This article was downloaded by: On: 25 January 2011 Access details: Access Details: Free Access Publisher Taylor & Francis Informa Ltd Registered in England and Wales Registered Number: 1072954 Registered office: Mortimer House, 37- 41 Mortimer Street, London W1T 3JH, UK



# Journal of Wood Chemistry and Technology

Publication details, including instructions for authors and subscription information: <http://www.informaworld.com/smpp/title~content=t713597282>

## Reactions of ß-ARYL Lignin Model Quinone Methides with Anthrahydroquinone and Anthranol

John Ralphª; Richard M. Edeʰ; Nicholas P. Robinsonʰ; Lyndsay Mainʰ <sup>a</sup> New Zealand Forest Research Institute, Kotorua, New Zealand <sup>b</sup> Chemistry Department, University of Waikato, Hamilton, New Zealand

To cite this Article Ralph, John , Ede, Richard M. , Robinson, Nicholas P. and Main, Lyndsay(1987) 'Reactions of ß-ARYL Lignin Model Quinone Methides with Anthrahydroquinone and Anthranol', Journal of Wood Chemistry and Technology, 7: 2, 133 — 160

To link to this Article: DOI: 10.1080/02773818708085258 URL: <http://dx.doi.org/10.1080/02773818708085258>

# PLEASE SCROLL DOWN FOR ARTICLE

Full terms and conditions of use:<http://www.informaworld.com/terms-and-conditions-of-access.pdf>

This article may be used for research, teaching and private study purposes. Any substantial or systematic reproduction, re-distribution, re-selling, loan or sub-licensing, systematic supply or distribution in any form to anyone is expressly forbidden.

The publisher does not give any warranty express or implied or make any representation that the contents will be complete or accurate or up to date. The accuracy of any instructions, formulae and drug doses should be independently verified with primary sources. The publisher shall not be liable for any loss, actions, claims, proceedings, demand or costs or damages whatsoever or howsoever caused arising directly or indirectly in connection with or arising out of the use of this material.

#### **JOURNAL OF WOOD CHEMISTRY AND TECHNOLOGY, 7(2), 133-160 (1987)**

#### REACTIONS OF 6-ARYL LIGNIN MODEL QUINONE METHIDES WITH ANTHRAHYDROQUINONE AND ANTHRANOL 1.2

**John Ralph\* New Zealand Forest Research Institute,**  Private Bag. Rotorua. **New Zealand** 

**and** 

**Richard n. We. Wicholas P.** *pobinson.* **and Lyndsay -in**  Chemistry Department. University of Waikato. **Private Bag, Hai1ton. New Zealand** 

#### **ABSTRACT**

Quinone methides prepared in situ from phenylcoumaran and **B-C-1 lignln models** whlch **did not contain a 8-hydrorylsthyl group. readily formed addition products with anthranol but not**  with anthrahydroquinone. For **B-aryl** lignin models containing **the hydroryrsthyl group, the retro-aldol reaction (llberatlng formaldehyde) was so facile under the conditions used that rtilbene fomatiar fra the quimne methide took precedence over adduct formation.** 

#### DITRODUCTION

**A** great deal of activity has been directed toward the study **of the reactions of anthrahydroquinone** *(AHQ)* **and anthrarrol (reduction products of anthraquFnona,** *AQ)* **with quFnonc mthidcs of the B-aryl ether type. It is primarily the reactions of this lignln unit which are responsible for the accelerated cleavage of the Lignln macrmlecule in alkaline-additive pulping.** 

**Copyrbht** *0* **<sup>1987</sup>by Yd Dekksr, Inc.** 

**Although there is considerable speculation' as to whether adducts between** *AHQ* **(or anthranol) and B-ether quinone methides are intermediates in the catalytic cleavage of B-ether bonds under soda-A0 pulping conditions. there** Is **no doubt that such adducts are readily formed. 4** 

**B-Aryl ether quLnona rathides are not the only quinone methides which can Corn under pulping conditions: any freephenolic unit with an a-leaving group (OH. mr, or oa) can** *Cora*  quinone methides. Indeed, when anthranol. <sup>13</sup>C labelled at **the 9** and **10 positions. was reacted In base with acctylated milled-wood 1ignin (conditions which generate lignin quinone methides readily at roa temperature), anthraml-lignin adducts uere ~btained.~" Only two of a multiplicity of peaks** *in* **the C-10 region oC the 13C spectrm could be attributed to adducts with B-aryl ether units.**  *7* 

*AHp* **or anthranol. If so. it is 00 interest to deternine. firstly, ha the presence of** *AHQ Md* **other species affects the**  reaction pathways of the 8-aryl units and. secondly, whether reactions involving **B-aryl quinone methides may help account for the considerable loss of** *'AQ.* **fra the pulping cycle.**  *8*  **We wished to know if other quinone methides could also trap** 

**The rain objective of the work described in this paper was to deteralne if qulnom methides fra B-C linked structures**  react with AHQ and anthranol and to characterise any adducts **€om. Only phenylcornaran (B-C-5) and B-C-1 structures are considered here, although studies on other B-C linked models are also in progress in our laboratories.** 

#### RESULTS AND DISCUSSION

#### **Hode 1s**

The most readily available phenylcoumaran models are **dehydrodiisoeugenol 1,'** and **its reduction product dihydro-**









B

 $rac{H}{A}$ 

	ß	R,	$\mathsf{R}_2$	8	
ھڪا <b>14b</b> ម 15 $\mathbf{v}$ 18 19 29	cm <sub>3</sub> CH <sub>3</sub> CH <sub>2</sub> OH CH <sub>2</sub> OAc CH <sub>2</sub> OSi <sup>t</sup> BuMe <sub>2</sub> CH <sub>2</sub> OH <b>CH<sub>2</sub>OAc</b> CH <sub>2</sub> OSi <sup>t</sup> BuMe <sub>2</sub>	H н н н н CH <sub>3</sub> CH <sub>3</sub> CH <sub>2</sub>	CH=CH-CH3 $CH_2$ -CH <sub>2</sub> -CH <sub>3</sub> Ħ н н н н н	н 21 22 Ac	23 24

**FIGURS <sup>2</sup>**- *Quinarc* **methides** 

**136 RALPH El' AL.** 

dehydrodiisoeugenol <u>2</u> (Figure 1). Use of <u>2</u> rather than <u>1</u> removes **the caplication of further reactions of the vinyl side chain which** *are* **not characteristic of the phenylcoumaran roiety of lignin.** 

**Although the use of these easily synthesisad models is valuable in developing methods for the study of adduct**  formation, it is preferable to use a more representative model **such as 2 (Figure 1) which possesses the hydroryrathyl group present in aost lignin side chains. The presence of this group markedly influences the course of important reactions.** 

**<sup>A</sup>mdel representing a 'ring-opned'** *0-C-5* **unit (which could not cycllse to a phenylcorraaran) was also required in** *our*  **studies. The ervthro isoaer of model** *5* **(Figure 1).** In **which the B-ring phenolic group is aethylated, was prepared using essentially the method of Brunaw** and **Lundquist. LO** 

A base-stable t-butyldimethylsilyl protecting group in **capamds 2.** *9,* **and** *0.* **Increased the stability of the qulnone methides with respect to polyrerFsation and removed the possibility of retro-aldol reactions.** 

