Citation: Deary, Michael, Durrant, Marcus and Davies, Martin (2013) A kinetic and theoretical study of the borate catalysed reactions of hydrogen peroxide: the role of dioxaborirane as the catalytic intermediate for a wide range of substrates. Organic & Biomolecular Chemistry, 11. pp. 309-317. ISSN 1477-0520

Published by: Royal Society of Chemistry

URL: http://dx.doi.org/10.1039/C2OB26842F <http://dx.doi.org/10.1039/C2OB26842F>

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A kinetic and theoretical study of the borate catalysed reactions of hydrogen peroxide: the role of dioxaborirane as the catalytic intermediate for a wide range of substrates†

Michael E. Deary,*a Marcus C. Durrantb and D. Martin Daviesb

Our recent work has provided new insights into the equilibria and species that exist in aqueous solution at different pHs for the boric acid – hydrogen peroxide system, and the role of these species in oxidation reactions. Most recently, (M. C. Durrant, D. M. Davies and M. E. Deary, Org. Biomol. Chem., 2011, 9, 7249–7254), we have produced strong theoretical and experimental evidence for the existence of a previously unreported monocyclic three membered peroxide species, dioxaborirane, that is the likely catalytic species in borate mediated electrophilic reactions of hydrogen peroxide in alkaline solution. In the present paper, we extend our study of the borate–peroxide system to look at a wide range of substrates that include substituted dimethyl anilines, methyl-p-tolyl sulfoxide, halides, hydrogen sulfide anion, thiosulfate, thiocyanate, and hydrazine. The unusual selectivity-reactivity pattern of borate catalysed reactions compared with hydrogen peroxide and inorganic or organic peracids previously observed for the organic sulfides (D. M. Davies, M. E. Deary, K. Quill and R. A. Smith, Chem.–Eur. J., 2005, 11, 3552–3558) is also seen with substituted dimethyl aniline nucleophiles. This provides evidence that the pattern is not due to any latent electrophilic tendency of the organic sulfides and further supports dioxaborirane being the likely reactive intermediate, thus broadening the applicability of this catalytic system. Moreover, density functional theory calculations on our proposed mechanism involving dioxaborirane are consistent with the experimental results for these substrates. Results obtained at high concentrations of both borate and hydrogen peroxide require the inclusion the diperoxodiborate dianion in the kinetic analysis. A scheme detailing our current understanding of the borate–peroxide system is presented.

Introduction

Our recent work has provided new insights into the equilibria and species that exist in aqueous solution at different pHs when hydrogen peroxide and boric acid are mixed together in concentrations of up to 0.5 M borate and 2.0 M hydrogen peroxide.1–3 In addition, through density functional theory (DFT) calculations, we have determined the species that are likely to be responsible for observed catalysis of electrophilic reactions of hydrogen peroxide with dimethyl sulfide and substituted phenyl methyl sulfides.3

Fig. 1 details our current understanding of the hydrogen peroxide–borate system and Table 1 lists the corresponding formation constants; the nomenclature has been retained and expanded, from our previous work.1–3 It is well known that both boric acid, 1, and peroxyboric acid, 3, which are both present at low pH, act primarily as Lewis acids, however our recent DFT study has shown that they are also likely to exhibit a small degree of Brønsted acidity,3 sufficient for the respective anions, structures 4 and 5, to exist in solution at low concentrations. The peroxyborate anion, 5, has a tautomeric form, 8, a previously unreported monocyclic three membered peroxide species named dioxaborirane.3 It is this species that we have proposed as being responsible for the catalysis that is observed for borate mediated electrophilic reactions of hydrogen peroxide with sulfides in alkaline conditions,3 rather than the previously assumed monoperoxoborate, 6, and diperoxoborate, 7, anions, that dominate in alkaline solution, the latter when there are high ratios of hydrogen peroxide to borate. Dioxaborirane, which is in equilibrium with monoperoxoborate via the addition of a water molecule, has a particularly low energy barrier for the reaction with dimethyl sulfide (ca. 2.8 kcal mol⁻¹), though it is likely to be present in only small
concentrations. For the same reaction monoperoxoborate has an activation barrier of 17.5 kcal mol\(^{-1}\), compared to 10.1 kcal mol\(^{-1}\) for the uncatalysed reaction of hydrogen peroxide, hence monoperoxoborate is unlikely to be catalytically active. The low activation barrier for the dioxaborirane is likely to be due to the intrinsic reactivity of the three membered ring, as has been reported elsewhere for the carbon analogue of dioxirane, together with the lack of any steric hindrance in the transition state. At low pH, peroxoboric acid, 3, with an activation barrier of 7.4 kcal mol\(^{-1}\), is a plausible catalyst, but it has a low formation constant and so the observed rate enhancement for its reaction with dimethyl sulfide is small. At higher ratios of borate to hydrogen peroxide, the unreactive peroxodiborate dianion, 9, is formed, and in this paper we adduce evidence that this species can complex with an additional hydrogen peroxide molecule to form the diperoxodiborate dianion, 10.

