Direct Detection of the Radical Cation of 2,6-Di-tert-butyl-4-methylphenol Generated by Electron-Transfer Oxidation with Matrix Alkyl Halide Cation Radicals. A Low-Temperature EPR and UV–VIS Optical Absorption Study†

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Electron-transfer oxidation of 2,6-di-tert-butyl-4-methylphenol (ArOH, 1) by matrix (RHal) radical cations (RHal = Freon-113, BuCl and BuCl) results in the formation of the radical cation ArOHH+ (2) as a primary intermediate directly observed during EPR and UV–VIS optical absorption measurements at 77 K. In the glassy matrix of BuCl the radical cation ArOHH+ exhibits a broad absorption band with λmax = 450 nm (ε/dm3 mol−1 cm−1 = 2.1 × 104). The EPR spectrum of ArOHH+ at 77 K is characterised by isotropic splitting from the freely rotating CH3 group and anisotropic splitting from the hydrogen atom in the OH group. Upon warming in the temperature range 77–103 K, deprotonation of ArOHH+ and deuterium-substituted analog ArODH+ resulting in the formation of the 2,6-di-tert-butyl-4-methylphenoxyl radical ArO• (4), was directly observed by EPR and optical absorption measurements in all matrices studied.

The antioxidative properties of (sterically hindered) phenols can be understood when it is considered that phenols can act as electron and as hydrogen atom donors. Abstraction of a hydrogen atom from the OH group results directly in a phenoxyl radical.1 Under electron transfer oxidative conditions, in most cases phenoxyl radicals were also detected as the first observable transient,2 which can only be explained by suggesting a radical cation as a very unstable intermediate.

Because of the very low pK valuea,3 of phenol radical cations they could be observed in aqueous or polar solutions only under strongly acidic conditions. Thus, cation radicals of various substituted phenols were characterised by their steady-state solution EPR spectra in sulfuric acid solutions3 or in the presence4c of strong Lewis acids (AlCl3 in CH3NO2, or CH3Cl). In both cases3,4a-c radical cations are in a reversible protonation equilibrium with the phenoxyl radicals and their formation in the primary electron transfer oxidative step could not be proved. Cyclic voltammetry on some phenolsd in acidic media (CH3Cl with 10% FSO3H) has also provided evidence for the formation of radical cations.

In the case of phenols that contain para-substituents with a high affinity for protons, such as amino groups, photoionisation combined with optical matrix isolation spectroscopy has indicated the existencef,g of radical cations. Furthermore, p-aminophenol radical cations could also be identified in strongly acidic solution usingh time-resolved Raman resonance spectroscopy. The photolytic and radiolytic ionisation of phenol itself in argon matrices yielded optical absorption spectra which have been interpretedi to be due to PhOHH+ and PhO•.

To avoid structural peculiarities leading to isomerisation involving the ortho-position and charge localisation in the para-position, we studied the ionisation of sterically hindered phenols by means of pulse radiolysis. Electron transfer oxidation of some of these phenols in non-polar solutions at room temperature indicates that phenoxyl radicals are really subsequent products of very short-lived cation radicals.7 In order to gain deeper insight into the chemistry and structure of the primary oxidised...
intermediates we have performed a low temperature matrix isolation investigation of the radical cations derived from the sterically hindered 2,6-di-tert-butyl-4-methylphenol (ArOH, 1). The radical cations were generated by \( \gamma \)-radiolysis of ArOH dissolved in glass-forming matrices of CF\(_2\)CICF\(_2\)_2 (Freon-113), Bu'Cl and BuCl. Using two different detection techniques – EPR and UV–VIS optical absorption spectroscopy – we report here upon the characterisation of the radical cations ArOH\(^{+}\) (2).