**node1 U was syntheslsed to represent free phmolic 0-C-1 units in lignin in which the B-ring phenol is etherifled.** 

## Anthranol and *AHQ Adducts with Quinone Methides of 8-aryl Models*

Alpha-aryl ethers such as compounds 1-5 (Figure 1) are **<sup>5</sup>**knam **to (reversibly) generate winone methides at a**  significant rate even at 10°C in 1M NaOH (Scheme 1). Therefore, **attempts to form adducts fra quinone mthides 14-17 (Figure 2) were made by addition of models** 1 **to 2 directly to solutions of anthranol or** *AHQ* **in base. sumarised** In **Table 1. These reactions gave products as** 

The anthranol adduct 27a from dehydrodiisoeugenol 1 **polymerised slowly on standing to give the polystyryl derivative** 



#### TABLE 1

Approximate yield (%) data for adduct reactions.<sup>a</sup>

- $\ddot{\phantom{a}}$ Conditions: 2 eq anthranol or AHQ; 1M NaOH (except 0.3 M for 4 to minimize hydrolysis of the  $\gamma$ -OAc group); 50°C; 1 hr  $(15$  min for  $\frac{8}{3}$ .
- Þ Ratio of erythro: threo isomers (in brackets) determined from H-1 MMR after acetylation.
- $\mathbf c$ 75% yield after flash chromatography.
- đ 80% after flash chromatography. Isolated erythro 41% and threo 39% as pure fractions.
- e Single reaction only; yield not optimised; other products not characterised. NRTR of crude material indicated possibility of threo isomer. approximately 90:10 erythro: threo.
	- Product not observed.

- **37. quantities of butylated hydroxytoluene. Polymerisation could be prevented by addition of trace** 

**Generation of the quinane methide fra the ring-opened**  phenylcoumaran model 6 at moderate temperatures required the w-OH to be replaced with a better leaving group. Attempts to form <u>7</u>, the a-bromide, from <u>6</u> using bromotrimethylsilane<sup>11</sup> were **unsuccessful due to spontaneous loss of formaldehyde, and**  formation of 4-hydroxy-3,2',3'-trimethoxystilbene from the **braide. Harsver. quinOna methide** *2* **could be generated in sltu fra the free phenolic diacetylated rode1 4 in base.**  *As* **there was** no **possibility of quincme methide** *9* **reverting to a phenylcouraran by an Internal cyclisation. the yield (Table 1) of anthranol adduct uas substantially higher Era model 4 than Era the true phenylcwmarans** *I-2.* 

**Analogously. the 8%-1 quinone methide** *22,* **generated Era the free phenolic diacetylated me1 13, was used for adduct reactions.** 

**silane" in chloroform. Treatrant of this solution with aqusow potusirn carbonate gave relatively stable solutions of quinone methide** *a.* **Alternatively. quimXre methide** *a* **was**  generated from the free-phenolic  $\alpha$ -acetate <u>10</u> in base. Reaction of anthranol with quinone methide 20 gave adduct 34 (after **acetylation) in very high yield as a** *50:50* **mixture OE ervthro**  and <u>threo</u> isomers. Unlike the parent acetylated adducts 32, **these were readily separated by tlc.**  Silylated model 9 could be brominated using bromotrimethyl-

A further point is apparent from Table 1. Whereas **anthranol adducts form readily fra these qulrwne methides, attempts to EON the corresponding** *AHp* **adducts (8.g.. R'roH of**  *m.* **Figure 3) using both aqueous and organic solvents. at temperatures ranging froa O°C to** *80.C.* **were unsuccessful, prestmably for a cabination of steric and electronic reasons. It has been noted previously ''12 that** *AliQ* **adds less readily** 







PIGURE 4 - Partial 200 MHz H-1 NMR spectrum of 8-C-5 adducts 32  $t = threq$ ,  $e = grythro$ 

**than anthranol to 8-aryl ether quinone methides and it has also been shown** 4.12 that. in competition studies. anthranol adducts **are formd** In **weruhelring preference.** 

#### Stereochemistry and NPIR Spectra of Adducts

**4,13 Despite the high kinetic steraoselectivity observed**  for' threo adducts from B-ether quinone methides (e.g., 23-24). both adduct isomers can be detected from attack of anthranol on most of the **B-aryl quinone methides (Figure 4 and Table 1)**.

**4.14.15** have fascinating NPUR characteristics due to  $\overline{a}$ **The B-aryl adducts 27-36 (Figure 3). like the B-aryl ether their conformations in solution. For example. in the ervthro lsoaar of** *jz* **(which is analogous to the threo lsosar in B-ether adducts because of the convention of group assignments) the A-ring is clearly situated over the anthracenyl ring systea. as**  is evidenced by the highly shielded ring A methoxyl and the ring A protons (Figure 4). However, the **minor** threo isomer is quite unlike the erythro isomer<sup>14</sup> of B-ether adducts in that ring A protons are more intensely shielded. The threo Y-acetate methyl **chemical shift is also anomalous. appearing at6 2.0 compared**  with a normal shift of 61.8. This methyl group is presumably in **a deshielding region of the anthracenyl ring system and/or the**  other aromatic rings. In addition, the B-ring methoxyl protons **experience substantial shielding, indicating their positions within the shleldlng regions of the anthracenyl ring system and/or ring A. Conformations of ervthro- and threo-32. postulated froa the information In their spectra. are shown in Figure 5.** 

Similar features appear in the spectra of B-C-1 adducts  $(e.g., Figure 6)$  indicating that small changes in the **substitution on ring B do not significantly alter the adduct conformations. Selected** *m* **data for these adducts are given In** 



erythro







adducts 36 in acetone-d6, (t) indicates peaks from the threo isomer

Solocted In the data for adducts

T. THEFT



÷

sindered rotation aschanging these positions, m. 3.59-3.05  $\ddot{\phantom{a}}$ 

\*\* Nimésicé rotation exchanging these positions. m. 3.79-3.85

Indicates resonances which are mot resolvable due to coincident chemical ahifts and/ox to line broadening caused by hindered<br>rolation.  $\ddot{\phantom{0}}$ 

