85.8</td>
</tr>
<tr>
<td>195</td>
<td>34.5</td>
<td>12.8</td>
<td>86.3</td>
</tr>
</tbody>
</table>
is also extracted during this phase. As the separation of the wood components dissolved in the black liquor is one of the major difficulties for the biorefinery approach in kraft pulp mills, a more selective process during initial phase would allow better utilisation and value-adding to these extracted fractions.

Arrhenius’ activation energies were determined for the different phases identified through both linear and non-linear approaches. Fig. (3) shows the variation of rate constants, natural logarithm as a function of the reciprocal cooking temperature. The activation energy calculated from the slope of the straight lines is presented in Table 4. Insignificant differences between the activation energies for the two kinetic treatments were observed, despite the slight differences between the rate constants.

Since the delignification process mainly occurs through the breakdown of α-0-4 and β-0-4 ether bonds, we can relate the activation energies obtained here for the delignification to the values published for the acidic hydrolysis of the lignin model containing these chemical bonds. Meshigini and Sarkanan [34] have reported activation energies from 79 to 102 kJ.mol\(^{-1}\) for the acidic hydrolysis of α-0-4, depending on the substituents, while Sarkanan and Hoo [35] have found values around 150 kJ.mol\(^{-1}\) for β-0-4 lignin model compounds. Based on these values, it is reasonable to suppose that part of α-O-4 linkages is hydrolysed during initial delignification and that the remaining α-0-4 and β-0-4 ether linkages are cleaved in the bulk phase.

The activation energy values obtained for the residual phase remain incomprehensible when related to model lignin hydrolysis because the extension of delignification occurring during this phase is not too small for kinetics analysis.

### Table 3. Rate Constants and Weight Fractions for the Sum of Exponential Kinetics Model (eq.2) of Acetone-Water Pulping of *Eucalyptus urograndis* Wood

<table>
<thead>
<tr>
<th>T (°C)</th>
<th>(k_i.10^3) (min(^{-1}))</th>
<th>(A_i.10^2)</th>
<th>(k_i.10^3) (min(^{-1}))</th>
<th>(A_i.10^2)</th>
<th>(k_f.10^3) (min(^{-1}))</th>
<th>(A_f.10^2)</th>
<th>R</th>
</tr>
</thead>
<tbody>
<tr>
<td>145</td>
<td>2.83</td>
<td>0.085</td>
<td>1.32</td>
<td>0.809</td>
<td></td>
<td></td>
<td>0.106</td>
</tr>
<tr>
<td>155</td>
<td>5.39</td>
<td>0.073</td>
<td>2.35</td>
<td>0.774</td>
<td></td>
<td></td>
<td>0.153</td>
</tr>
<tr>
<td>165</td>
<td>10.8</td>
<td>0.070</td>
<td>5.69</td>
<td>0.796</td>
<td>2.45</td>
<td>0.071</td>
<td>0.063</td>
</tr>
<tr>
<td>175</td>
<td>14.6</td>
<td>0.891</td>
<td>4.81</td>
<td>0.071</td>
<td>0.038</td>
<td></td>
<td></td>
</tr>
<tr>
<td>185</td>
<td>26.5</td>
<td>0.899</td>
<td>6.41</td>
<td>0.074</td>
<td>0.027</td>
<td></td>
<td></td>
</tr>
<tr>
<td>195</td>
<td>40.1</td>
<td>0.900</td>
<td>12.7</td>
<td>0.095</td>
<td>0.005</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

### Table 4. Arrhenius’ Activation Energy for the Acetone-Water Delignification Processes of *Eucalyptus urograndis* Wood