Finally we should note that a large number of polyborate species have been proposed to exist at higher borate concentrations, though literature studies have shown that the trimer, 11 and the tetraborate ion, 12 are the major species present. In addition, we cannot rule out the existence of polyperoxyborates: there is no reason why species 11 and 12 should not complex with hydrogen peroxide in an analogous way to boric acid, 1.

At alkaline pHs the predicted high reactivity of dioxaborirane is consistent, under the Hammond Postulate, with its low selectivity (taken as the Hammett \(\rho\) value, \(\rho = -0.65\), for its reaction with a range of the sulfides) when compared with other hydroperoxides such as hydrogen peroxide (\(\rho = -1.5\)).

### Table 1 Formation constants for the equilibria shown in Fig. 1

<table>
<thead>
<tr>
<th>Formation constant</th>
<th>Value</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>(K_{BOH})</td>
<td>(1.0 \times 10^{-9}) M</td>
<td>14</td>
</tr>
<tr>
<td>(K_{BO})</td>
<td>(2.51 \times 10^{-10}) M (^{-1})</td>
<td>3</td>
</tr>
<tr>
<td>(K_{(BOOH)OH})</td>
<td>(0.01) M (^{-1})</td>
<td>14</td>
</tr>
<tr>
<td>(K_{(BOOH)O})</td>
<td>n.d. (^{2})</td>
<td>3</td>
</tr>
<tr>
<td>(K_{(BOOH)2})</td>
<td>(2.0 \times 10^{-7})</td>
<td>14</td>
</tr>
<tr>
<td>(K_{BOOH})</td>
<td>(2.0) M (^{-1})</td>
<td>14</td>
</tr>
<tr>
<td>(K_{BOO})</td>
<td>n.d. (small)</td>
<td>3</td>
</tr>
<tr>
<td>(K_{BOO})</td>
<td>(4.3) M (^{-1})</td>
<td>2</td>
</tr>
<tr>
<td>(K_{BOOB})</td>
<td>(61) M (^{-1})</td>
<td>This work</td>
</tr>
<tr>
<td>(K_{(BOO)2B})</td>
<td>(1.44 \times 10^{-7}) M (^{-1})</td>
<td>8</td>
</tr>
<tr>
<td>(K_{PB1})</td>
<td>(1.52 \times 10^{-8}) M (^{-1})</td>
<td>8</td>
</tr>
<tr>
<td>(K_{PB2})</td>
<td>(1.52 \times 10^{-8}) M (^{-1})</td>
<td>8</td>
</tr>
</tbody>
</table>
peroxymonocarbonate, $\text{HOOCO}_2^-$ ($\rho = -1.4$ in 1.76 : 1 ethanol : water$^{25}$), peroxynbenzoic acids ($\rho = -1.15$, $-1.09$ and $-1.13$ for 3-chloro, 4-nitro and 4-methyl perbenzoic acids respectively$^{16}$), and peroxomonosulfate, $\text{HOOSO}_4^-$ ($\rho = -1^{-17}$).

The experimental and theoretical work described in the present paper extends the range of substrates for the hydrogen peroxide–borate system to include substituted dimethyl anilines, methyl-$p$-tolyl sulfide, iodide, bromide, thiourea, the anion of hydrogen sulfide, $\text{HS}^-$, and hydrazine.

The substituted dimethyl anilines are included in order to determine whether the same low selectivity observed for reaction with sulfides is also seen for reactions at a tertiary nitrogen. A similar observation would lend further support to the reactive species being the dioxaborirane since it would rule out the potentially complicating possibility that the observed effects were in part due to the ability of sulfides to act both as nucleophiles and electrophiles with peroxoborates.

Substituted phenyl methyl sulfoxides are included as a comparison because they are considerably less nucleophilic toward hydroperoxides.$^{18,19}$ Thiourea, $\text{HS}^-$, iodide and bromide, all anions, are compared with thiocyanate, which was originally used to demonstrate the electrophilic reactivity of peroxoborates.$^{20}$ Hydrazine is included because borate has previously been reported to be a better catalyst than Cu(II) for the reaction between hydrazine and hydrogen peroxide, and gives the highest hydrogenation efficiency in the diimide hydrogenation of nitrile-butadiene rubber latex using hydrazine and hydrogen peroxide (although these properties were not attributed to peroxoborate formation).$^{21,22}$