![Chemical Structures](image)

**Experimental**

First-derivative X band EPR measurements and \( \gamma \)-irradiation (dose 5 kGy) of the frozen, degassed solutions were performed as described in our previous work.\(^8\) A copper cryostat was used for the annealing of the irradiated samples at various temperatures. After annealing (cf. Ref. 9) the samples for 10 min they were recooled to 77 K for EPR or UV–VIS optical absorption measurements. Absorption spectra in frozen glassy matrices of Bu'Cl at 77 K were measured for samples in Suprasil quartz cells (1 x 10 x 30 mm\(^3\)) using a Shimadzu UV–VIS Scanning Spectrophotometer UV-2101PC according to procedure outlined in Ref. 10(a). The experimental error in the absorbance measurements was estimated to be 10–15%.

![Chemical Structures](image)

**Results and discussion**

*Low temperature EPR studies.* Since the ionisation potential (\( E_i \)) of ArOH (ArOD) is lower\(^\ddagger\) than that of the matrices studied,\(^10\) \( \gamma \)-irradiation of ArOH in frozen halogenated matrices (RHal) is expected to lead to the formation of the radical cations ArOH\(^{+}\)\(^10,11\) as a result of an electron transfer from the solute molecules to the matrix radical cations RHal\(^{+}\) according to eqn. (1).

\[
\text{ArOH} + \text{RHal}^{+} \rightarrow \text{ArOH}^{+} + \text{RHal}
\]  

(1)

Secondary electrons generated in matrices during \( \gamma \)-irradiation at 77 K are completely scavenged by matrix molecules resulting in the formation of matrix-derived radicals as well as Cl\(^-\) anions.\(^10,11\) In the case of Freon-113, matrix-derived CF\(_2\)CICFCl\(^-\) radicals could not strongly contribute to the observed EPR signal because of the anisotropic line broadening.\(^11\)

Table 1 shows the hyperfine coupling constants for all investigated species. The EPR spectrum of irradiated ArOH (0.05 mol dm\(^{-3}\)) in a Freon-113 matrix [Fig. 1(a)] we assign to the radical cation ArOH\(^{+}\). This asymmetric, near to 1:3:3:1 quartet signal exhibits splitting from three equivalent protons in the CH\(_3\) group. The asymmetry of the EPR spectrum and the additional doublet features are due to the anisotropic dipole–dipole interaction of the unpaired electron on the 2p oxygen orbital with the H atom in the OH group (cf. Refs. 12).

![Chemical Structures](image)

**Chemicals:** 2,6-Di-tert-butyl-4-methylphenol (ArOH, 1) from Bayer (Leverkusen) was used as received. 2,6-Di-tert-butyl-4-methyl-1-deuteriophenol (ArOD, 3) was obtained from ArOH by means of a deuterium exchange reaction with excess C\(_2\)H\(_2\)OD. NMR spectral data confirmed more than 90% deuterium in ArOD. Freon-113 (puriss. Genetron® 113, Fluka), Bu'Cl and BuCl (both from Aldrich) were used as described in Ref. 8.

\(^\ddagger\) \( E_i \) for ArOH and ArOD are not known. In this work we assume that they have the same \( E_i \) (8.52 eV) as PhOH (see Ref. 6).

\(^\ddagger\) The hyperfine coupling constant for deuterium should be 6.514 times smaller\(^14\) and hence it contributes to the linewidth of the EPR spectrum.
Table 1. EPR parameters for ArOH*-, ArOD*+ and ArO* in Freon-113.

<table>
<thead>
<tr>
<th>Radical</th>
<th>$g_0$</th>
<th>$g_{iso}$</th>
<th>$a_{1/2}(\text{CH}_3)$</th>
<th>$a_{1/2}(\text{meta-H})$</th>
<th>$a_{1/2}(\text{OH})$</th>
<th>T/K</th>
<th>Ref.</th>
</tr>
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<tr>
<td>ArOH*+</td>
<td>2.0035</td>
<td>14.3</td>
<td></td>
<td></td>
<td>A$_h$ = 10.0</td>
<td>77</td>
<td></td>
</tr>
<tr>
<td>ArOD*+</td>
<td>14.3</td>
<td>0.89 (1 H)</td>
<td>3.72</td>
<td></td>
<td></td>
<td>77</td>
<td></td>
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<tr>
<td>ArOH*+</td>
<td>14.19</td>
<td>0.8 (1 H)</td>
<td>3.6</td>
<td></td>
<td></td>
<td>213</td>
<td>4(c)</td>
</tr>
<tr>
<td>ArO*</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td>77</td>
<td></td>
</tr>
<tr>
<td></td>
<td>11.1</td>
<td>1.70</td>
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<td></td>
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<td>298</td>
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</tr>
<tr>
<td></td>
<td>11.0</td>
<td>1.65</td>
<td></td>
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</tr>
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</table>