<table>
<thead>
<tr>
<th>Kinetics approach</th>
<th>Initial phase (kJ.mol(^{-1}))</th>
<th>Bulk phase (kJ.mol(^{-1}))</th>
<th>Residual phase (kJ.mol(^{-1}))</th>
</tr>
</thead>
<tbody>
<tr>
<td>Linear model (eq. 1)</td>
<td>118.3±4.7</td>
<td>87.1±5.3</td>
<td></td>
</tr>
<tr>
<td>Non-linear model (eq. 2)</td>
<td>102.3±3.7</td>
<td>117.7±6.1</td>
<td>89.0±9.6</td>
</tr>
</tbody>
</table>
Arrhenius’ activation energies, calculated from slope of the straight lines shown in Fig. (5), are 119.4 ± 7.6 and 176.9 ± 6.3 150 kJ.mol⁻¹ for, respectively, the faster and the slower hydrolysis rates. Polysaccharide removal seems to be much more sensitive to the solvent media than delignification process, especially the rapid hydrolysis. This can be confirmed when the activation energy values obtained here are compared to certain results reported in literature. The activation energy published for the faster hydrolysis kinetics of the acetic acid-water pulping, is in the range 84-88 kJ.mol⁻¹ [19, 25-27]. However, when small amounts of hydrochloric acid are added, the activation energy dramatically drops. For example, when the HCl concentration in the acetic acid-water liquor is 0.014 M or 0.027 M, the activation energies are 39.5 and 44.6 kJ.mol⁻¹, respectively. On the other hand, the activation energy for slower hydrolysis kinetics remains unchanged (around 124 kJ.mol⁻¹). By comparing the values obtained during this work (Table 6) and those mentioned above, it seems that the presence of acetone in the liquor protects the polysaccharides from degrading, since the activation energies are considerably higher. This observation clearly confirms earlier results obtained on the effect of the acetone-water liquor composition on pulping performance, delignification and polysaccharide removal [11]. Eucalyptus urograndis wood was cooked at 185°C for 2 hours in acetone-water liquor containing acetone from 0 to 100% in volume. We observed that the amount of polysaccharides extracted during the cooking trials with liquor, varying from 0 to 30% in acetone, is almost equal at about 30%. However, if the liquor composition used in this work is 50% acetone, the selectivity is increased and the polysaccharide removal is reduced to 21%.

### 3. Selectivity

By plotting both lignin and polysaccharide removal as a function of the pulp yield, a temperature-independent analysis of the process becomes possible, as showed in Fig. (6). The format of the curves suggests there is an interesting correlation between delignification and polysaccharide extraction during the process. Polynomial fitting of the data was used to generate the lines shown in Fig. (6).

In order to evaluate the selectivity during pulping, the ratio of extracted wood component (lignin over polysaccharides) was adopted. Fig. (7) shows the accumulated selectivity of the process as function of pulp yield. The selectivity parameter was found to be very high at the beginning of the processes but quickly descends during the range of 95 - 85% yield, tending to a levelling-off value at around 1.5.

Although accumulated selectivity is important to the control of both pulping conditions and pulp properties, Fig. (7) does not give any idea of the actual selectivity at a given moment of the pulping. This can be found from the first derivative of the fitted curves included in Fig. (6), according to Masura’s procedure [18]. However, as the curves were obtained by data fitting and not by modelling, only a qualitative analysis of the actual lignin and polysaccharide extraction was performed (Fig. 8). The delignification rate is very high from the beginning of the process until 85% yield, it decreases to a minimum at 76% yield, and increases to a maximum centered at 58% yield. The polysaccharide extraction rate is very low in the beginning, passes throughout a maximum at around 74% and then decreases to a minimum at 56% pulp yield. At the end of the pulping, delignification becomes very difficult and the high values of polysaccharide degradation tend to suggest cellulose degradation. The format of the lignin and polysaccharide extraction distribution curves confirms the important degree of correlation existing between the two processes. When the extraction of one component reaches a maximum, the other is at a minimum and vice-versa. The two maxima of the delignification selectivity are present along the cooking process: one from the begin-

