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PULPING REACTIONS OF VINYL ETHERS

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ABSTRACT

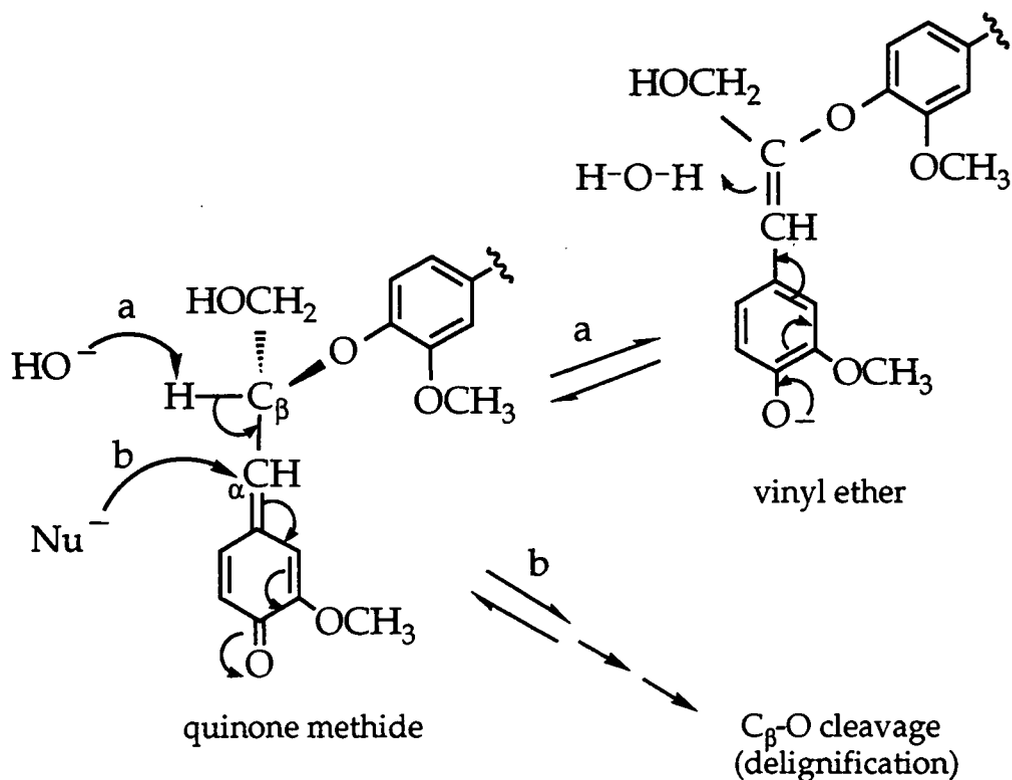
Model studies suggest that vinyl ethers are in equilibrium with quinone methides at 170°C in aqueous NaOH and that 25-30% of the vinyl ethers undergo fragmentation during kraft and AQ pulping.

INTRODUCTION

The delignification of wood chips during alkaline pulping consists of a complex set of reactions.^{1,2} The goal of alkaline pulping is to fragment the lignin into water soluble pieces and separate the aqueous phase from the insoluble carbohydrate fibers. Many undesirable lignin reactions also occur in the process. One such reaction is vinyl ether formation from a quinone methide intermediate (Scheme 1).¹

