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X. Pan, Dominique Lachenal, Catherine Lapierre, Bernard B. Monties. Structure and reactivity of spruce mechanical pulp lignins. Part I: bleaching and photoyellowing of in situ lignins. Journal of Wood Chemistry and Technology, 1992, 12 (2), pp.135-147. hal-02709050

HAL Id: hal-02709050 https://hal.inrae.fr/hal-02709050

Submitted on 1 Jun 2020

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STRUCTURE AND REACTIVITY OF SPRUCE MECHANICAL PULP LIGNINS PART I. BLEACHING AND PHOTOYELLOWING OF IN SITU LIGNINS

Xiaoqi Pan and Dominique Lachenal
Ecole Française de Papeterie, Institut National Polytechnique de Grenoble,
Domaine Universitaire, BP 65, 38402 Saint Martin d' Hères Cedex, France

Catherine Lapierre and Bernard Monties
Laboratoire de Chimie Biologique, INRA / CBAI,
Institut National Agronomique Paris-Grignon, 78850 Thiverval-Grignon, France

ABSTRACT

This paper reports the structural investigation of a series of *in situ* spruce lignins in wood, thermomechanical pulp (TMP) and the corresponding bleached (BTMP) and photoyellowed (YBTMP) samples. This was achieved by using an original two-step degradative technique, thioacidolysis followed by Raney nickel desulfuration. The determination of thioacidolysis monomeric and dimeric products allowed an estimate of various lignin building units and interunit bonds, respectively. It was observed that the thermomechanical and bleaching treatments did not affect spruce lignin structure to an appreciable extent. On the contrary, the photoyellowing treatment caused marked structural changes in lignin, particularly when run in severe conditions. These changes were essentially a decrease in the amount of β-O-4 and β-1 interunit bonds and a relative increase in catechol units and in vanillin end-groups.

INTRODUCTION

Interest in high yield pulps, such as thermomechanical pulp (TMP), is growing because these pulps can be produced with a low capital investment and without pollution problems associated with the manufacture of chemical pulps.

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However, their widespread use as substitutes for chemical pulps is hampered by their rapid colour reversion upon light exposure. Although the detailed mechanism of this photoyellowing is not yet clearly understood, a number of studies have shown that lignin is primarily involved in this process, according to the following sequence 1, 2: i) absorption of the near ultraviolet portion of sunlight by photosensitizers like α -carbonyl groups in lignin, 3, 4 and ii) light-induced oxidation leading to coloured products from lignin, 1, 2, 5, 6

The present study, together with forthcoming papers of series, aims at obtaining a better knowledge of the structure of TMP lignins and of their chemical changes during bleaching and photoyellowing processes.

Despite the complexity of a whole lignocellulosic sample, we chose to examine in situ TMP lignins, because isolated lignin preparations, which are often used when soluble samples without analytical interferences from polysaccharides are looked for, may not adequately reflect the behaviour of in situ lignins towards bleaching and photoyellowing. Accordingly, a spruce wood was successively subjected to thermomechanical defibration, bleaching treatment and photoyellowing experiments of variable duration. We thus recovered one TMP sample together with one bleached TMP (BTMP) and three photoyellowed BTMP (YBTMP) samples.

The comparative structural evaluations of the *in situ* spruce TMP, BTMP and YBTMP lignins were achieved by a chemical degradation method thioacidolysis, which leads to lignin depolymerization by selective cleavage of alkyl aryl ether bonds (except methyl aryl ether). The monomeric and dimeric degradation products analyzed are representative of lignin building units and interunit bonds, respectively. 7-10

The thioacidolysis technique first allows an estimate of the amount of the most frequent lignin structures, i.e. arylglycerol- β -aryl ether (β -O-4), through the determination of the main monomeric product 1. As shown in Figure 1, the compound 1 represents the non condensed β -O-4 linked structure in lignin.

In addition, analysis of other thioacidolysis monomeric products makes it possible to quantitate the amount of some minor structural constituents of lignin which may play a key role in the bleaching and photoyellowing processes. These structures are coniferaldehyde, vanillin and coniferyl alcohol end-groups and also the β-O-4 linked catechol units, which lead to the specific thioacidolysis products 2 - 6, respectively, as outlined in Figure 2.