Fig. 1. EPR spectra of radical cations ArOH*+ (a) and ArOD*+ (b) in Freon-113 matrix compared with computer-simulated spectra (c) and (d), respectively. Deprotonation of both radical cations results in the phenoxyl radical ArO* (e). All spectra were recorded at 77 K.

2p orbitals of the oxygen atom as described by the mesomeric structures 2a, 2b and 2c. An increase (about 29%) of the methyl coupling constant $a_{1/2}(\text{CH}_3)$ in ArOH*+ and ArOD*+, in comparison with that for the corresponding phenoxy radical ArO* (Table 1), indicates increased spin density at the C-4 position of the aromatic ring in the radical cation. This is in line with the results for other substituted phenol radical cations. The $g$-factor for the radical cation ArOD*+ is smaller than that for the phenoxy radical ArO* (which is observed at lower magnetic field, see below), thus further supporting the identification of the radical cation ArOH*+ (cf. Ref. 3).

With $A_{||}(\text{H})$ and $g_{||}$ determined directly from the experimental EPR spectrum of ArOH*+, and assuming an axially symmetric anisotropic hyperfine tensor for the hydrogen atom of the OH group [cf. Ref. 12(b)], the EPR spectra of the ArOH*+ and ArOD*+ radical cations were computer-simulated in agreement with experiment [Figs. 1(c) and (d), respectively] using $A_\perp = 2\text{ G}$, $g_{\perp} = 2.0050$ and a linewidth of 2.2 G.

EPR measurements performed with ArOH in a BuCl matrix also clearly demonstrate the formation of ArOH*+. Although in this case both ArOH*+ and matrix-derived butyl radicals contribute to the EPR spectrum, the characteristic 1:3:3:1 quartet of ArOH*+ is clearly observed (see insert in Fig. 2). Similar results were obtained in a BuCl matrix (spectra not shown).

Upon annealing of irradiated matrices at 88 K, the intensity of the EPR signal from the radical cations diminished and the slightly asymmetric 1:3:3:1 quartet signal (shifted to low field) of the phenoxy radicals ArO* (4) grew, in all matrices studied. Two different paramagnetic species – ArOH*+ (ArOD*+) and ArO* – were simultaneously observed in the EPR spectra upon annealing in the temperature range up to 107 K. After 10 min annealing at 107 K only the phenoxy radical ArO* was observed in the EPR spectrum in both ArOH and ArOD samples, as shown in Fig. 1(e) for the Freon-113 matrix. This indicates the deprotonation of the radical cations ArOH*+ and ArOD*+ according to eqs. (2a) and (2b). The deprotonation is nearly quantit-
ative, based on the EPR measurements in the Freon-113 matrix.

\[
\text{ArOH}^- \rightarrow \text{ArO}^- + H^+ \quad (2a)
\]

\[
\text{ArOD}^- \rightarrow \text{ArO}^- + D^+ \quad (2b)
\]

Upon further warming, the phenoxyl radicals ArO\(^-\) decayed (more than 90\%) in all matrices studied, owing to their reactions with matrix-derived radicals and to reversible dimerisation\(^{36,38}\) into molecular metastable dimeric products. After melting the irradiated matrices, a weak EPR signal due to the phenoxyl radicals ArO\(^-\) (Table 1) could be observed at room temperature in the first few minutes.