<table>
<thead>
<tr>
<th>Temperature (°C)</th>
<th>kₐ,10⁴(min⁻¹)</th>
<th>Aₛ</th>
<th>kₛ,10⁴(min⁻¹)</th>
<th>Aₛ</th>
<th>R</th>
</tr>
</thead>
<tbody>
<tr>
<td>145</td>
<td>1.31</td>
<td>0.252</td>
<td>0.244</td>
<td>0.042</td>
<td>0.705</td>
</tr>
<tr>
<td>155</td>
<td>2.25</td>
<td>0.290</td>
<td>0.952</td>
<td>0.052</td>
<td>0.657</td>
</tr>
<tr>
<td>165</td>
<td>7.54</td>
<td>0.306</td>
<td>2.65</td>
<td>0.065</td>
<td>0.628</td>
</tr>
<tr>
<td>175</td>
<td>16.5</td>
<td>0.315</td>
<td>7.63</td>
<td>0.082</td>
<td>0.602</td>
</tr>
<tr>
<td>185</td>
<td>28.9</td>
<td>0.381</td>
<td></td>
<td></td>
<td>0.619</td>
</tr>
<tr>
<td>195</td>
<td>42.5</td>
<td>0.421</td>
<td></td>
<td></td>
<td>0.579</td>
</tr>
</tbody>
</table>
**Fig. (5).** Arrhenius plot for the rapid (■) and slow (●) rate hydrolysis of polysaccharides during the acetone-water pulping of *Eucalyptus urograndis*.

**Table 6.** Arrhenius’ Activation Energy for the Polysaccharides Removal Using Acetone-Water Pulping of *Eucalyptus urograndis* Wood

<table>
<thead>
<tr>
<th>Kinetics Approach</th>
<th>Fast Hydrolysis (kJ.mol⁻¹)</th>
<th>Slow Hydrolysis (kJ.mol⁻¹)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Non-linear model (eq. 3)</td>
<td>119.4 ± 7.6</td>
<td>176.9 ± 6.3</td>
</tr>
</tbody>
</table>

**Fig. (6).** Temperature-independent profile of lignin (top) and polysaccharide (bottom) extraction during acetone-water pulping of *Eucalyptus urograndis*. Temperature icons: 145°C(●); 155°C(●); 165°C(●); 175°C(●); 185°C(●); 195°C(●).

The behaviour of wood component removal during acetone-water, organosolv pulping is quite different from that observed for alkaline (kraft and soda) delignification as reported by Masura [36,37]. The author showed that serious polysaccharides removal occurs in the beginning of the process (100 - 85% of pulp yield) when the extraction of lignin is almost negligible. However, in the range of 85 - 50% pulp yield, the rate of lignin removal rapidly increases while the polysaccharide extraction rate becomes very low. Lignin selectivity is consequently very low in the range of 100 - 85% yield, followed by a maximum at around approximately 60% and decreases until the end of cooking. The accumulated selectivity is totally different in the two processes. For organosolv pulping, the format of the curve is an exponential decay (Fig. 6), while for the kraft process, a bell-shape curve is observed: the selectivity being very low until 85% pulp yield, suddenly rising to a maximum around 65-70%, and then decreasing again to very low values at around 50%.

Some factors can be invoked here to explain the differences observed between the processes. The first one is the so-called ‘sieve effect’ theory of delignification proposed by Göring and co-workers [17,37,38]. According to the authors, there is a correlation between the molecular weight of extracted lignin fragments and the size of the pores in cell walls.
during the delignification process. Thus, in the beginning of the kraft pulping process, polysaccharides removed from the cell walls enlarge the pores permitting extraction of the lignin fragments. Moreover, it is known that the strongly alkaline kraft liquor easily dissolves part of the hemicelluloses. In neutral (beginning of the acetone-water, organosolv process) and slightly acidic (after part of acetate groups present in the wood have been hydrolysed) media, the dissolution of hemicelluloses is less effective. During organosolv delignification, the better solubility of lignin fragments in the organic solvents, compared to the alkaline liquor, especially at the beginning of the process, might be a major difference. This could explain the high selectivity observed here at the beginning of the process. After this initial behaviour, the removal of polysaccharides is required to let the lignin fragments be extracted as illustrated by the low selectivity between the 80 and 70% yield, followed by the maximum at 58% yield.