HCH oCH₃

CH₂SEt

CHSEt

C

FIGURE 1. Main monomeric product 1 formed from the thioacidolysis of guaiacyl β-O-4 structures.

FIGURE 2. Monomeric degradation products issued from the thioacidolysis of coniferaldehyde (2), vanillin (3), coniferyl alcohol (4 and 5) end-groups and β -O-4 linked catechol units (6).

Finally, a recent extension of the thioacidolysis method, namely thioacidolysis followed by Raney nickel desulfuration, allows an estimate of the amount of various carbon-carbon and diaryl ether interunit linkages, through the quantitative determination of the dimeric degradation products illustrated in Figure

FIGURE 3. Main dimers recovered from the thioacidolysis, then desulfuration of guaiacyl lignins.

3. These dimers represent so-called 5-5 (compounds 7 and 8), β -5 (compounds 9 and 10), β -1 (compounds 11 and 12), 4-O-5 (compound 13) and tetrahydrofuran ring (THF compound 14) lignin interunit linkages. In the degradative method used herein, dimer 11 may originate from 1,2-diguaiacylpropane structures and / or p,p'-diguaiacylstilbene ones.

RESULTS AND DISCUSSION

The lignin content and the brightness value of the various samples investigated are reported in Table 1.

TABLE 1. Lignin Content and Brightness Value of Spruce Wood and Pulp Samples

	Wood	TMP	BTMP	BTMP YBTMP-1 YBTMP-2 YBTMP-	ҮВТМР-2	YBTMP-
Brightness (% ISO)	n. d.	50	76	59	56	41
Klason lignin (%)	28.5	29.0	28.4	26.2	26.1	24.8
Acid-soluble lignin (%)	0.3	0.4	0.4	0.5	0.8	2.0

n.d.: not determined

It can be seen that the lignin content is not affected by the thermomechanical and bleaching treatments, indicating that lignin removal is not significant in these processes. The photodegradation does not cause any substantial variations in this lignin content either. Only the more severe yellowing experiment (YBTMP-3) induces a small decrease in Klason lignin content and simultaneously a slight increase in the acid-soluble lignin fraction. This result supports the hypothesis that UV irradiation might increase solubility of lignin in sulfuric acid.² This phenomenon could be linked to the formation of small acid-soluble chromophoric fragments from the lignin polymer.

The absolute and relative yields of the thioacidolysis monomers 1 - 6 recovered from the spruce wood meal and pulp samples are reported in Table 2. It can be observed that the number of non condensed β-O-4 structures, essentially reflected by the yield of compound 1, is similar in wood, TMP and BTMP, which indicates that the thermomechanical and bleaching treatments do not affect these prominent structures to any large extent. The only bleaching effect revealed by the data of Table 2 is the decrease in coniferaldehyde end-group and the concomitant increase in vanillin one. According to Gierer ¹¹, alkaline hydrogen peroxide leads to oxidative cleavage of the ring-conjugated double bond of coniferaldehyde. This mechanism may be of significance to the bleaching process since coniferaldehyde is known as one of the major coloured species present in lignin. ^{12, 13}

TABLE 2. Absolute (*) and Relative (**) Yields of Monomeric Products Recovered from the Thioacidolysis of in situ Lignins in Spruce Wood and Various TMP Samples.

Compound	Origin in Lignin	Wood	TMP	ВТМР	YBTMP-1	YBTMP-2	YBTMP-3
* 1	β-O-4 linked guaiacyl unit	1232	1081	1003	1043	859	149
** 2	coniferaldehyde	2	3	1	2	2	3
** 3	vanillin	n.d.	1	3	4	4	14
** 4 and 5	coniferyl alcohol	10	11	13	11	12	8
** 6	catechol unit	n.d.	2	4	3	5	15

^{*} and **: The values are expressed as µmol / g lignin and as relative molar percentages based on compound 1, respectively.

n.d.: not determined because of small amount

one. According to Table 2, main observations are the following: structural changes in lignin, particularly in the case of the more drastic YBTMP-3 On the other hand, the photoyellowing treatments induce more pronounced

