Synthesis and characterization of 2,6-di-*tert***-butyl-1-thio-1,4-benzoquinone, the first isolable monothio-1,4-benzoquinone**

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The title compound, a labile but isolable substance owing to appreciable steric protection, shows substantially higher electron affinity than the corresponding 1,4-benzoquinone and is reduced by water.

While p -benzoquinones have been extensively studied, 1 little has been known about the corresponding sulfur analogues. A major reason may be the general lability of thiocarbonyl compounds that makes their handling and isolation in pure form difficult.2 Bock and co-workers have reported the pyrolytic generation and spectroscopic characterization of the parent dithio- and monothio-1,4-benzoquinones, **1** and **2**, in low temperature matrices.3 Recently 2,6-dialkyl-4-thio-1,4-benzoquinones **3** have been generated and trapped by cycloaddition with 1,3-dienes.⁴ The only isolable, related monothioquinone so far known is monothioanthraquinone **4**.5 We have recently reported the synthesis and properties of thioquinone methides **5** in which *tert*-butyl groups give steric protection to a considerable extent.6 Here we report the synthesis and characterization of 2,6-di-*tert*-butyl-1-thio-1,4-benzoquinone **6**, the first isolable, monothio-1,4-benzoquinone.

We have already described the generation of lithium 2,6-di*tert*-butyl-4-lithiothiophenoxide **7** as the key synthon for **5**.6 The exposure of a THF solution of 7 to O_2 at -70 °C followed by the usual work-up afforded bisphenol disulfide **8**† in 50% yield. A major byproduct was 2,6-di-*tert*-butylthiophenol, a reduction product. Obviously, the two *tert*-butyl groups at the *ortho*-positions are not bulky enough to hinder the coupling of the intermediate thiophenoxy radical as reported for the 2,6-di*tert*-butylthiophenoxy radical itself.7 The initially expected mercaptophenol **9**† was obtained by reduction of **8** with zinc powder in acetic acid (60%). The attempted oxidation of **9** to **6** with DDQ in benzene or acetone resulted, however, in the formation of **8**. We therefore expected that oxidation of **8** to its bis(phenoxy radical) **10** would cause cleavage of the rather weak disulfide bond, leading to the generation of **6**. In fact, NMR monitoring of a mixture of **8** and excess DDQ (3.5 equiv.)

in deaerated benzene- d_6 in a sealed tube at room temperature gave rise to new proton signals as singlets at δ 1.14 and 6.48 and a ¹³C signal at δ 233.45 (C=S) which are reasonable for the desired **6**. However, in spite of the use of excess DDQ, the reaction seemed to form a 2+1 equilibrium mixture of **6** and **8** probably by mediation of DDQ and its hydroquinone. Although the signals of **6** remained almost unchanged in the solution for at least two months, the attempted isolation of **6** from the mixture failed because of its lability. This failure suggests that the isolation of **6** from solutions should be difficult. In the end, the isolation of **6** in almost pure form was achieved in 70% yield by its sublimation to a water-cooled cold finger from a solidstate mixture of **8** and PbO₂ (1:30 w/w) heated at 140 °C under vacuum.

The monothiobenzoquinone **6**† thus obtained is a greenish yellow solid. Although relatively stable in the solid state as well as in neutral aprotic solvents such as benzene and acetone at room temperature, **6** is sensitive to moisture and other protic substances. The IR spectrum of **6** shows an intense carbonyl absorption at 1639 cm⁻¹ and a less intense thiocarbonyl absorption (intense in the Raman spectrum) at 1141 cm^{-1} . This thiocarbonyl stretching frequency is considerably lower than that of $4(1212 \text{ cm}^{-1})^5$ and even lower than the significantly dipolar $4H$ -pyran-4-thione (1168 cm⁻¹).⁸ The electronic spectrum of 6 exhibits a weak absorption ($\varepsilon = 32$) at 758 nm, which is assignable to the $n \rightarrow \pi^*$ transition of the thiocarbonyl group, as well as a strong absorption at 321 nm. The visible absorption of **6** is at an appreciably longer wavelength than those of **2** (500 nm)³ and $4(697 \text{ nm})$.⁵ The IR and visible absorption spectral results suggest a substantial polarization of the thiocarbonyl group of **6**. In this context, a semiempirical theoretical calculation (PM3) predicts that **6** should take the boat form, due to steric congestion, with bow and stern angles of 50° and 27°, respectively. This molecular deformation from planarity could be responsible for the enhanced polarization (elongation) of the thiocarbonyl bond.