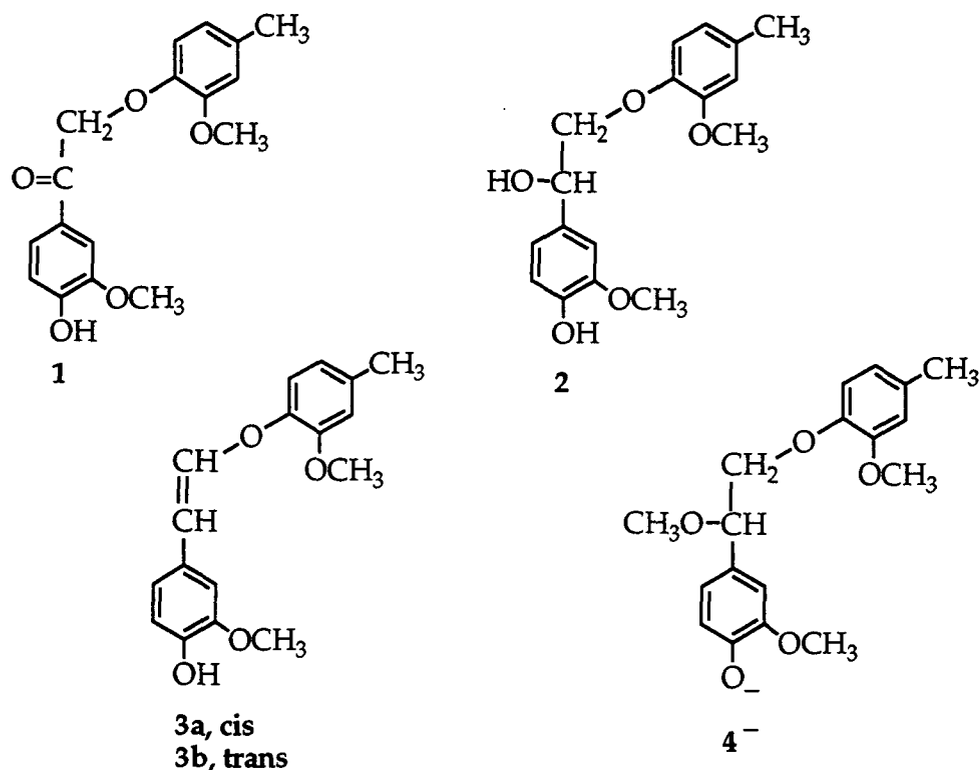
Vinyl ether structures are considered to be relatively stable to alkali and, thus, impede the desirable C_β-aryl ether fragmentation processes.^{1,3} However, previous studies in our laboratory indicated that vinyl ethers are slowly fragmented in ~1M NaOH at 170°C.⁴ The results suggested that vinyl ethers are in equilibrium with quinone methides, as indicated in step "a" of Scheme 1. The following study verifies the reversibility of this step and defines more clearly the reactivity of vinyl ethers under pulping conditions.

Scheme 1



RESULTS AND DISCUSSION

A lignin model vinyl ether was synthesized by (1) coupling α -bromoacetovanillone with sodium creosolate to give ketone dimer **1**, (2) reducing the ketone with sodium borohydride, and (3) treating the resulting alcohol (**2**) with $NaOCH_3$ in methanol at $150^\circ C$ for three hours. The procedure provides a relatively clean ~2:1 mixture of cis/trans isomers **3a/3b**. A key to the synthesis is that the quinone methide from **2** probably has only two reaction pathways available: (1) it can form a vinyl ether by C_β -proton abstraction by CH_3O^- (step a, Scheme 1) or (2) reversibly capture CH_3O^- to give **4⁻**; the α - OCH_3 of **4⁻** cannot be ionized and, therefore, cannot participate in a neighboring group displacement which would break the adjacent C_β -aryl ether bond.¹ A creosol "B" ring was chosen to differentiate this ring from the guaiacol "A" ring, since both A and B ring phenols can be generated under pulping conditions.²



Treatment of the vinyl ether mixture **3a/3b** with different pulping chemicals (NaOH, NaSH, AQ, glucose, and AQ/glucose) at 150°C for 15-120 minutes produced no creosol fragmentation product. This low reactivity is in contrast to compounds similar to **2**, which give 80+% yields of B ring phenols in 30 minutes at 150°C.² However, the vinyl ethers displayed some reactivity when the temperature was raised to 170°C (Table 1).

Table 1. Alkaline degradations of vinyl ether mixture **3a/3b** at 170°C.