Downloaded At: 13:23 25 January 2011

Between The man data for adducts 

MANGEL	$\lim_{n\to\infty} \alpha_n$	$\mathbf{u}_{\mathbf{1}\mathbf{v}}. \mathbf{G}$	Į	Nothing 16	<b>Acatate methyle</b>	ů	8	$\ddot{\phantom{0}}$	ដឹ	Ø	Montate C-9	i	÷
ã													
₿		Ù		22.8 59.6. M.O	$\bullet$	$\frac{1}{2}$	$\ddot{a}$	$\frac{1}{2}$	$\ddot{ }$	183.4	ı	¢	
		v	$\frac{3}{2}$	$-3.5.7$	$\ddot{\bm{x}}$	$\ddot{a}$	$\ddot{x}$	$\ddot{a}$	$\ddot{\bullet}$	183.1	$\frac{1}{2}$		
Ħ		ü		2.54 55.64 87.2	『宮 1918 1918	$\ddot{i}$	$\ddot{ }$	$\ddot{ }$	$\vec{•}$	182.9	167.4. 169.4. 171.4		
			x	$\frac{1}{2}$	<b>地震 医胃 医胃</b>	$\ddot{x}$	$\ddot{ }$	Į	$\ddot{\bullet}$	183.0	1414, 1414, 141		
	ŝ		z		1978 1978	$\frac{1}{3}$		$\ddot{ }$	$\ddot{\phantom{a}}$	182.9			
ă			3	$-2.7.82$ <b>GRI)</b> 2-11	$-2.5 - 2.8$	$\frac{1}{2}$	e. R	$\ddot{a}$	$\vec{a}$	$\frac{1}{2}$	140.6.170.4		
	ŝ		3	<b>M.2. 89.6</b> $\ddot{a}$ , $\ddot{a}$ , $\ddot{a}$	<b>Altreation</b>	$\frac{2}{3}$	$\ddot{a}$	$\ddot{a}$	$\ddot{3}$	142.7	160.4, 171.1		
			S	$\frac{11.4}{41.5}$ (a)	あい おい	$\ddot{\phantom{a}}$	$\ddot{x}$	$\vec{3}$	$\ddot{ }$	$\frac{1}{2}$	144.5.119.5		
			7	$\frac{1}{2}$ $\frac{1}{2}$ , $\frac{1}{2}$ , $\frac{1}{2}$ , $\frac{1}{2}$	<b>P.12, 21.8</b>	$\ddot{\dot{\mathbf{z}}}$	a A	$\ddot{\mathbf{3}}$	$\ddot{\phantom{0}}$	182.9	166.3, 171.2		
N			3	$\frac{1}{22}$ , $\frac{1}{22}$ , $\frac{1}{22}$ , $\frac{1}{22}$	$\ddot{\mathbf{a}}$	$\ddot{ }$	្ទ	$\ddot{\bm{s}}$	$\vec{v}$	្ន	$\ddot{3}$	$-7.07$	$\ddot{x}$
			\$	33.6, 33.9.	$\ddot{\mathbf{a}}$	ĵ	$\ddot{a}$	$\ddot{\bm{s}}$	$\ddot{\bullet}$	183.2	148.5	$-3.4 - 3.7$	$\ddot{x}$
	ŝ		3	<b>(21) 9.74 M</b> 53.3. 55.5	$\ddot{\mathbf{z}}$						14.3	$-3.6 - -9.7$	$\ddot{\hat{\pi}}$
				55.3. 58.7. M.7 (M3)	$\ddot{\mathbf{a}}$	;	$\ddot{\bullet}$	$\ddot{\bm{3}}$	$\ddot{•}$	i.com	$\ddot{3}$	$-3.2. -5.7$	$\ddot{x}$
계			2.4	$\frac{2}{3}$ 33.7	<b>R.B. S.B.</b>	$\ddot{\mathbf{3}}$	$\ddot{\ddot{\bm{x}}}$	$\ddot{ }$	$\ddot{\ddot{\bullet}}$	$\ddot{a}$	<b>ENG.</b> 19.191		
		٠ ż	2	$\frac{2}{3}$ $\frac{3}{3}$ $\frac{4}{3}$	<b>9.8 1.8</b>	$\ddot{3}$	$\ddot{ }$	i	$\ddot{3}$	$\frac{1}{2}$	168.6.119.6		

**B-ARYL LIGNIN MODELS** 

143

(1)  $\epsilon$  - anthus,  $\epsilon$  - jame (2)  $\epsilon$  - containing (3)  $\epsilon$  - containing (3)  $\epsilon$  - which are also use of  $\epsilon$ 

(\*) from a 75129 ott minister<br>(5) two rocamers freezes out on MR time scale

**144 RALPH ET AL.** 

**Tables 2 and 3. Full spectral details and assigments may be given in a separate publication.** 

**The 13C WaZ C-10 cheaical shifts of these adducts in**  acetone-d allow a more complete analysis of the C-10 region  $\frac{1}{100}$ **of the 13C** *NlQt* **of the anthranol-lignin adduct reported pre~ioruly.~ Figure** *7* **is a highly expanded plot of the C-10 region of the anthranol-lignin adduct on which is shown the**  positions of **8-aryl ether, phenylcoumaran, and 8-C-1 adducts. There Is little doubt that adducts between these three unit**  types account for most of the resonances in this region.

phenomena<sup>1123</sup> in their proton and <sup>23</sup>C MPUR spectra. The effect of temperature on the spectra of these adducts was far more complex **than that observed for certain B-aryl ether adducts." and indicated that the behaviour was not slmply due to rotation about a single** *bond.* **One compound. threo-34, was fully resolved at**  ambient probe temperature into two rotamers (60:40 ratio) on the **I3C llBl timescale (Figure** *8).*  **The broader resonances Era**  the major rotamer indicate that either the T<sub>2</sub> relaxation times **are shorter or that further fluxional behaviour is occurring in**  the major rotamer that is not significant in the minor rotamer. **Several of the adducts exhibited hindered rotation** 

**The observed stereoselectivity of anthranol attack can**  reasonably be attributed to the quinone methide conformations. **Figure 9 shows the two major conformers expected** for 8-aryl quinone methides. If the conformation does not significantly **alter the charge density at the alpha carbon** and **R is sterically less demanding than Ar (as expected for R = Me, CH<sub>2</sub>OH, CH<sub>2</sub>OAc), wwld be the major canformer. Attack Era the less hindered**  side would lead to the erythro adduct. As **R** becomes larger (e.g.,  $\texttt{CH}_2$ OSi<sup>t</sup>BuMe<sub>2</sub>), rotamer <u>B</u> becomes significant, resulting in more threo product.

For reasons that are not clear, these B-aryl quinone methides are markedly less stable toward polymerisation than **8-aryl ether quinone wthldes and, at modest concentrations. are observed only transiently at rocm temperature. Attempts to characterise them by techniques''** *are* **in progress.** 



**Pxme 7** - **1%** *NHR* **(in acetone-dg) of the C-LO region of an anthranol-lignin adduct7 and chemical shifts of model adducts (E** = **ervthro, T threo. B** = **B-aryl ether, PIC** = **phenylcouraran)** 



**FIGURE** *<sup>8</sup>*- **Partial 13C** *Mut* **of adduct threo-34 shawing hindered rotation** 

## **The Phenylcoumaran Quinone Methide: Competition Between Anthranol Adduct Formation and the Reversal Reaction to the Pheny 1 coumar an**

**Formation of quintme methides Era phenylcomarans 1-2 in**  base is clearly reversible (Scheme 1). The assumed steady state **quinone wthide concentration is too low to be detectable by cawentiara1 W-vlsible or** *HI¶R* **spectroscopy. The reverse**  reaction to the phenylcoumaran must therefore be very rapid. **Nevertheless. nucleophiles such as anthranol, obviously compete**  for the quinone methide.