Hydrogen peroxide is an environmentally benign and atom efficient oxidant, and the catalysis of its reactions by borate/boric acid is fundamental chemistry that, as well as offering environmental applications for the destruction of pollutants including dimethyl sulfide, has wider potential importance since sodium perborate is used as a source of peroxide in synthetic chemistry.$^{23-25}$ Indeed, Chaudhuri et al. have recently reported on the borate catalysed oxidation of organic sulfides by hydrogen peroxide in different media to achieve selective oxidation to sulfones and sulfones in high yields, though selectivity in this context arose from the significant difference in oxidation rates of sulfides and sulfones towards peroxides and the time allowed for the reaction to take place.$^{26}$

Results

Kinetic study

Whilst dioxaborirane is a plausible reactive species for the hydrogen peroxide–borate system in alkaline conditions, it does present a problem in terms of analysis of the kinetic data since the equilibrium constant, $K_{\text{HOBO}}$, which is likely to be small, is not known. Nevertheless, making the assumption that all of the species depicted in Fig. 1 are in rapid equilibrium with each other, it is valid to analyse the kinetic data relative to the overall concentration of the significant species in solution, i.e. the monoperoxoborate, and mono-

peroxodiborate for these experiments. This also has the advantage of retaining comparability with our earlier work. Eqn (1) was, therefore, used to analyse the kinetic data for reactions carried out at high ratios of borate to hydrogen peroxide, as with our recent study of the reaction with dimethyl sulfide. Here, $k_{\text{obs}}$ is the observed first order rate constant, $k_0$ is the rate constant for the uncatalysed reaction with hydrogen peroxide, $k_{\text{PHIBOH}}$ is the rate constant for the reaction of monoperoxoborate, including any reactive species in equilibrium with it, and $k_{\text{PHIBOH2}}$ is the rate constant for the reaction of monoperoxodiborate. The dependence of $k_{\text{obs}}$ on the total concentration of boric acid is fitted using the approach described previously,$^{1,2}$ according to eqn (1), with the rate constant for the reaction of monoperoxodiborate, $k_{\text{PHIBOH2}}$ set equal to zero. The concentrations of the peroxide species are calculated using the mass balance equations for total boron (2) and peroxide (3) concentrations and the associated equilibrium constants listed in Table 1. The presence of the polyborate species, $11$ and $12$ were accounted for in the curve fitting analysis using a bisec-

![Fig. 2](Image)

Fig. 2 Effect of total boric acid concentration on the reaction of $8 \times 10^{-3}$ M sodium thiosulfate and $0.087$ M hydrogen peroxide (pH 9.5) and of $2.1 \times 10^{-3}$ M hydrogen peroxide and $0.0225$ M sodium sulfide (pH 10.0), ionic strength 0.2 M with sodium sulfate, curves are best fit values according to eqn (1) or (4) with the rate constants shown in Table 2 and $k_{\text{PHIBOH2}}$ set to zero.
concentration on the observed rate constants.

The primary amino group. Fig. 4 shows the e

group and the slower phase represents reactions involving the

assigned to oxidation at the more nucleophilic tertiary amino

aniline, which was biphasic. The initial, faster, phase is

measured as the Ti(IV) complex, were linear for the duration of

time, corresponding to the loss of hydrogen peroxide

strates hydrazine, bromide and iodide, plots of ln("

stants for the reaction of the substrate with hydrogen peroxide and monoperox-

formation.28,29 The absorbance change is fitted to two exponen-

maximum polysulfide formation, followed by sulfate for-

bance change reaches a maximum corresponding to

carried out with the sulfide in excess. In this case the absor-

Table 2. Rate constants ± standard deviations for the reactions with hydrogen

p

excess hydrogen peroxide showed simple monoexponential

loss of absorbance, except in the case of

p

concentration causes a decrease in kobs. The corresponding

values of kP1 and kP1BOH, obtained using eqn (1), with kP1BOH2 set to zero are included in Table 2.

For the reaction of HS− with hydrogen peroxide in the pre-

sence of borate, practicalities of analysis required this to be

carried out with the sulfide in excess. In this case the abso-

bance change reaches a maximum corresponding to

maximum polysulfide formation, followed by sulfate for-

The absorbance change is fitted to two exponen
tial terms, and the e

calculated from the data of Wilson.20 a Adjusted for the statistical factor of two for the two nucleophilic nitrogens.

For the reaction of HS− with hydrogen peroxide and the sub-

strate, S, (in this case sulfide) is in excess.

Using eqn (4) as before, which is the equivalent form of

eqn (1) when the substrate, S, is in excess.

kobs = (kP1 + kP1BOH[(HO)3BOOH−]/[H2O2] + kP1BOH2[(HO)3BOOB(OH)3−]/[H2O2][S])

(4)

For the reaction between hydrogen peroxide and the sub-

strates hydrazine, bromide and iodide, plots of ln(A−A∞) versus

time, corresponding to the loss of hydrogen peroxide measured as the Ti(IV) complex, were linear for the duration of

measurements, up to three half-lives. Fig. 3 shows the effect of

total boric acid concentration on kobs. The corresponding rate

constants, obtained using eqn (4) as before, are included in

Table 2.