The kinetics and selectivity information described here demonstrate the potential of organosolv acetone-water pulping in the frame of a biorefinery approach. Lignin-like fragments can be selectively extracted in the beginning of the pulping process and these fractions can be easily separated from the spent black liquor, purified and used as started material for different applications. Then, during the bulk phase, polysaccharides are mainly extracted and this fraction, after the solvent separation, can be used for the production of biofuel via saccharification and fermentation. The last fractions obtained during the residual phase of delignification being constituted of a mixture of lignin and polysaccharides, including lignin-carbohydrates complexes, would be very difficult to purify and could be preferentially be used to provide energy for the solvent evaporation.

CONCLUSIONS

Organosolv, acetone-water pulping of Eucalyptus urograndis (a hybrid of Eucalyptus urophylla and Eucalyptus grandis) wood was carried out at temperatures ranging from 145°C to 195°C. Studies were performed on kinetics of lignin and polysaccharide removal, and on selectivity.

The pseudo, first-order kinetics of delignification were analysed through two mathematical approaches. First, the rate constants were estimated from linear fitting of natural logarithms as a function of cooking time. Then, the rate constants obtained from the linear fitting were used as initial values in a non-linear fitting using a sum of three exponential terms related to initial (phase I), bulk (phase II) and residual (phase III) delignification. According to this latter procedure, the phase I occurs when cooking is carried out at 145°C, 155°C, and 165°C, while phase II was observed for all the temperatures studied. Phase III was identified only for temperatures above 165°C.

Arrhenius’ activation energies of initial, bulk and residual delignification phases, were estimated respectively at 102.3, 117.7 and 89.0 kJ.mol⁻¹ from the rate constants obtained by the non-linear fitting approach. These values can be related to the cleavage of α-O-4 linkages during the initial phase, and to the remaining α-O-4 and the β-O-4 linkages during in the bulk phase.

Kinetics of polysaccharide removal were studied by using a sum of two exponential equations related to the hydrolysis of wood carbohydrates occurring at two different rates. The faster reaction was observed at all the temperatures studied while the slower is observed for temperatures below 175°C. However, for temperatures above 185°C, both reactions occur with the same rate. Activation energies of 119.4 and 177.9 kJ for slower and faster reactions, respectively, were obtained.

The ratio of extracted lignin over extracted polysaccharide was adopted as a selectivity parameter. We observed that, in contrast to kraft pulping, high selectivity is observed in the beginning of acetone-water pulping and decreases along the pulping process. The overall selectivity at the end of the process was 1.5. In order to get the actual selectivity at a given moment of the cooking process, the first derivative of the delignification and polysaccharide removal curves was obtained. The format of the distribution curves reveals two regions of high selectivity: the first one from the beginning of the process until 85% pulp yield, and the second at 65% to 54% pulp yield. Three different fractions of extracted material can be obtained using this process: the first one rich in lignin at the beginning of the process can be used for the production of high added-value lignin based products, the second one rich in polysaccharides can be proposed for the production of biofuel, and finally the third one for providing energy for the pulping process.

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REFERENCES


Procter AR, Yean WQ, Göring DAI. The topochemistry of delignification in kraft and sulphite pulping of spruce wood. Pulp Pap Mag Can 1967; 68(9): T445-60.


Allgren PA, Yean WQ, Göring DAI. Chlorite delignification of spruce wood - Comparison of molecular weight of the lignin dissolved with the size pores in the cell wall. TAPPI J 1971; 54(5): 736-40.