Communications to the Editor 5981

Si 
$$K^+$$
 $O^ A_{\alpha}^{H} = 7.33 (2 H), 1.40 (2 H) G$ 

 $a_{R}^{H} = 0.48 (2 \text{ H}) \text{ G}$ 

g = 2.00494

silacyclononane-1,2-semidiones as the ion-paired (3) and free ion (4), respectively.

## References and Notes

(1) Aliphatic Semidiones. 37.

- G. A. Russell, G. Wallraff, and J. L. Gerlock, J. Phys. Chem., 82, 1161
- With Li+ only the cis ion pair can be detected, whereas with Cs+ only the trans semidione is observed: G. A. Russell, D. F. Lawson, H. L. Malkus, R. D. Stephens, G. R. Underwood, T. Takano, and V. Malatesta, J. Am. Chem. Soc., 96, 5830 (1974).
- (4) H. C. Heller, J. Am. Chem. Soc., 86, 5346 (1964); C. Corvaja, P. L. Nordio, and G. Giacommeti, Ibid., 89, 1751 (1967).
- Prepared by the disproportionation of  $\alpha$ -hydroxycyclopentanone in Me<sub>2</sub>SO in the presence of K<sup>+</sup>CH<sub>3</sub>SOCH<sub>2</sub><sup>-</sup>. To obtain the spectra in the presence of Li<sup>+</sup>, Na<sup>+</sup>, or (C<sub>2</sub>H<sub>5</sub>)<sub>A</sub>N<sup>+</sup> the K<sup>+</sup> salt of the semidione was treated with a slight excess of [2.2.2] cryptand to give the free semidione ion to which the new counterion was added as the iodide or perchlorate salt.
- (6) Experiment performed by Mr. B. Graether.
   (7) a<sup>K</sup> is not observed for 1,2-semidiones even in the presence of crown ethers (which for Na<sup>+</sup> greatty reduces the rate of the exchange; semidione<sup>-</sup>, Na<sup>+</sup> + \*Na<sup>+</sup> = semidione<sup>-</sup>, \*Na<sup>+</sup> + Na<sup>+</sup>).<sup>2</sup> a<sup>K</sup> can be observed in Me<sub>2</sub>SO for semidione radical dianions (unpublished work with Mr. T. Takano), such as that shown.

$$cH_3$$
 $C=C$ 
 $C=C$ 
 $K^+$ 
 $C=C$ 
 $C=C$ 

- $a_{\rm CO}^{\rm C}$  in *cis*-dimethylsemidione increases from 1.1 (K<sup>+</sup>, Me<sub>2</sub>SO) to 1.4 (Na<sup>+</sup>, Me<sub>2</sub>SO), 1.8 (Li<sup>+</sup>, Me<sub>2</sub>SO), 1.9 (K<sup>+</sup>, *tert*-butyl alcohol). In Me<sub>2</sub>SO  $a^{\rm H}$  increases from 7.0 (K<sup>+</sup>) to 7.1 (Na<sup>+</sup>) to 7.4 (Li<sup>+</sup>).<sup>3</sup> G. A. Russell and E. T. Strom, *J. Am. Chem. Soc.*, **86**, 744 (1964).
- (10) National Science Foundation Fellow, 1975-1978

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## Conversion of Allyl Alcohols to Dienes by Sulfoxide and Selenoxide Syn Elimination. Synthesis of PCB Arene Oxides

Sir:

Of the many procedures and reagents available for the dehydration of alcohols, few work well for the conversion of allyl alcohols to dienes, and even fewer provide for regiospecific dehydration. We report here a new procedure (Scheme I, Y = S, Se) for the 1,4 dehydration of allyl alcohols, and application of this method as a key step in the synthesis of chlorinated and brominated biphenyl oxides.