- cross-linked structures. This point will also be discussed below. responsible for the reduction in the yield of compound 1 because of formation of lignin condensation reaction upon light exposure may be, to a larger extent, as irradiation time and spectral distribution of the light source. On the other hand as demonstrated by the decrease in the yield of compound 1, after a drastic homolytic cleavage of β-O-4 linkages and the formation of phenoxy and phenacy. The extent of this lignin degradation is dependent on the treatment conditions such radicals which could be further degraded or condensed to coloured photoproducts. yellowing treatment. Based on literature data, 14 - 16 this could involve the i) The number of non condensed β -O-4 structures is considerably reduced
- end-groups could be explained by the mechanism of ring-conjugated double bond cleavage proposed by Gellerstedt and Pettersson. 20 consistent with the repeated observations of vanillin and vanillic acid in irradiated lignocellulosic materials. 6, 17 - 19 Besides, the destruction of coniferyl alcohol C_{α} - C_{β} bond oxidative cleavage occurring during lignin photodegradation, which is compounds 3 and 4, 5, respectively. The former change could be related to the alcohol ones slightly decreases, as shown by the relative determinations of iii) The relative amount of the catechol compound 6 increases, especially for ii) The amount of vanillin end-groups increases whereas that of coniferyl
- in lignin photodegradation. 5, 13, 14, 21 23 leads to results that are in agreement with previously reported literature data reported studies which stressed the role of ortho-quinonoid and catechol structures reactions occur during photoyellowing and convert β -O-4 linked guaiacyl units to the corresponding catechol ones. This is also in accordance with previously the more severe YBTMP-3 treatment. This change suggests that demethylation As discussed above, the analysis of the thioacidolysis monomers essentially

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TABLE 3. Total Yield and Relative Molar Percentage of Main Dimeric Products Obtained from the Thioacidolysis of *in situ* Lignins in Spruce Wood and Various

Sample	Total yield	Re	lative mola	Relative molar percentages (%)	:s (%)	
	(μmol/g lignin)	5-5 (7,8)	β-5 (9, 10)	5-5 β-5 β-1 4-0-5 (7,8) (9,10) (11,12) (13)	4-0-5 (13)	TH (14)
Wood	352	30.9	32.3	26.0	6.7	4.2
TMP	380	29.5	34.1	25.6	7.7	3.1
BTMP	337	31.6	33.7	25.1	6.1	3.6
ҮВТМР-1	280	33.1	35.8	19.8	6.6	4.8
YBTMP-2	264	33.4	36.1	18.0	8.9	3.6
ҮВТМР-3	24	36.4	40.5	17.4	5.8	n.d.

n.d.: not determined (too small amount)

cause any significant changes in the lignin polymer. According to model compound carbon-carbon interunit linkages in spruce lignin. The values obtained for the experimental conditions. As stated in Introduction, the β-1 dimer 11 may originate undergo alkali-promoted conversion to the corresponding stilbenes under bleaching experiments, 24, 25 however, the phenolic 1,2-diguaiacylpropane structures may indicates that the thermomechanical treatment and the bleaching process do not wood, TMP and BTMP in situ lignins are very similar. This result once again the formation of p,p'-stilbenes in lignin. method used herein is not able to detect any structural variation corresponding to from 1,2-diguaiacylpropane and / or p,p'-diguaiacylstilbene structures. Hence, the In this Table, it can be seen that the 5-5, β -5 and β -1 bonds are the main

In contrast, the yellowing treatments cause marked changes, summarized as

drastic YBTMP-3 treatment. This markedly low yield is indicative of the high 7 - 14 is reduced after the yellowing experiments, particularly in the case of the i) Similarly to the case of thioacidolysis monomers, the yield of the dimers