Thiocarbonyl compounds are usually more easily reduced than the corresponding carbonyl compounds.9 Upon cyclic voltammetry, **6** shows two irreversible reduction waves at -0.60 and -1.33 V (both peak potentials *vs*. Ag/Ag⁺, in 0.1 M $n-Bu_4NClO_4-CH_3CN$, ferrocene/ferrocene+ = 0.10 V). These reduction potentials are about 0.5 V lower than those of 2,6-di*tert*-butyl-1,4-benzoquinone **11** (the corresponding peak potentials: 1.07 and 1.88 V) measured under the same conditions, indicating that **6** is a substantially strong electron acceptor. Surprisingly, **6** was found to be reduced rather than hydrolysed by water: when a small amount of D_2O was added into an acetone solution of **6** either in the dark or light at room temperature, the signals of **6** at δ 1.37 (s) and 6.46 (s) disappeared within 1 h with the concurrent appearance of new signals at δ 7.00 (s), 6.93 (br s), 6.73 (br s) and 6.52 (s) in the olefinic to aromatic region. The products were identified to be monothiohydroquinone **9**, disulfide **8** and benzoquinone **11**, formed in a $45:50:5$ ratio (¹H NMR and TLC comparison with those of the authentic samples). The formation of **8** (probably also **9**) points to the intermediate formation of the corresponding thiophenoxy radical. In addition, the ferricyanide test as a preliminary qualitative test afforded a positive result, conforming the formation of hydrogen peroxide in the solution. Thus, hydrolysis of **6** to **11** is a very minor process, different from the usually easy hydrolysis of thioketones to ketones.

In conclusion, although the two *tert*-butyl groups at the *ortho* positions are not bulky enough for steric protection, 2,6-di-*tert*butyl-1-thio-1,4-benzoquinone **6** was synthesized as a labile but isolable compound at around room temperature and was found to show a considerably high electron affinity and to undergo an unusual reaction with water. We are now investigating the detailed chemical properties of **6**.

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Notes and references

† *Selected physical and spectroscopic data*; **6**: mp 87–88 °C; MS (EI) *m/z* $238 (M^+ + 2H, 100\%)$, $236 (M^+, 20)$, $221 (M^+ - CH_3, 27)$, $180 (M^+ - C_4H_8)$ 25), 165 (M⁺ - 71, 69); IR: v (KBr)/cm⁻¹ 2963 (s), 1639 (s, C=O), 1556 (m), 1456 (m), 1366 (m), 1302 (m), 1261 (m), 1232 (m), 1200 (m), 1141 (m, C=S), 1096 (m), 1072 (m), 1022 (m), 910 (m), 801 (m); $\delta_{\rm H}$ (400 MHz, C_6D_6) 1.13 (s, 18H), 6.49 (s, 2H); $\delta_C(100 \text{ MHz}, C_6D_6)$ 30.98, 36.87, 121.94, 163.58, 189.94 (C=O), 233.45 (C=S); UV–Vis (cyclohexane) $\lambda_{\text{max}}/ \text{nm}$ (ε) 321 (14 300), 456 sh, 758 (32); **8**: mp 210–213 °C; MS (EI) *m/z* 474 (M+, 12%), 238 (M⁺/2 + 1, 100), 181 (M⁺/2 - C₄H₉, 16); δ_H (270 MHz, CD₂Cl₂) 1.04 (s, 18H), 1.60 (s, 18H), 4.84 (br s, 2H, OH), 6.65 (s, 2H), 6.86 (s, 2H); The NMR spectra show restricted rotation of the S–S or C–S bond with an estimated energy barrier of ΔG [‡] = 16.5 ± 0.2 kcal mol⁻¹ (T_c = 75 °C in benzene); see also ref. 10; **9**: mp 124–127 °C; MS (EI) *m/z* 238 (M+, 100%), 181 (M⁺ - C₄H₉, 17); $\delta_H(270 \text{ MHz}, \text{CDCl}_3)$ 1.58 (s, 18H), 3.24 (s, 1H), 4.57 (s, 1H), 6.89 (s, 2H).

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