Treatment of alcohol 1a with 2-nitro-4-methylbenzeneselenenyl chloride in refluxing dichloromethane leads to diene

Table I. 1,4-Dehydration of Allyl Alcohols Using 2,4-Dinitrobenzenesulfenvl Chloride-Triethvlamine

Dimi	roochizehesanen ji em	oride Triethylanine	
run	allyl alcohol	diene	yields, %ª
1	<u>la</u>	<u>Za</u>	83 <sup>b</sup>
2	15	<u>2b</u>	66 <sup>b</sup>
3	le	<u>2</u> ç	79 <sup>b</sup>
4	CH OH	Ph	68 <sup>C</sup>
5	ОН	Ph	77 <sup>C</sup>
6	HO	Ar C61	1 <sub>5</sub> 71°
7	$\cup$		1 <sub>2</sub> -C <sub>6</sub> H <sub>3</sub> <10 <sup>C</sup>
8	<b>√</b> ОН	70 Ar C <sub>6</sub> 1	i <sub>s</sub> 75 <sup>b</sup>
9			2-C6H3 58b
10	OH	Ph	<sub>74</sub> b
11	Ph	Ph	73 <sup>b</sup>
12	Ph OH	Ph	54 <sup>c</sup> ,d

<sup>a</sup> Reactions were carried out on 1-10 mmol of allyl alcohol; products were purified chromatographically or by distillation. b Reaction in CH<sub>2</sub>Cl<sub>2</sub> at 25-40 °C. c Reaction in ClCH<sub>2</sub>CH<sub>2</sub>Cl at 80 °C. d The diene was a 60:40 mixture of E,E and E,Z isomers.

#### Scheme I

2a (65% yield). The reaction must proceed by [2,3]-sigmatropic rearrangement of selenenate ester 3 to the allyl selenoxide 4, which fragments to diene. There are several reported examples of allyl selenoxide to selenenate rearrangements, 2,5 but this is the first demonstration of the allyl selenenate to selenoxide conversion.

In the course of attempting to generalize the process described above a number of readily available selenenyl (PhSeCl, 2,4-(NO<sub>2</sub>)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>SeBr) and sulfenyl (CCl<sub>3</sub>SCl, 2-NO<sub>2</sub>C<sub>6</sub>H<sub>4</sub>SCl, 2,4-(NO<sub>2</sub>)<sub>2</sub>C<sub>6</sub>H<sub>3</sub>SCl) halides were examined. It was hoped that the powerful electron-attacting substituents in the sulfenyl halides, together with the acceleration provided by the allylic nature of the sulfoxide intermediate, would lead to sufficiently fast sulfoxide syn elimination<sup>6</sup> to accommodate the mild reaction conditions desirable for the synthesis of reactive dienes. 2,4-Dinitrobenzenesulfenyl chloride is the reagent currently showing the most promise. Of a variety of reaction conditions tried, treatment with 2.4 equiv<sup>7</sup> of the sulfenyl chloride and 3 equiv of triethylamine in dichloromethane (-30 to 40 °C) or 1,2-dichloroethane (20 to 80 °C) gave the highest yields. Table I presents several examples using these conditions.8 At its current stage of development, the reaction proceeds in acceptable yield for a variety of allyl alcohols. One exception is run 7 in Table I, for which the sulfenate-sulfoxide equilibrium may be particularly unfavorable (note, however, that the less hindered alcohol in run 6 works satisfactorily). The sulfenate esters apparently can undergo decomposition reactions if the electrocyclic reactions are too slow.

Runs 3, 4 and 5 demonstrate that all of the isomeric phenylcyclohexadienes can be prepared without loss of regiospecificity. The dienes prepared in runs 8 and 9 were especially susceptible to aromatization and isomerization. Optimum yields were obtained when sulfenate ester formation was performed at -30 °C (3 h), followed by warming to complete the reaction.

The diene prepared in run 1 is a key intermediate in the synthesis of several arene oxides derived from polychlorinated biphenyls (PCB's). 2,5,2',5'-Tetrachlorobiphenyl has been extensively studied as a model for the toxic, mutagenic, and carcinogenic effects of PCB mixtures. 10 The arene oxide 5a has been implicated as an intermediate during the metabolic degradation of this tetrachlorobiphenyl in rhesus monkeys.10a