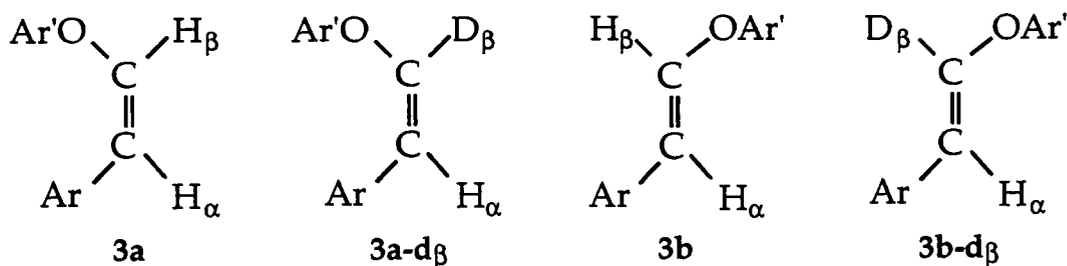
Additive	% Creosol ^a			
	Set 1- 60 min	Set 1- 120 min	Set 2- 60 min	Set 2- 120 min ^b
none	11.6	12.5	15.9	14.0, 16.0
AQ	22.3	---- ^c	18.9	23.5, ---- ^c
AQ/glucose	19.4	26.8	21.3	29.3, 29.6 ^d
NaSH	---- ^c	24.6	18.8	25.3, 26.6

^adifferent sets refer to different experiments; all employed 100 equiv. of NaOH and 5 equiv. of additive; ^bduplicate determinations; ^cvessel leaked; ^dglucose (5 equiv.) in NaOH gave 17.2% creosol after 120 min.

A relative comparison of the data sets in Table 1 shows fairly close agreement, meaning the temperature was reasonably constant between different days. The mixture of glucose and anthraquinone (AQ) generates anthrahydroquinone (AHQ).² Carbohydrates by themselves can promote lignin model fragmentation;⁵ however, this was not the case for the vinyl ether (Table 1, footnote "d" and other unreported data). Glucose is rapidly degraded in NaOH at 170°C⁶ and, thus, may not be present to any appreciable extent during this slow developing reaction.

The data indicate that good pulping additives promote fragmentation of the model. They probably accomplish this by reacting with the quinone methide, which is equilibrium with the vinyl ether, and promoting fragmentation by addition to C_α and then a neighboring group displacement mechanism¹ or by an electron transfer mechanism in the case of AHQ.⁷ The better additives are about twice as effective as NaOH in promoting vinyl ether fragmentation.

To further demonstrate the existence of a quinone methide/vinyl ether equilibrium, we heated the vinyl ether mixture **3a/3b** at 170°C for 4 hr with 0.86M NaOD in D₂O and analyzed the resulting product by ¹H-NMR. The product mixture was composed mainly of vinyl ethers of the types shown below, with a ratio of **3a/3a-d_β** and **3b/3b-d_β** of about 60/40.



The partial exchange of the β-proton in **3a** and **3b** was indicated by a reduced signal area for H_β and the appearance of the H_α signal - a singlet, sandwiched between a doublet. The H_α doublet is due to coupling with a neighboring β-proton; the H_α singlet indicates the absence of a β-proton in the deuterium exchanged compound. The α- and β-cis and trans proton assignments were based on expected H_α-H_β coupling constants and chemical shifts values for previously studied vinyl ethers.⁸

The incorporation of deuterium at the C_β-position is explained by the transfer of a deuterium from D₂O to C_β when the vinyl ether is con-

verted to the quinone methide (the reverse of step "a" in Scheme 1) and then removal of the H from the $-C_{\beta}HD-$ unit of the quinone methide during its conversion back to a vinyl ether (step "a" forward in Scheme 1).

Additional support for the enhanced reactivity of vinyl ethers in the presence of AHQ comes from research results of Hise and co-workers.⁹ The ratio of two lignin fragmentation products, vinyl guaiacol to coniferyl alcohol, changes with different cooking stages in the AQ pulping of pine; also, the absolute levels of vinyl guaiacol are higher in AQ pulping than in kraft and soda pulping. These results have been interpreted to mean that formaldehyde is lost from C_{γ} -units of lignin quinone methides to give vinyl ethers, which upon hydration and reaction with AHQ produce vinyl guaiacol. Coniferyl alcohol is a fragmentation product of a β -aryl ether phenolic lignin end unit which has not lost formaldehyde.

In a very brief report, Yaguchi and co-workers observed a low yield of β -aryl ether fragmentation products after heating a vinyl ether lignin model at 160°C for 1 hour in the presence of THAQ, a reduced form of AQ.¹⁰

CONCLUSIONS

The greater amount of fragmentation with additives AHQ and NaSH (as compared to NaOH alone), together with the deuterium exchange results, indicate that vinyl ethers are in equilibrium with quinone methides at 170°C in aqueous NaOH. Based on the vinyl ether model results, we predict that about 30% of the lignin vinyl ethers produced during kraft and AQ pulping will fragment.