> reactions might increase the yields of trimers and other oligomers, not analyzed similar within the whole series of samples studied (Table 1). Condensation cross-linked degree of the YBTMP-3 lignin, since the total lignin contents are contain coloured structures herein, in the thioacidolysis product mixture. Such highly condensed lignins might

experiments performed by Castellan and coworkers. 26, 27 They demonstrated that situ lignins. Actually, this role is indirectly supported by some model compound experimental evidence of the role of β -1 structures in the photodegradation of in β -1 interunit bond. This observation indicates that the β -1 structure is targetted by dimers brought about by the photoyellowing appears to be the relative decrease in may be converted to stilbene intermediates via alkali-promoted reaction. 24, 25 The phenanthrenes in the presence of oxygen, particularly under the influence of β -1 structure was prone to photodegradation leading to coloured products. the photodegradation process. To our knowledge, this is the first direct latter can readily be oxidized to strongly chromophoric stilbenequinones and / or leucochromophoric and chromophoric systems. Upon bleaching, the β -1 structures According to Nimz, 28 this β -1 structure is potential precursors of leading to formation of chromophores. light. 28 Our results suggest that the mechanism proposed might be a possible way ii) The results reported in Table 3 shows that the major modification of the

CONCLUSION

photoyellowing. changes occurring to in situ spruce TMP lignins during bleaching and The results presented in this paper provide evidence about the structural

appreciable extent. chromophoric coniferaldehyde end-groups. This conclusion supports the hypothesis that bleaching can be performed without degrading lignin to an lignin polymer. The main reaction which can be detected is the destruction of the The bleaching treatment does not substantially change the structure of the

of the lignocellulosic samples under conditions simulating solar irradiation. This On the contrary, a marked degradation of lignin occurs upon light exposure

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degradation is demonstrated by the following structural changes: a substantial decrease in the amount of non condensed β-O-4 structures, a higher condensation degree reflected by the low yield of the thioacidolysis monomeric and dimeric degradation products, a relative increase in vanillin end-groups as well as in catechol units and a noticeable relative decrease in β-1 lignin bonding patterns.

All these detailed structural informations have been quantitatively obtained by means of an original two-step degradative technique, thioacidolysis followed by Raney nickel desulfuration, applied to a microanalytical scale amount of spruce wood and TMP samples.

EXPERIMENTAL

1. Pulp Sample Preparation

The TMP samples were prepared from spruce (picea abies) in a Sprout-Waldron pilot installation at the Centre Technique du Papier in Grenoble. The following conditions were used: temperature 120 °C, pressure 1.4 bar, retention time 15 min. The pulps were then refined in a Sprout-Bauer refiner to a freeness of 45 °SR. The pulp brightness was 50 % (ISO).

Bleaching treatments were performed in polyethylene bags under the following conditions: pulp consistency 15 %, H₂O₂ 5 %, NaOH 2.5 %, Na₂SiO₃ (39° Be¹) 3 %, DTPA (commercial solution) 0.4 % (all chemicals being charged on the basis of oven-dried pulp), temperature 70 °C, time 4.5 hours. The brightness of the resulting bleached TMP (BTMP) was 76 % (ISO).

BTMP handsheets (about 50 g/m^2) were irradiated in an Original Hanau suntest apparatus using a Xenon lamp and with or without an UV filter which eliminates the light of wavelength below 290 nm. Three yellowed pulp samples were prepared:

- YBTMP-1: 2 hours irradiation (1 hr. each side), with UV filter, final brightness 59 % (ISO);
- YBTMP-2: 2 hours irradiation (1 hr. each side), without UV filter, final brightness 56 % (ISO);
- YBTMP-3: 24 hours irradiation (12 hrs. each side), with UV filter, final brightness 41 % (ISO).

Brightness measurements were conducted using an Elrepho 2000 instrument at the wavelength of 457 nm. Brightness data reported herein are average values of the measurements performed on both sides of ten different pulp handsheets.

2. Lignin Content Determination

Both acid-insoluble and acid-soluble lignin contents in the wood meal and aforementioned pulp samples were determined according to the TAPPI standard methods (T 222 os-74 and Useful method 250, respectively).

3. Thioacidolysis

The analyses of the monomeric 7,8 and dimeric 9,10 degradation products recovered from the thioacidolysis of the wood meal and various pulp samples were performed as previously described. Standard errors between duplicate experiments were 5% and 10%, for the monomeric and dimeric products, respectively.

ACKNOWLEDGEMENT

Financial support and a Ph D thesis grant to X. P. from the Centre Technique du Papier-Grenoble is gratefully acknowledged.

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