The allyl alcohol 1a was prepared in 83% yield by addition of unstable 2,5-dichlorophenyllithium<sup>11</sup> to cyclohexenone. Conversion to diene 2a (Scheme I) and epoxidation gave the

intermediate epoxide 6. This compound could be monobrominated (22% yield of crystalline 7b, mp 145-146 °C), dibrominated (50% yield of crystalline 7c, mp 146-147 °C, based on 2a), and tribrominated under progressively more vigorous conditions using N-bromosuccinimide. The desired tetrachlorobiphenyl oxide 5a<sup>13</sup> was prepared in highest purity by conversion of 7c to a mixture of isomeric dichlorides 7a (R<sub>4</sub>N<sup>+</sup>Cl<sup>-</sup>, CH<sub>3</sub>CN) followed by chlorination (t-BuOCl,  $CCl_4$ ,  $h\nu$ ) and dehydrochlorination (DBU,  $CH_2Cl_2$ , 25°). The compound showed remarkable chemical stability. It could be chromatographed on silica gel with no detectable decomposition and solutions in methanol and dimethyl sulfoxide had half-lives of 20 and 50 days, respectively, at 25 °C.14

The structure of 5a was demonstrated by conversion to the phenols 8 and 9, whose substitution pattern was unambiguously assigned by analysis of the 270-MHz NMR spectra. 15

The arene oxides 5c 5d, and 5e were similarly prepared by dehydrobromination of the appropriate bromide.

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- (7) The sulfenic acid generated by syn elimination apparently reacts with 1 equiv of sulfenyl chloride.
- (8) The allyl alcohol starting materials were prepared by addition of organometallic reagents to enones (runs 1–3, 8–10), reduction of enones (runs 4, 5), alkylation of the lithium reagent prepared by deprotonation of phenyl prenyl selenide followed by oxidation<sup>2a</sup> (run 11), treatment of aldehyde with α-lithioselenoxide (run 12),9 or acid-catalyzed rearrangement of allyl alcohols (runs 6, 7).
- conois (runs 6, 7).

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  (13) 5a was purified by TLC on silica gel and crystallization: mp 75-76 °C; 38% yield from 7a; NMR ( $\delta$ , CDCl<sub>3</sub>) 4.23 (dd, J = 3.9, 2.6 Hz, 1 H), 4.31 (d, J =3.9 Hz, 1 H), 6.34 (d, J = 2.6 Hz, 1 H), 7.21 (dd, J = 2.4, 0.4 Hz, 1 H), 7.29(dd, J=8.6, 2.4 Hz, 1 H), 7.37 (dd, J=8.6, 0.4 Hz, 1 H). Anal. Calcd C<sub>12</sub>H<sub>6</sub>Cl<sub>4</sub>O: C, 46.79; H, 1.96. Found: C, 46.73; H, 2.06. If crude dibromide 7a was used, 5a could be prepared in 36% yield starting from diene 2b.
- (14) The monochlorobenzene oxides also show increased stability toward aromatization (Selander, H. G.; Jerina, D. M.; Piccolo, D. E.; Berchtold, G. A. J. Am. Chem. Soc., 1975, 97, 4428).
- (15) Phenol 8 was isolated in pure form: mp 101–102 °C; NMR ( $\delta$ , acetone- $d_6$ ) 3.85 (br s, 1 H), 7.20 (d, J = 0.4 Hz, 1 H), 7.33 (d, J = 0.4 Hz, 1 H), 7.38 (dd, J = 2.6, 0.4 Hz, 1 H), 7.45 (dd, J = 8.5, 2.6 Hz, 1 H), 7.52 (dd, J = 8.5, 0.4 Hz, 1 H). Anal. Calcd. for C<sub>12</sub>H<sub>6</sub>Cl<sub>4</sub>O: C, 46.79; H, 1.96. Found: C, 46.71;
- (16) A. P. Sloan Fellow, 1975-1979.

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# Intramolecular Ligand Reorganization in Five-Membered-Ring Tetraoxyselenuranes

Sir:

The intramolecular ligand reorganization of phosphoranes has received much attention, that of sulfuranes considerably less, and reorganization of selenuranes hardly any at all.<sup>3</sup> Paetzold and Reichenbacher<sup>4</sup> have prepared a number of tetralkoxyselenuranes. In the case of the tetramethoxy compound, the 1H NMR spectrum indicates that all of the hydrogens of the methoxy groups are equivalent. It was suggested that rapid intramolecular exchange may account for this observation.5