Oxidation of Tetraphenylborate by Hexachloroiridate(IV)

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Summary The reaction, $BPh_4^- + 2IrCl_6^{2-} + H_2O \rightarrow Ph_2BOH + Ph_2 + 2IrCl_6^{3-} + H^+$, has been shown to occur quantitatively in aqueous solution; the kinetics are interpreted in terms of a mechanism involving BPh_4^- radicals which are also proposed as intermediates in certain electrochemical and photochemical reactions of BPh_4^- .

We have found that BPh₄⁻ and IrCl₆²⁻ react quantitatively in aqueous solution in accord with equation (1). The stoicheiometry was confirmed by (i) spectral titrations which exhibited sharp end points at IrCl₆²⁻: BPh₄⁻ ratios of 2·0:1, (ii) the quantitative (>95%) isolation of Ph₂ (identified by m.p., ¹H n.m.r., and g.l.c.), and (iii) quantitative estimation of Ph₂BOH by ¹H n.m.r., as well as conversion of the latter into the corresponding anhydride, Ph₂BOBPh₂, which was isolated and characterized by its m.p. (115°) and i.r. spectrum.

$$\begin{array}{c} {\rm BPh_4^-} + 2{\rm IrCl_6^{2-}} + {\rm H_2O} \rightarrow \\ {\rm Ph_2BOH} + {\rm Ph_2} + 2{\rm IrCl_6^{3-}} + {\rm H^+} \end{array} \eqno(1)$$

The kinetics of the reaction were studied by monitoring the ${\rm IrCl_6}^2$ - concentration spectrophotometrically at 490 nm (ϵ 3·2 × 10³). Kinetic measurements, encompassing the initial concentration ranges $1\cdot2\times10^{-4}$ — $1\cdot5\times10^{-3}$ M-IrCl₆²-, $1\cdot1\times10^{-4}$ — $1\cdot5\times10^{-3}$ M-BPh₄-, and 0—0·1M-HClO₄, vielded the rate-law, —d[BPh₄-]/dt = k[BPh₄-][IrCl₆²-]

with $k = (1.2 \pm 0.3) \times 10^2 \, \text{l mol}^{-1} \, \text{s}^{-1}$ at 25° (independent of the H+ concentration), the ionic strength being maintained at 0.5M with NaClO₄.

The form of the rate-law, together with the virtually universal tendency of $IrCl_6^{2-}$ to react as a 1-electron outersphere oxidant, 1,2 favour the mechanism in equations (2)—(4), in which an intermediate BPh₄ radical, generated by the rate-determining electron-transfer step, undergoes a further (rapid) 1-electron oxidation to yield the observed products.

$$BPh_4^- + IrCl_6^{2-} \xrightarrow{k} BPh_4^{\cdot} + IrCl_6^{3-}$$
 (2)

$$BPh_{4} + IrCl_{6}^{2-} \xrightarrow{fast} BPh_{4} + IrCl_{6}^{3-}$$
 (3)

$$BPh_4^+ \xrightarrow{fast} > [BPh_2^+] + Ph_2$$

$$\downarrow H_2O \rightarrow Ph_2BOH + H^+$$

$$(4)$$

These results and conclusions are of interest in connection with several other aspects of boron chemistry. (1) Radical displacement reactions of boranes of the type, 3 R· + BR' $_3$ \rightarrow BR' $_2$ R + R'·, in which the possibility of neutral tetraco-ordinated intermediates of the type BR' $_3$ R· has previously been considered but not demonstrated. (2) The electrochemical oxidation of BPh $_4$ -, which has also been found to yield Ph $_2$ BOH and Ph $_2$ as products and which

has previously been interpreted in terms of a primary 2-electron oxidation to BPh₂+ and Ph₂.5 Our results suggest that an alternative mechanism, involving successive 1-electron steps with the intermediate formation of BPh₄·, is not unlikely. (3) The photochemical decomposition of the BPh₄- which yields Ph₂BOH and a mixture of phenylcyclohexadienes in the absence of oxygen, as well as biphenyl in the presence of oxygen. The photochemical observations, in conjunction with our chemical oxidation studies, can be interpreted in terms of a mechanism involving the photochemical electron-detachment step (5), followed by groundstate reactions of the BPh₄· intermediate. The fact that Ph₂ is formed in quantitative yield by chemical oxidation of

BPh₄- with IrCl₆²-, but in yields of only up to 77% (along with compensating yields of phenylcyclohexadiene) in the photochemical decomposition of BPh₄⁻ in the presence of air can be attributed to the (not unexpectedly) greater efficiency of $IrCl_6^{2-}$, compared with O_2 , in intercepting and oxidizing BPh₄· (equation 3).

$$BPh_4^- \xrightarrow{hv} BPh_4 \cdot + e^-$$
 (5)

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