**In order to ascertain haw eEEectively anthranol can capeta with the qulnone methide reversal reaction, the a-iodide 38 was prepared Ira dihydrodehydrodihoeugenol.** *2.* **by treatment with**  iodotrimethylsilane in chloroform. It was assumed that, on **treatment with base. the phenolic** TWS **group would rapidly**  hydrolyse and that the quinone methide 14b would rapidly form by **eliaination of iodide. The phenylcorrnraran 2 was isolated in high yield fra such a reaction. Addition of the iodide 38 to a solution of anthranol in base and workup within 5 minutes gave a**  ratio of adduct  $\frac{27b}{21}$  to phenylcoumaran 2 of approximately 10:90. Since only traces of adduct <u>27b</u> were detected when phenylcoum. **3. is reacted with anthranol under the same conditions for 5 minutes, it is ccmcluded that anthranol attack is about 10% as rapid as the recyclisation reaction.** 





FIGURE 9 - Major conformers of 8-aryl quinone methides

## The Phenylcoumaran Quinone Methide: Competition Between Adduct Formation and the Retro-Aldol Reaction

During much of this work, quinone methides were most conveniently generated from free-phenolic acetates or t-butyldimethylsilylated compounds, but the use of these derivatives precludes one very important reaction - the retro-aldol elimination of formaldehyde. To relate the results of model studies to reactions involving lignin itself, it is essential that the model contains the hydroxymethyl group.

Despite its surprising effectiveness at trapping quinone methides. Not even anthranol could compete against the retro-aldol elimination of formaldehyde from quinone methides 15. 18 or 21 to give stilbenes. The reaction therefore follows the same course as in the absence of the additive (Schemes 1 and  $2)$ .  $17$ 

This point illustrates the pitfalls of using inappropriately substituted or derivatised models. Acetylated models, 4,18 or acetylated lignin itself.<sup>6.7</sup> are frequently used to allow in-situ generation of guinone methides at room temperature. But the protection afforded to the hydroxymethyl group by acetylation means that formaldehyde loss by a retro-aldol reaction cannot compete in subsequent quinone methide reactions. This is not so







## **B-ARYL LICNIN MODELS** *149*

**critical** In **the case of B-aryl ether quinone methides since**  anthranol and AHQ can add to quinone methide 23 efficiently before the retro-aldol reaction (to give the styryl ether) can **occur.12 However, in phenylcoumaran and &c-1 models, the retro-aldol reaction entirely dominated the reaction of**  representative quinone methides 15, 18, or 21 (with the **B-CH<sub>2</sub>OH substituent) whereas adducts readily formed from the acetylated quinone methides 16,** *19,* **or** *22,* **or the silylated**  derivatives 17 or 20. Thus. reactions on acetylated milled-wood **lignin with anthranol. where adducts other than 8-ether adducts appear to be observed in the C-13** *NMR* **spectra,<sup>6,7</sup> may not be** representative of reactions of the underivatised quinone  $nethides.$ 

#### **coucLusrows**

**Mduct formation betmen 8-dry1 quinone methldes and anthranol. but** *not AHQ,* **occurs readily. but only when the possibility of formaldehyde ellmlnation fra the quinone methide is removed. Consequently. the reactivity of lignin phenylcoraaran and B-C-1 units in soda-AQ pulplng is not expected to be altered by the presence of** *AQ* **species. Conversely. it is not expected that** *KQ* **losses can be attributed to reactions of**  its reduction products with  $\beta$ -aryl quinone methides.

#### **EXPERIMENTAL**

<sup>1</sup>H NPIR spectra were determined in  $CDCl_3$  or acetone-d<sub>6</sub> **on a CU Varlan T-60. a JBOL Rt30Q. or a Bruker** *ACMQ* **R spcctro**meter using tetramethylsilane as an internal reference. **NHR** spectra were determined in CDCl<sub>3</sub> or acetone-d<sub>6</sub> on a JWOL **FX9OQ (22.5 IQfz) or Bruker AC200 (50 rocZ) FT spactrarster Using tetran#thylsilane as an internal reference.**  *29*  **sl bRB spectra**  were determined in CDCl<sub>3</sub> on a Bruker AC200 (39.8 MHz) spectrometer using a broadband proton-decoupled <sup>29</sup>si DEPT<sup>19,20</sup> pulse sequence with 1/2J delays of 72 ms. and a 0 pulse of approximately  $17^{\circ}$  (optimised).  $29^{11}_{10}$  coupling constants  $\binom{2}{J}$  = 6.53,  $\frac{3}{J}$  = 6.96 Hz) were determined from the  $\frac{29}{J}$  Si satellites in  $\frac{1}{1}$  HMHR spectra. Mass spectra were determined on an HP 5985 quadrupole GC/MS under electron impact conditions using 70 eV ionizing energy (direct insert probe).

## General Methods

#### Preparation of TBDMS derivatives<sup>21</sup>

The primary alcohol (1.0 eq), t-butyldimethylsilylchloride (1.5 eq). and diazabicyclo[5.4.0]undec-7-ene (DBU, 1.4 eq) in CH<sub>2</sub>Cl<sub>2</sub> (10 ml/mmole 1.2-diarylpropane-1.3-diol) were stirred for 40 minutes at 40°C. The products were extracted into CH<sub>2</sub>Cl<sub>2</sub> and washed three times with saturated aqueous MH<sub>4</sub>Cl. The organic phase was dried over MgSO<sub>4</sub> and the solvent removed to give the silyl ethers in ca. 95% yield after purification.

### Quinone Methides

Quinone methides 14-24 were generated in situ either from the free phenolic phenylcoumarans  $(\frac{1}{2}, \frac{5}{2})$ , or from the free phenolic ¤-acetylated derivatives (8, 10, 13).

## Anthrahydroguinone (AHQ and  $AHQ^{2-}$ )

Solutions of  $M_0^2$  in aqueous 1 M or 0.3 M MaOH containing, or free from, sodium dithionite were prepared as described in reference 4.

#### Anthranol

Solutions of the anion of anthranol were prepared by refluxing a mixture of anthrone in 1 M or 0.3 M NaOH (20-50 ml) **under nitrogen until dissolution was colplete (.bout 1** hour). The solutions were then cooled to the required temperature.

### *Am)* **or Anthrurol** *lldduc* **tS**

**To a solution of** *AHQ* **or anthranol (2** *eq)* **in base (at the tampstature given In Table 1) was added the qulnone methide**  precursor in solid form or as a solution in a small volume of **Q12C12. After the appropriate tbe (usually 1 hour for**  phenylcoumarans and 15 minutes for Y-acetate quinone methide **precursors). the mixture was neutralbed with 5% H** *so* **and 24 extracted with aC13 (2 XI. The chloroform extract was dried**   $over$  **HgSO4** and the solvent removed.

## **Specific Syntheses**

**pehvdtodiisoeuqenol (1). Dehydrodiisoeugenol** was **prepared 9 in 50% yield from boeugenol.** 

 $Dihydrodehydrodi}$ isoeugenol (2). Dehydrodiisoeugenol (1, 1.0 g. 3.07 mmole) in 95% ethanol was hydrogenated at atmospheric **pressure over 5% W/c (2.5 mg) for 2 hours (yield quantitative).**  Crystallisation from petroleum ether (100-120°C fraction) gave white needles m.p. 89-91°C.

