((Trimethylsilyl)methyl)-1,4-benzoquinones. Generation and Trapping of o-Quinone Methides

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In this communication we report the conversion of ((trimethylsilyl)methyl)-1,4-benzoquinones 1 to reactive electrophilic o-quinone methides 3, a transformation that takes place when 1 is treated with various nucleophilic species under neutral conditions (Scheme I). This unusual reaction is envisaged to involve nucleophilic attack on the trimethylsilyl group of 1 to give the corresponding vinylogous enolate anion 2 which leads to o-quinone methide 3 upon O-protonation. These reactive intermediates then proceed to hydroquinones 4 or related products via Michael additions or cycloadditions to the enone moiety. To our knowledge, no direct precedent for the conversion of 1 to 3 has previously been reported. This is of particular interest since the chemistry of o-quinone methides of structural type 3, i.e., those having a hydroxyl or alkoxy group in conjugation with the methylene group, have not received extensive attention.2

Besides synthetic applications, the electrophilic character of o-quinone methides of structural type 3 has important biological implications. Specifically, such intermediates have been suggested as alkylating agents generated in vivo from appropriately substituted quinones in a bioreductive activation process.3 The results presented here compliment evidence that such intermediates are electrophilic alkylating agents.4

Synthesis of the ((trimethylsilyl)methyl)quinones 8 and 9 rests upon the previously reported ring expansion of cyclobutenones to quinones and related aromatic compounds (Scheme II).5-7 These specific examples were prepared starting with the cyclobutenediones 5a,b, which were converted to 6a,b and 7a,b, respectively, upon treatment with 1-lithio-3-(trimethylsilyl)propyne and 2-lithio-3-(trimethylsilyl)propene in THF at -78 °C.8 Thermolysis of **6a** and **6b** in p-xylene at reflux gave the respective quinones 8a and 8b in good yields. Analogously, thermolysis of 7a and 7b gave the corresponding hydroquinones which were

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Scheme I

Scheme II

Reagents: (a) 1-Lithio-3-(trimethylsilyl)propyne,THF, -78 °C; (b) 2-Lithio-3-(trimethylsilyl)propen, THF, -78 °C; (c) ρ -Xylene, reflux; (d) Ag₂O, ρ -Xylene, rt.

Scheme III

a:R=R'=OMe

63-82%

Scheme IV

converted directly to the quinones 9a and 9b upon oxidative workup $(Ag_2O/p$ -xylene).

A set of reactions illustrating the intermediacy of the o-quinone methides and their ease of formation is outlined in Scheme III. When ethanolic solutions of the quinones 10 were heated at reflux for 0.5-6 h and the reaction worked up under oxidative conditions,

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the corresponding ethoxymethyl derivatives 13 were isolated (63-82%) along with minor amounts (5-7%) of the symmetrical dimer 14. Interestingly, these reactions also proceed at ambient temperatures but at a slower rate. For example, a solution of 10a in absolute ethanol stirred at room temperature for 24 h resulted in 13a and 14a in 45% and 22% yields, respectively.

These data agree with the proposed o-quinone methides 11 as intermediates. In this regard, the alkoxyquinones are envisaged to arise via Michael addition of ethanol to the enone followed by oxidation of the resulting hydroquinone.10 The dimer 14 could result from a number of possible pathways, but an attractive possibility involves Diels-Alder dimerization of the o-quinone methides 11 to the corresponding spiropyrans. 11 The symmetrical dimer, in its half-reduced oxidation state, would then result directly from an intramolecular elimination. Subsequent oxidation of this would then give the symmetrical diquinone dimer 14.

Further evidence for the o-quinone methide intermediate comes from the observations that 10a gave the hydroquinone 12a (85%) in refluxing glacial acetic acid/sodium acetate and that this hydroquinone gave a similar product distribution as observed for 10a when subjected to refluxing ethanol. Thus, the conversion of 10a and 12a to 13a and 14a strongly suggests the o-quinone methide 11a as a common transient intermediate.