Low temperature UV–VIS optical absorption measurements. In contrast with the pure matrices,\(^{10a}\) glassy matrices containing dissolved ArOH became intensely yellow-green in color upon irradiation. For the optical absorption measurements Bu\(^{15}\)Cl matrix was chosen as the matrix, due to the fact that upon being frozen it results in good glasses.\(^{10a}\) Fig. 2 (bold curve) shows the absorption spectrum immediately after irradiation of ArOH (0.05 mol dm\(^{-3}\)) in a glassy Bu\(^{15}\)Cl matrix. Based on the EPR results (above) we assign the broad absorption band with \(\lambda_{\text{max}} = 450 \text{ nm}\) and a shoulder at \(\lambda = 295 \text{ nm}\) (not shown) to the absorption spectrum of the radical cation ArOH\(^{+}\). This double-band absorption spectrum is quite similar to that of the phenol radical cation PhOH\(^{+}\) in a frozen argon matrix\(^{8}\) — a broad absorption band at \(\lambda = 420 \text{ nm}\) and a shoulder at 275 nm. The red shift of the ArOH\(^{+}\) absorption could be due to matrix and alkyl-substitution effects.

The changes in optical absorption upon warming of irradiated solutions parallel those in the EPR spectra. Two characteristic peaks\(^{9}\) of the phenoxyl radical ArO\(^-\) at \(\lambda_{\text{max}} = 401 \text{ nm}\) and 382 nm are clearly seen in the absorption spectrum taken after the decay of the radical cation ArOH\(^{+}\) upon warming up to 107 K (Fig. 2, dotted curve). This is in agreement with the EPR results described above.

Assuming that ArO\(^-\) resulted only from the deprotonation [see eqns. (2a) and (2b)] of the radical cations ArOH\(^{+}\), an estimate of the extinction coefficient (\(\varepsilon\)) for the radical cation ArOH\(^{+}\) could be made directly from the absorption spectra. Using\(^{10a}\) for ArO\(^-\) an \(\varepsilon\)-value at \(\lambda_{\text{max}} = 401 \text{ nm}\) of \(4.3 \times 10^3 \text{ dm}^3 \text{ mol}^{-1} \text{ cm}^{-1}\), this gives, for the broad absorption band of the cation radical ArOH\(^{+}\) at \(\lambda_{\text{max}} = 450 \text{ nm}\), \(\varepsilon = 2.1 \times 10^3 \text{ dm}^3 \text{ mol}^{-1} \text{ cm}^{-1}\). 

\(\uparrow\) The \(\varepsilon\)-value for ArO\(^-\) at \(\lambda_{\text{max}} = 401 \text{ nm}\) was calculated from the \(k/t\) ratio determined by Land and Porter [Ref. 2(b)] and the (extrapolated to room temp.) decay rate constant measured by Rüege and Fischer (Ref. 16). This value is in good agreement with that (4.13 \times 10^3 \text{ dm}^3 \text{ mol}^{-1} \text{ cm}^{-1}) measured\(^{17}\) for the analogous 2,4,6-trimethylyphenoxyl radical.

**Conclusions**

Electron transfer oxidation of simple hindered phenols such as ArOH and ArOD by matrix radical cations results in the formation of the radical cations ArOH\(^{+}\) and ArOD\(^{+}\) as primary intermediates directly observed during low-temperature EPR and UV–VIS optical absorption measurements. Deprotonation of ArOH\(^{+}\) and ArOD\(^{+}\) according to eqns. (2a) and (2b) yields phenoxyl radicals ArO\(^-\). This reaction was observed by EPR and optical absorption spectroscopy in different halogenated matrices in the temperature range 77–103 K.

Thus, electron transfer oxidation of phenols followed by deprotonation of the resultant radical cation is a two-step reaction route that explains the appearance of phenoxyl radicals under conditions excluding hydrogen abstraction from the OH group of phenol molecules. Both steps are of importance for phenol chemistry in biological systems as well as for the chemistry of sterically hindered phenolic antioxidants and stabilisers.

**References**


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