**method of Brunow and Lundquist.<sup>10</sup> The major by-product of the BP3.Rt 0 rearrangeaent of the chalcone epoxide was identified 2**  as fluorohydrin  $\frac{39}{2}$  (yield 42%). **md 3 was prepared in 30% overall yield uslng the** 



Compound 39 was a white crystalline solid: m.p. 120-122°C;  $\frac{1}{2}$   $J_{\text{OH}-P}$  = 0.9 Hz, B-OH), 2.24 (1H, dd,  $J_{\text{OH}-\alpha}$  = 4.9 Hz,  $J_{\text{OH}-P}$ = 1.5 Hz,  $\alpha$ -OH), 3.81 (3H, s, OCH<sub>3</sub>), 3.89 (3H, s, OCH<sub>3</sub>). 4.06 (1H, dddd, J<sub>BP</sub> = 25.6 Hz, J<sub>B</sub><sub>Q</sub> = 6.8 Hz, J<sub>B</sub><sub>OH</sub> = 5.4 Hz,<br>J<sub>BT</sub> = 2.4 Hz, HB), 4.74 (1H, dd, J<sub>QB</sub> = 6.8 Hz, J<sub>G-OH</sub> = 4.9 Hz, Ha), 4.97 (lH, d, J = 10.9 Hz, B ring benzyl CH), 5.04 (lH, d,  $J = 10.9$  Hz. B ring benzyl CH). 5.12 (2H. s. A ring benzyl CH<sub>2</sub>). 5.98 (IH. dd. J<sub>TP</sub> = 45.9 Hz. J<sub>TB</sub> = 2.4 Hz. HT). 6.70-7.20 (6H. m. Ar-H). 7.30-7.50 (10H. m. benzyl Ar-H). Mass Spectrum m/z: 518 ( $m^2$ , 0.1), 256 (5), 242 (3), 238 (3), 227 (3), 165 (3), 137 (4), 92 (8), 91 (100), 65 (10).  $^{13}$ C NRTR of diacetate (22.49 MHz, CDCl<sub>3</sub>) 6: 20.1 (a-OCOCH<sub>3</sub>), 20.8 ( $\beta$ -OCOCH<sub>3</sub>), 55.7, 55.8 (3,3' OCH<sub>3</sub>), 70.8 (4-benzyloxy CH<sub>2</sub>), 72.5 (d,  $3J_{\text{QF}} = 7$  Hz, a), 73.6 (d,  $2J_{\text{BF}} = 18$  Hz, 8), 74.3 (2'-benzyloxy CH<sub>2</sub>). 86.7 (d. <sup>PT</sup>J<sub>TF</sub> = 178 Hz. 7). 111.3 (2).<br>112.8 (4'). 113.4 (5). 118.8 (d. <sup>3</sup>J<sub>6'F</sub> = 9 Hz. 6'). 120.3 (6), 123.1 (5'), 127.2 (benzyl 3.5), 127.7 (benzyl 4), 128.4 (benzyl 2.6), 129.5 (1), 129.6 (d,  $2J_{1+P} = 20$  Hz, 1'), 136.9 (benzyl 1), 137.8 (benzyl 1'), 144.0 (d,  $3J_{21g} = 9$  Hz, 2'), 148.1 (3), 149.2 (4), 152.0 (3'), 168.7 ( $\alpha$ -ogoc $H_3$ ), 169.7  $(0.020CH, )$ .

Compound  $\overline{4}$ . Acetylation of compound  $\overline{3}$  to give the diacetate, followed by removal of the phenolic acetate<sup>22</sup> with pyrrolidine (1.2 eq) in CHCl<sub>2</sub> for 30 minutes, gave  $\frac{1}{2}$  as a clear oil in 90% overall yield.  $\frac{1}{2}$  New (200 MHz, CDCl<sub>3</sub> 6: 2.00 (3H, s,  $\gamma$ -OCOCH<sub>3</sub>) 3.30 (1H, m, HB), 3.78 (3H, s, OCH<sub>3</sub>), 3.88 (3H, s, OCH<sub>3</sub>), 4.36 (1H, dd, J<sub>B71</sub> = 7.6 Hz, J<sub>7172</sub> = 11.1 Hz,  $\gamma_1$ ), 4.47 (1H, dd,  $J_{\beta\gamma2} = 5.5$  Hz,  $J_{\gamma1\gamma2} = 11.1$ Hz,  $\gamma_2$ ), 5.72 (1H, d,  $J_{\text{eff}} = 6.9$  Hz, Ha), 5.80 (1H, s,  $Ar$ -OH), 6.8-7.2 (6H, m, Ar-H).

Compound 5. t-Butyldimethylsilylation prior to debenzylation in the synthesis of compound 3 gave 5 (88% overall yield).  $\frac{1}{11}$  MMMR (200 MMHz, CDCl<sub>3</sub>) 6: 0.04, 0.06 3H, (2 x 3H, 2s, SiMe<sub>2</sub>), 0.89 (9H, s, SiBu<sup>t</sup>), 3.65 (1H, m, HB), 3.84 (3H, s. OCH<sub>3</sub>). 3.90 (3H. s. OCH<sub>3</sub>). 3.80-4.00 (2H. m. H $\gamma$ 's). 5.55 (1H, d. J<sub>og</sub> = 5.8 Hz, H<sup>o</sup>), 5.60 (1H, s, Ar-OH), 6.80-7.00 (6H,<br>m, Ar-H).  $^{13}$ C NMR (50 MHz, CDC1<sub>3</sub>) 6: -5.31, -5.28 (SiMe<sub>2</sub>), 18.3 (Me<sub>3</sub>CS1), 25.9 (Me<sub>3</sub>CS1), 54.2 (B), 56.0  $(ORe's), 65.5 (7), 87.9 (a), 108.7 (A2), 112.1 (B4), 114.3 (A5),$ 117.1 (B6), 119.1 (A6), 121.2 (B5), 128.0 (A1), 133.7 (B1), 144.5 (B2), 145.7 (A4), 146.7 (A3), 148.3 (B3).

Compound 6 was prepared by modifying the procedure for synthesis of compound 3.<sup>10</sup> Condensation of 2,3-dimethoxybenzaldehyde 40 with the benzyl ether of acetovanillone following the method of reference 10 gave the chalcone 1-(4-benzyloxy-3-methoxyphenyl)-3-(2,3-dimethoxyphenyl)-2-propen-1-one 41 as a yellow oil. Crystallisation from hot ethanol gave 41 as yellow crystals m.p. 94-95°C (yield 76%).  $\frac{1}{2}$  <u>H NAME</u> (60 MHz, CDCl<sub>3</sub>) 6: 3.80 (6H, s, OCH<sub>3</sub>), 3.90 (3H, s.  $OCH_2$ ), 5.18 (2H, s, CH<sub>2</sub>Ph), 6.70-7.80 (11H, m, Ar-H). 7.40 (IH, d J = 16 Hz, HB), 8.08 (IH, d, J = 16 Hz, HY).  $^{13}$ C MMR (22.5 MMz, CDCl<sub>3</sub>) 6: 55.8, 55.9 (3.3'-OCH<sub>3</sub>), 61.6  $(2'-OCH<sub>3</sub>)$ , 70.7 (benzyl CH<sub>2</sub>), 111.3 (C2), 112.2, (C4'). 114.0 (C5), 119.6 (C6'), 122.8, (C6) 123.2 (C5'), 124.1 (CB), 127.2 (benzyl C3, C5). 128.0 (benzyl C4), 128.6 (benzyl C2, C6), 129.2 (Cl), 131.6 (Cl'), 136.2 (benzyl Cl), 138.7 (CY), 148.8  $(C2')$ , 149.6  $(C3)$ , 152.3  $(C3')$ , 153.1  $(C4)$ , 188.7  $(\alpha - C = 0)$ . Mass spectrum  $m/z$ : 404 ( $m^{\dagger}$ , 12), 373(13), 254(19), 91(100).