Additional studies illustrating the synthetic utility of the ((trimethylsilyl)methyl)quinone/o-quinone methide conversion are given in Scheme IV. Thermolysis of 10a in 5% aqueous acetonitrile followed by oxidative workup afforded three products: the alcohol 15 (12%), the xanthen derivative 16 (22%), and the ethylene dimer 14a (46%). The xanthen derivative is viewed as arising from the Diels-Alder cycloaddition of the quinone 10a with the o-quinone methide 11a.12 Interestingly, the amount of water had a dramatic effect on the product distribution. For example, when 10% rather than 5% aqueous acetonitrile was employed, a 67% yield of 14a and 18% of 15 were isolated. Only traces of 16 were detected. These results are consistent with a mechanism in which water functions as the reagent to induce o-quinone methide formation (Scheme I, NuH = H_2O).¹³ At low water concentration (5%), the concentration of the o-quinone methide is also low, and thus it is easily intercepted by an unreacted quinone in a Diels-Alder cycloaddition to give 16. As the water concentration is increased (10%) so follows the o-quinone methide concentration, and dimerization as described previously takes place to give 14a.

Finally, an example of Diels-Alder trapping of the o-quinone methide is illustrated by the synthesis of 17. That is, generation of the o-quinone methide in 10% aqueous acetonitrile in the presence of excess *n*-butyl vinyl ether gave the chromanol 17 in 72% yield.14,15

In conclusion, the most significant aspects of this study include the following: (a) ((trimethylsilyl)methyl)-1,4-benzoquinones 1 function as excellent precursors to o-quinone methide intermediates 3; (b) the o-quinone methides 3 do not undergo tautomerization, which suggests that ketonization is slower than nucleophilic addition; (c) the position of the trimethylsilyl group in 1 dictates the specific quinone methide formed; and (d) the conversion of

1 to 3 can be accomplished under mild and neutral conditions and thus allows the synthetic utilization of these reactive electrophilic intermediates.

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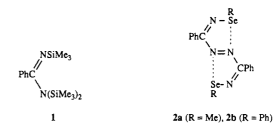
Supplementary Material Available: A table of spectroscopic data (IR, MS, ¹H and ¹³C NMR) for compounds 6-17 (2 pages). Ordering information is given on any current masthead page.

Synthesis and Structure of Azo Dyes with Short, Intramolecular Selenium-Nitrogen Contacts

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Recently there has been considerable interest in the synthesis and structures of both organic and inorganic selenium-nitrogen (Se-N) compounds. Some important milestones include the structural characterization of (a) dimers of the cyclic seven- π electron radicals 1,2,4,6-selenatriazinyl, (Ph₂C₂N₃Se)₂, 1,2,3,5diselenadiazolyl, $(PhCN_2Se_2)_2$, and 1,2,4-triselena-3,5-diazolylium, $(Se_3N_2^{\bullet+})_2$, and (b) metal complexes of the acyclic, binary Se-N anions, Se_3N^- and $Se_2N_2^{-4.5}$ We have prepared eightmembered 1,5-P₂VN₄Se₂ rings by the cyclocondensation reaction of Ph₂P(NSiMe₃)[N(SiMe₃)₂] with organoselenium trichlorides.⁶ In an attempt to obtain the unknown 1,5-C₂N₄Se₂ ring, we have found that the reaction of PhC(NSiMe₃)[N(SiMe₃)₂] (1) with RSeCl₃ (R = Me, Ph) unexpectedly produces the intensely colored diazenes 2a and 2b rather than the expected eight-membered ring. The X-ray structure of 2a reveals a nearly planar structure in which the hypervalent selenium atoms are connected by short, intramolecular contacts (ca. 2.65 Å) to one nitrogen of the azo group. A better synthesis of 2b (and its sulfur analogue) involves the reaction of 1 with 3 mol of PhEC1 (E = S, Se).



The dark red azo compound 2a was obtained in 44% yield by the slow addition of 1 to an equimolar amount of MeSeCl₃ in acetonitrile at 23 °C.7,8 An X-ray structural analysis confirmed

⁽¹⁰⁾ Subjecting 10a to isopropyl alcohol (10 h, reflux) furnished the corresponding isopropoxymethylquinone and 14a in 67% and 10% yields, respectively (oxidative workup). Analogously, the reaction of 10a in tert-butyl alcohol (44 h) gave 45% of the tert-butoxymethylquinone and 22% of 14a.

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⁽¹³⁾ Heating 10a in dry benzene at reflux for 90 h gave >98% of recovered

starting material.
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