EXPERIMENTAL

Materials. Oxygen-free water was prepared by boiling distilled water for 30 min, bubbling nitrogen in the water as it cooled, and sealing the container until needed. Oxygen-free NaOH solutions were prepared in a similar manner, using O_2 -free water for dilution to the right molarity.

β -(2'-Methoxy-4'-methylphenoxy)-4-hydroxy-3-methoxyacetophenone (1). The procedure for coupling 25.2g of β -bromo-4-hydroxy-3-methoxyacetophenone with eight equivalents of the sodium salt of 2-

methoxy-4-methylphenol was identical to previously described couplings of this type in our laboratory.¹¹ The product was separated by chromatography¹¹ and crystallized from toluene to give 5.8 g of **1**; m.p. 109-111°C; IR (mull) cm^{-1} 3300 (OH), 1670 (C=O), and 1590 (aryl); $^1\text{H-NMR}$ (CDCl_3) δ 2.28 (s, 3, ArCH_3), 3.85 (s, 3, OCH_3), 3.91 (s, 3, OCH_3), 5.23 (s, 2, CH_2), 6.2 (br s 1, OH), 6.6-7.0 (m, 4, ArH) and 7.3-7.6 (m, 2, C_2 and C_6 -ArH); $^{13}\text{C-NMR}$ (d_6 -acetone) ppm 21.0 (q, ArCH_3), 56.1, 56.2 (q, OCH_3), 72.5 (t, CH_2), 111.5, 114.2, 115.2, 115.5, 121.3, 123.7 (d, $\underline{\text{Ar-H}}$), 128.0, 131.9, 146.5, 148.0, 150.2, 152.3 (s, $\underline{\text{Ar-R}}$ and $\underline{\text{Ar-O}}$) and 193.3 (s, C=O); MS m/e (%) 302 (M, 30), 151 (100), 137 (18), 91 (11), and 77 (10).

1-(4'-Hydroxy-3'-methoxyphenyl)-2-(2''-methoxy-4''-methylphenoxy)-1-ethanol (2). To a stirred slurry of 2.0g (6.6 mmoles) of β -(2'-methoxy-4'-methylphenoxy)-4-hydroxy-3-methoxyacetophenone (**1**) in 100 mL of ethanol was added dropwise 2.5g (66 mmoles) of sodium borohydride dissolved in 100 mL of water. After stirring at room temperature for 2 hr, the reaction mixture was neutralized with 3M HCl and extracted three times with CHCl_3 . The combined CHCl_3 extracts were dried (anh. Na_2SO_4) and evaporated. The residue was crystallized from toluene to give colorless crystals, mp 96-99° (one experiment), 86.5-88.5° (another experiment); both sets of crystals had identical spectra: IR (mull) cm^{-1} 3100-3600 (OH), 1600 (aryl); $^1\text{H-NMR}$ (CDCl_3) δ 2.30 (s, 3, ArCH_3), 3.65 (br s, 1, OH), 3.84 (s, 3, OCH_3), 3.86 (s, 3, OCH_3), 3.85 (d of d, $J_{\text{AX}} = 9.3$ and $J_{\text{AB}} = 10.0\text{Hz}$, 1, $\text{ArCH}_X\text{CH}_A\text{H}_B$), 4.12 (d of d, $J_{\text{AB}} = 10.0$ and $J_{\text{BX}} = 3.2\text{Hz}$, 1, $\text{ArCH}_X\text{CH}_A\text{H}_B$), 4.98 (d of d, $J_{\text{AX}} = 9.3$ and $J_{\text{BX}} = 3.2\text{Hz}$, 1, $\text{ArCH}_X\text{CH}_A\text{H}_B$), 5.73 (s, 1, OH), 6.6-7.0 (m, 6, ArH); $^{13}\text{H-NMR}$ (d_6 -acetone) ppm 20.9 (q, ArCH_3), 56.0 (q, OCH_3), 72.4 (d, $\underline{\text{ArCH}}$), 76.3 (t, $\underline{\text{ArCHCH}_2}$ -), 110.5, 114.1, 115.0, 115.7, 119.5, 121.5 (d, $\underline{\text{Ar-H}}$), 131.5, 133.5 (s, $\underline{\text{Ar-C}}$), 146.3, 146.9, 147.6, and 150.2 (s, $\underline{\text{Ar-O}}$); MS m/e (%) 304 (M,19), 153 (45), 152 (27), 138 (100), 137 (22), 123 (28), and 93 (25).