Spoxidation was more simply carried out by stirring chalcone 41 (0.05 mole) in pyridine (160 ml) with aqueous NaOCl  $(5.43, 160$  ml) for 3 hours.<sup>23</sup> Workup gave a white solid which was recrystallised from hot ethanol to give the chalcone epoxide 1-(4-benzyloxy 3-methoxyphenyl)-3-(2.3-dimethoxyphenyl)-2.3epoxypropan-1-one 42 as white needles m.p. 123-124°C (76%) yield).  $\frac{1}{2}$  MMMR (60 MHz, CDCl<sub>3</sub>) 6: 3.79 (3H, s, OCH<sub>3</sub>).

3.88 (3H, s, OCH<sub>3</sub>), 3.94 (3H, s, OCH<sub>3</sub>), 4.20 (1H, d, J = 2 Hz, HB), 4.32 (IH, d, J = 2 Hz, H $\gamma$ ), 5.21 (2H, s, CH<sub>2</sub>Ph), 6.70-7.70 (11H. m. Ar-H).  $^{13}$ C MMR (22.5 MHz. CDCl<sub>3</sub>)  $\delta$ : 54.9 (CB), 55.3, 55.4 (3-, 3'-OCH<sub>3</sub>), 59.5 (CY), 60.5  $(2'-OCH<sub>3</sub>)$ . 70.2 (benzyl CH<sub>2</sub>). 110.1 (C2). 111.8 (C4'). 112.3 (C5), 116.5 (C6'), 122.7 (C6), 124.0 (C5'), 126.8 (benzyl C3, C5), 127.6 (benzyl C4), 128.1 (benzyl C2, C6), 128.4, 129.1 (C1, Cl'), 135.6 (benzyl Cl), 147.6 (C2'), 149.6 (C3), 152.0 (C3'),  $152.7$  (C4),  $191.0$  (C=0).

Rearrangement of  $\underline{42}$  with BF<sub>3</sub>.Et<sub>2</sub>0 in Et<sub>2</sub>0 followed by reduction of the crude mixture with NaBH /OH gave a colorless oil which crystallised on standing. Recrystallisation from CH<sub>2</sub>Cl<sub>2</sub>/pet. ether (40-60°C) gave 1-(4-benzyloxy-3-methoxyphenyl)-2-(2,3-dimethoxyphenyl)-propan-1,3-diol 43, m.p. 122.5-123.5°C (yield 61%, c.f. 38% in synthesis of  $\underline{3}^{10}$ ). No fluorohydrin was detected among the by-products.  $\frac{1}{11}$  MMR (60 MHz, CDCl<sub>2</sub>) 6: 2.50 (2H, br s, OH), 3.55 (3H, s, OCH<sub>3</sub>), 3.60 (3H. s. OCH<sub>3</sub>). 3.78 (3H. s. OCH<sub>3</sub>). 3.20-4.20 (3H. m. HB.  $Hy's$ ), 4.90 (1H, d, J = 6 Hz, Ha), 5.05 (2H, s, CH<sub>2</sub>Ph). 6.60-7.60 (11H. m. Ar-H).  $^{13}$ C Meg (22.49 MHz, CDCl<sub>3</sub>) 6: 47.8 (CB), 55.6, 55.7 (3-OCH<sub>3</sub>, 3'-OCH<sub>3</sub>), 60.5 (2'-OCH<sub>3</sub>), 63.8 (CY), 71.0 (benzyl CH<sub>2</sub>), 74.8 (Ca), 110.5 (C2), 111.0  $(C4')$ , 113.7 (C5), 118.8 (C6), 121.1 (C6'), 123.7 (C5'), 127.3 (benzyl C3, C5), 127.6 (benzyl C4), 128.4 (benzyl C2, C6), 132.7 (Cl), 135.9 (Cl'), 137.1 (benzyl Cl), 147.3 (C2'), 147.7 (C3), 149.3 (C4), 152.6 (C3').

Debenzylation of 43 in wet dioxane with a catalytic amount of 5% Pd/C under 1 atm  $H_2$  for 2 hours gave 6 as a colorless oil, yield 100%.  $\frac{1}{2}$  MMR (60 MMz, CDCl<sub>3</sub>) 6: 2.60 (2H, br s. OH), 3.62 (3H, s, OCH<sub>3</sub>), 3.66 (3H, s, OCH<sub>3</sub>), 3.80 (3H, s, OCH<sub>3</sub>), 3.30-4.40 (3H, m, HS, H<sup>T</sup>'s), 4.90 (1H, d, J = 6 Hz, Hz), 6.60-7.00 (6H, m, Ar-H).  $^{13}$ C MHZ (50 MHz, CDCl<sub>3</sub>) ô: 48.2 (CB), 55.6, 57.7 (3.3'-OCH<sub>3</sub>), 60.7 (2'-OCH<sub>3</sub>), 63.9

 $(C\gamma)$ , 75.1  $(C\alpha)$ , 109.4  $(C2)$ , 111.1  $(C4')$ , 114.1  $(C5)$ , 119.5  $(C6)$ , 121.0  $(C6')$ , 124.0  $(C5')$ , 132.8  $(C1)$ , 134.3  $(C1')$ , 145.2  $(C4)$ , 146.5  $(C3)$ , 147.8  $(C2')$ , 152.8  $(C3')$ .

Compound 7. Attempted bromination of 6 using bromotrimethylsilane<sup>11</sup> gave only 4-hydroxy-3.2'.3'trimethoxystilbene, presumably via HBr and formaldehyde elimination from bromide 1.  $\frac{1}{2}$  if MPR (60 PHz, CDCl<sub>2</sub>) 6: 3.80 (9H, s. OCH<sub>3</sub>). 6.40-7.20 (8H, m. ArH and vinyl H).

Compound 8. Acatylation prior to debenzylation in the above synthesis of compound  $6$  gave  $8$  in 74% overall yield for the two steps.  $\frac{1}{2}$   $\frac{1}{$  $\gamma$ -OCOCH<sub>3</sub>), 1.86 (3H, s, a-OCOCH<sub>3</sub>), 3.68 (3H, s, OCH<sub>3</sub>), 3.80 (3H, s, OCH<sub>3</sub>), 3.85 (3H, s, OCH<sub>3</sub>), 4.05 (2H, m HY<sub>1</sub>, B). 4.25 (1H. dd. J<sub>872</sub> = 5.8 Hz. J<sub>7172</sub> = 9.8 Hz. H7<sub>2</sub>). 5.60 (lH. s. Ar-OH), 6.05 (lH. d.  $J_{\alpha B}$  = 5.8 Hz. HB), 6.70-7.05 (6H, m, Ar-H).  $^{13}$ C MMR (22.5 MHz, CDCl<sub>3</sub>) 6: 20.7  $(\gamma$ -OCOCH<sub>2</sub>), 21.0 (a-OCOCH<sub>2</sub>), 42.1 (B), 55.6, 55.8 (A3, B3 OCH<sub>3</sub>), 60.6 (B2-OCH<sub>3</sub>), 64.7 (7), 75.5 (a), 110.0 (B4), 111.2  $(A2)$ , 114.2  $(A5)$ , 120.3, 120.3  $(A6, B6)$ , 123.6  $(B5)$ , 130.6 $(A1)$ , 131.9 (B1), 145.6, 146.3, 147.8 (A3, A4, B2), 152.5 (B3), 169.8 (а-ососн<sub>3</sub>), 170.7 (т-ососн<sub>3</sub>).

Compound 9. t-Butyldimethylsilylation prior to debenzylation in the above synthesis of compound 6 gave 9 in 92% overall yield.  $\frac{1}{2}$  MMRR (200 MHz, CDCl<sub>3</sub>) 6: -0.049, -0.043  $(2 \times 3)$ H, 2s, SiMe<sub>2</sub>), 0.88 (9H, s, Bu<sup>t</sup>Si), 3.65 (3H, s, OCH<sub>3</sub>), 3.65 (1H, m, HB), 3.68-3.78 (2H, m, H  $\gamma$  s), 3.76 (3H, s, OCH<sub>3</sub>), 3.85 (3H, s, OCH<sub>3</sub>), 5.14 (1H, d, J = 4.5 Hz, Ha), 5.55 (1H. s. ArOH). 6.60-7.00 (6H. m. Ar-H).

Compound 10. t-Butyldimethylsilylation, followed by acetylation prior to debenzylation in the above synthesis of compound 6 gave 10 in 85% overall yield over the three steps.  $\frac{1}{2}$  (200 MHz, CDCl<sub>3</sub>) 6: 0.15, 0.18 (2 x 3H, 2s, SiMe<sub>2</sub>). 1.05 (9H, s, Bu<sup>t</sup>Si), 1.98 (3H, s,  $\alpha$  -OCOCH<sub>3</sub>), 3.82 (3H, s,

OCH<sub>2</sub>), 3.86 (1H, m, HB), 3.94 (3H, s, OCH<sub>2</sub>), 3.90-4.10 (2H, m, HY's), 3.98 (3H, s, OCH<sub>3</sub>), 6.00 (1H, s, Ar-OH), 6.34 (1H, d.  $J_{\alpha R}$  = 7.8 Hz. Ha). 6.80-7.40 (6H. m. Ar-H).

Compound 11. Compound 11 was prepared by amalgamating synthetic schemes from Berndtsson et al.<sup>24</sup> and Nakatsubo and Higuchi.<sup>25</sup> Thus, benzyl vanillin was condensed with homoveratric acid to give an acid 44 corresponding to compound 6 in ref. 24 in 90% yield as a mixture of threo and erythro isomers in a 3:1 ratio. The two isomers were separated by fractional crystallisation from acetone-hexane. <u>Brythro-44</u> was a white crystalline solid; m.p. 175-182°C. <sup>1</sup>H NPER (90 MHz. CDCl<sub>3</sub>) 6: 3.67, 3.68, 3.73 (9H, 3s, methoxyls), 3.83 (lH, d,  $J_{\alpha}$  = 9 Hz, HB). 5.05 (2H, s, CH<sub>2</sub>Ph), 5.32 (1H, d, J<sub>20</sub> = 9 Hz, Hz), 6.63-7.31 (11H, m, aromatics), 9.60 (1H, bs, COOH). Threo-44 was a white crystalline solid, m.p. 150-152°C; <sup>1</sup>H NPER (90 MHz, CDCl<sub>3</sub>)  $\delta$ : 3.67, 3.68, 3.73 (9H, 3s, methoxyls), 3.83 (unresolved, Hß), 5.00 (2H, s, CH<sub>2</sub>Ph), 5.14 (1H, d, J<sub>20</sub> = 9 Hz, Ho), 6.63-7.31 (11H, m, aromatics), 9.60 (1H, bs, COOH).

As the diborane reduction of acids 44 did not proceed well. each isomer of the acid 44 was methylated with diazomethane to give the ester 45 corresponding to compound 5 of ref. 25 in quantitative yield (approximately 70% yield after 1 recrystallisation). Erythro-45 was a white crystalline solid; m.p. 145.5-147°C;  $\frac{1}{2}$  MPR (60 MHz, CDCl<sub>3</sub>) 6: 3.55 (3H, s, acetate methyl) 3.80 (lH. d.  $J_{\beta\alpha} = 7$ . Hß). 3.83. 3.86. 3.86 (9H. 3s. methoxyls) 5.12 (2H, s, CH<sub>2</sub>Ph), 5.18 (1H, d,  $J_{\alpha\beta} = 7$ , Ho). 6.65-7.37 (11H. m. aromatics):  $^{13}$ C MRR (22.5 MHz, CDCl<sub>3</sub>) 6: 51.9 (ester methyl), 55.9 (methoxyls), 59.2 (CB), 71.0 (CH<sub>2</sub>Ph), 74.9 (Ca), 172.9 (ester carbonyl); mass spectrum m/z 434 ( $M^{\frac{1}{2}}$  -H<sub>2</sub>O, 0.5), 242 (13), 210 (32), 151 (41), 91 (100). Three-45 was a light yellow crystalline solid (ex acetane-hexane); m.p. 131.5-132.5°C;  $\frac{1}{2}$  MMMMMMM (60 MHz, CDCl<sub>3</sub>) 6: 3.72 (3H, s, acetate methyl), 3.75 (unresolved, Hß), 3.70, 3.70, 3.77 (9H, 3s,

methoxyls), 5.03 (2H, s, CH<sub>2</sub>Ph), 5.18 (1H, d, J<sub>og</sub> = 8 Hz,<br>Ha), 6.57-7.33 (11H, m, aromatics);  $^{13}$ <u>C NHR</u> (22.5 MHz, CDC1<sub>3</sub>) & 52.2 (ester methyl), 55.8 (methoxyls), 59.5 (CB), 71.0 (CH<sub>2</sub>Ph). 76.4 (Cu). 174.0 (ester carbonyl): mass spectrum. m/z 434 ( $M^{\dagger}$  - H<sub>2</sub>O, 0.5), 242 (11), 210 (44), 151 (43), 91 (100).

Debenzylation, acetylation and reduction, essentially as described in ref. 25. gave the required  $B-C-1$  model  $11$ . triacetate of threo-ll was a colourless oil; <sup>1</sup>H NAME (60 MHz, CDCl<sub>2</sub>) 6: 2.05, 2.14, 2.31 (9H, 3s, acetate methyls), 3.35 (1H, m. HB); 3.73, 3.80, 3.88 (9H, 3s, methoxyls), 4.49 (2H, m. HYs), 6.10 (lH, d, J = 9, Ha), 6.58-6.95 (6H, m, aromatics);  $\frac{13}{13}$ C NRTR (22.5 MHz, CDCl<sub>3</sub>) 6: 20.5, 20.8, 21.1 (acetate methyls), 49.5 (CB), 55.7 (methoxyls), 64.3 (C $\gamma$ , 75.9 (Cm), 168.6 (Рhoсосн<sub>3</sub>), 169.7 ( $\alpha$ -ососн<sub>3</sub>), 170.8 ( $\gamma$ -ососн<sub>3</sub>; mass spectrum m/z 460 ( $M^{\uparrow}$ , 9), 400 (6), 358 (5), 237 (14), 223  $(100)$ , 195 (50), 181 (37), 164 (96), 153 (64), 43 (56).