2-(2'-methoxy-4'-methylphenoxy)-1-(4'-hydroxy-3-methoxyphenyl)-ethene (3a/3b). To thirteen 4.5 mL bombs was added 40 mg of **2**. In a nitrogen atmosphere, 3.5 mL of a 0.1 M NaOCH_3 solution in dry methanol was added to each bomb. The bombs were sealed and placed in a 150°C oil bath for 3 hours. After being pulled and cooled, all bombs were emptied and rinsed twice with water. The bomb contents and rinsings were combined, acidified to pH 5 with dilute HCl, and CHCl_3 extracted (4

x 25 mL). The CHCl_3 extracts were dried over Na_2SO_4 and later evaporated to leave an oil (500 mg), which was a mixture of cis and trans vinyl ether isomers (**3a** and **3b**).

Column chromatography of the oil resulted in partial separation of the trans isomer; however, the yield was low. A $^1\text{H-NMR}$ of the separated cis isomer showed (CDCl_3) δ 2.32 (s, 3, Ar- CH_3), 3.85 (s, 3, OCH_3), 3.89 (s, 3, OCH_3), 5.48 (d, $J = 7\text{Hz}$, 1, ArCH=), 5.58 (s, 1, OH), 6.46 (d, $J = 7\text{Hz}$, 1, =CH-O), 6.6-7.1 (m, 5, Ar-H), and 7.57 (d, $J = 2.7\text{Hz}$, 1, Ar-H). The crude oil shows the same set of $^1\text{H-NMR}$ signals, plus additional ones assigned to the trans vinyl ether: (CDCl_3) δ 3.86 (s, OCH_3), 6.22 (d, $J = 12.4\text{Hz}$, ArCH=). Proton coupled and decoupled $^{13}\text{C-NMR}$ spectra of this oil were complex; however, the following signals were apparent: (CDCl_3) ppm 21.1 (q, Ar CH_3), 55.5, 55.7, and 55.8 (q, OCH_3), 140.7 and 143.4 (d, =CHO-), 108-150 (many signals, aryl and vinyl carbons). A mass spectrum of the oil (direct insertion probe) showed: m/e (% of base peak) 286 (M,100), 257 (14), 225 (36), 149 (14) 137 (18), 133 (16), and 77 (16). Integration of the $^1\text{H-NMR}$ spectrum of the crude oil indicated an ~ 2:1 mixture of cis/trans isomers.

Vinyl Ether Degradations. The reactions were conducted in a nitrogen atmosphere with oxygen-free water in 4 mL stainless steel pressure vessels (bombs). Typically, the bomb contained 30 mmoles of vinyl ether mixture **3a/3b** and 3.5 mL of 0.86M NaOH. The bombs were loaded in a glove bag with 1.0 mL of 0.5M NaOH containing **3a/3b** and mixed with 1 mL of 1.0M NaOH containing glucose or Na_2S , depending on the experiment, and diluted with 1.0M NaOH to a volume of 3.5 mL. In the appropriate experiment, 31.2 mg of anthraquinone was added. The sealed bombs were rotated for selected time periods in a 170°C oil bath, then cooled quickly, and opened. A small volume of 1.0M NaOH, containing a known amount of *p*-isopropylphenol (GC internal standard), was added; the contents of the bombs were then transferred to a flask.

The reaction mixture was treated with dimethyl sulfate, for conversion of the phenols to methyl ethers,⁴ and then extracted with chloroform. The CHCl_3 solution was washed with water, dried (Na_2SO_4), and analyzed by gas chromatography (Hewlett-Packard HP 5890 GC, HP-17 packed column). The amount of *m*-creosol in the sample was determined by comparing the GC signal area of methylated creosol to that of methy-

lated *p*-isopropylphenol. Standard mixtures of *m*-creosol and *p*-isopropylphenol were methylated and analyzed by GC to develop a response factor.

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