The free phenol diacetate 13 was prepared by acetylation of the ester 45 followed by LAH reductions, re-acetylation and catalytic removal of the benzyl group. Threo-13 was a colourless oil;  $\frac{1}{H}$  MMMR (60 MHz, CDCl<sub>3</sub>) 6: 2.03, 2.10 (6H. 2s. acetate methyls), 3.49 (lH. m. HB), 3.70, 3.75, 3.83 (9H. 3s, methoxyls), 4.43 (2H, m, H $\gamma$ s), 5.98 (1H, d, J<sub>oR</sub> = 8.4, Ho), 6.53-6.75 (6H, m, aromatics).

## Anthranol Adducts

Conditions, yields and erythro: threo ratios are given in Table 1; selected NRTR data are given in Table 2 and <sup>13</sup>C NRTR data in Table 3.

Adduct 27a: Prepared from model 1. The erythro isomer was isolated by flash chromatography using chloroform as eluant. Acetylation gave adduct 28a. On standing, compound 27a spontaneously polymerised, presumably to the styrene polymer 37 as indicated by the line broadening, the loss of the allylic

methyl resonance, and the appearance of a new (broad) aliphatic methyl resonance (6ca. l).

Adduct 27b: Prepared from model 2. The erythro isomer was isolated by prep. tlc using EtOAc:hexane as eluant. Acetylation gave adduct 28b.

Adduct 29: Prepared from model 4. Acetylation gave 31. Attempts to separate threo and erythro isomers were unsuccessful.

Adduct 30: Prepared from model 8. Acetylation gave 32. Small scale separation on analytical tic plates using multiple elution with EtOAc-hexane gave 300 µg pure erythro adduct.

Adduct 33: Prepared from model 5, followed by acetylation. Adduct 34: Prepared from model 10, followed by acetylation. 29<br>Si NMR spectra also showed hindered rotation features. Threo-34: 6 19.96, 19.99; erythro-34 6 19.39 (broad).

## Competition for the Quinone Methide Between Anthranol Adduct Formation and Reversal to the Phenylcoumaran

Model 2 (160 mg, 1.0 eq) in CDCl<sub>2</sub> was treated with trimethylsilyl iodide (117 mg, 1.2 eq) for 3 minutes to give the a-iodide 38 (as evidenced by <sup>1</sup>H MMTR, 6 5.5,  $J_{\kappa R}$  ca. 10 Hz). This solution was rapidly added to a solution of anthranol (190 mg) in aqueous base (1 M MaOH), stirred for 5 minutes, then neutralised and extracted with CHCl<sub>3</sub>, etc., as in the general method for adduct formation. The resultant product mixture was approximately 10% anthranol adduct 27b to 90% phenylcoumaran 2 by  $\frac{1}{2}H$  MPIR.

#### **ACKNOWLEDGEMENTS**

The authors are indebted to Drs J.M. Uprichard, I.D. Suckling, T.J. Fullerton (New Zealand Forest Research Institute) and L.L. Landucci (U.S. Forest Products Lab., Madison, Wisconsin) for helpful discussion, and to Dr G. Brunow

**(University of Helslnkl. Finland) for his helpful suggestions**  regarding his syntheses of the phenylcoumaran models. **for R.H.E. through Forest Research Institute Post Graduate Study Award Yo. 206 is gratefully acknowledged. Funding** 

#### **REPERENCES**

- **1. Ralph. J.** and **Me, R.H. Reactions of phenylcouraran llgnln rodel** quhone **methides with AHQ/anthranol. Presented at**  the 1985 International Symposium of Wood and Pulping **Chomlstry. Vancouver, 1985.**
- **2. me, R.H. and Ralph. J. Phenylcollraran qulnone wthides. Presented at the 1985 weu Zealand Institute of chemistry Conference. Paper Yo. 17-0, Chrlstchurch, New Zealand.**
- **3. Dlmel, D.R. J.** Wood *Chep.* **Technol., 3, 1 (1985).**
- **4.**  Landucci. L.L. and Ralph. J. J. Org. Chem., 47, 3486 **(1982).**
- **5.**  In Chemistry of Delignification with Oxygen, **\*am, and Peroxides. p. 107, mi Publlsherr Co.. Tokyo. Japan (1980).**
- 6. Landucci, L.L. J. Wood Chem. Technol., 1, 61 (1981).
- **7.**  Ralph. J. and Landucci. L.L. J. Wood Chem. Technol., 6, 73  $(1986)$ .
- **8.**  Landucci, L.L. and Ralph, J. J. Wood Chem. Technol., 4, **149 (1984) and references therein.**
- **9. Loopold. B. Acta Cham. %and.. 4, 1523 (1950).**
- **10.**  Brunow. G. and Lundquist. K. Acta Chem. Scand., B38, 335 ( **1984).**
- **11. Ralph. J. and Young, R.A. J.** Wood **Cham. Technol.. 3. 161 (1983).**
- **12.**  Ralph, J. Lignin model quinone methides, facts and **fallacies. Presented at the 1985 Internatlonal of** vood **and Pulplng Chemistry, Vancouver 198s.**

**13.**  It has recently been shown that a kinetically favoured **threo anthranol-B-aryl ether qufnone methide adduct**  isomerises to the erythro isomer in basic anthranol **solutlon:** 

Poppius, K. Acta Chem. Scand., 839, 861 (1985).

- **14. Ralph,** J. **and Landucci, L.L.** J. **Org. Cha., 48, <sup>3884</sup>** ( **1983** ) .
- **15. Ralph, J.. Landuccl. L.L., Mlcholson, B.K., and Yilkins, a.L. J. Org.** *char.. 2.* **3337 (1984).**
- **16. Ralph, J. and Ad-. 8.8.** J. **Wod Cha. Technol., 3 <sup>183</sup> (1983).**
- **17. Glerer.** J. **Svensk Papperstid..** *73.* **571 (1970).**
- **18. Gierer, J.. Lindeberg, 0.. and Noren. I. Holzforschung.**  Gierer, J., Lind<br>Gierer, J., Lind<br><u>33</u>, 213 (1979).
- **19. Doddrell, D.H., Pegg. D.T.. and Bendall, H.R. J. Hag. Res., 48. 323 (1982).**
- *20.*  **Doddrell, D.H.. Pegg, D.T., Brooks, V., and Bendall. H.R.**  *J. An. Chem. Soc., 103, 727 (1981).*
- **21. ~izpurua.** J.U. **and Palm.** *C.* **Tetrahedron Lett,, a, 475 (198s).**
- **22.**  *~san,* **P. Tetrahedron Lett.,** *23.* **184s (1982).**
- **23. mrpor, s.** J. **Org.** *cher.,* **a. 250 (1963).**
- **24. Berndtsson, I., Khanna, B. and Lundquist, It. Acta Cham. -and., BH, 453 (1980).**
- **25. Watsubo. P. and Hlguchi. T. Holzforschung,** *29,* **193**  ( **19711).**