The reactions of the substituted dimethyl anilines with

excess hydrogen peroxide showed simple monoexponential

loss of absorbance, except in the case of p-amino dimethyl aniline, which was biphasic. The initial, faster, phase is

assigned to oxidation at the more nucleophilic tertiary amino

group and the slower phase represents reactions involving the

primary amino group. Fig. 4 shows the effect of total boric acid concentration on the observed rate constants.

For the substituted dimethyl aniline reactions, the devia-
tion of the plots from linearity cannot be explained by eqn (1)

using the formation constant for monoperoxodiborate, KBOOB,

obtained from reaction with dimethyl sulfide. The latter reaction

was carried out at low peroxide concentration whereas

dimethyl aniline reactions are carried out at high concen-

trations of peroxide and the negative deviations occur at

high boron concentrations, conditions known to favour

the formation of the diperoxodiborate anion 10,30 which

can be can be represented as eqn (5). The dependence of kobs

on the total concentration of boric acid is, therefore, fitted according to eqn (6) with both kP1BOH2 and kP2BOH2 set equal
to zero and the value of KBOOB set equal to that obtained from the dimethyl sulfide data. This yields the best fit values

Table 2 Rate constants ± standard deviations for the reactions with hydrogen peroxide and monoperoxoborate. a, b, c kP1 and kP1BOH are the second order rate con-

stants for the reaction of the substrate with hydrogen peroxide and monoperox-
oborate (including any reactive species with which it is in equilibrium) respectively.

<table>
<thead>
<tr>
<th>Substrate</th>
<th>kP1/10^(-6) M^(-1) s^(-1)</th>
<th>kP1BOH/10^(-6) M^(-1) s^(-1)</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH3SCl3</td>
<td>34 600 ± 6400</td>
<td>338 000 ± 16 000</td>
</tr>
<tr>
<td>SO4^2−</td>
<td>24 200 ± 200</td>
<td>8120 ± 240</td>
</tr>
<tr>
<td>HS^−</td>
<td>299 000 ± 15 000</td>
<td>107 000 ± 9800</td>
</tr>
<tr>
<td>Br^−</td>
<td>19 700 ± 1600</td>
<td>94 600 ± 15 000</td>
</tr>
<tr>
<td>NH3NH2</td>
<td>0.30 ± 0.05</td>
<td>25.8 ± 0.3</td>
</tr>
<tr>
<td>p-NH2C6H5N(CH3)2</td>
<td>1.6 ± 0.7d</td>
<td>23.5 ± 0.6d</td>
</tr>
<tr>
<td>p-CH3C6H5N(CH3)2</td>
<td>1800 ± 60</td>
<td>7470 ± 860</td>
</tr>
<tr>
<td>p-BrC6H5N(CH3)2</td>
<td>136 ± 5</td>
<td>2640 ± 80</td>
</tr>
<tr>
<td>C6H5N(CH3)2</td>
<td>33 ± 1</td>
<td>1520 ± 25</td>
</tr>
<tr>
<td>p-BrC6H5S(O)CH3</td>
<td>9.1 ± 0.4</td>
<td>1040 ± 25</td>
</tr>
<tr>
<td>p-CH3C6H4S(O)CH3</td>
<td>3.5 (0.76)</td>
<td>124</td>
</tr>
<tr>
<td>SCN−</td>
<td>540</td>
<td>5000</td>
</tr>
</tbody>
</table>

a Measured at a pH in the range 9.5-10. b Measured at pH 4.3. c Calculated from the data of Wilson.20 d Adjusted for the statistical factor of two for the two nucleophilic nitrogens.
of $k_{p1}$ and $k_{P1BOH}$, shown in Table 2, together with $K_\text{B(OO)2B} = 61 \pm 16 \text{ M}^{-1}$.

$$\text{(HO)}_3\text{BOO}(\text{OH})_3^{2-} + \text{H}_2\text{O}_2
\to (\text{HO})_3\text{B}(\text{OO})_2\text{B}(\text{OH})_3^{2-} + 2\text{H}_2\text{O} \; K_{\text{B(OO)2B}}$$

(5)

Experimental runs for methyl-p-tolyl sulfoxide at total boric acid and hydrogen peroxide concentrations of 0.11 M and 0.42 M, respectively, yield the rate constants shown in Table 2 at pH 9.58. Sulfoxides are susceptible to nucleophilic attack by acid and hydrogen peroxide. A better estimate of $k_{p1}$ is obtained from the run at pH 4.3 shown in Table 2.

Theoretical study

In the absence of an experimental value for $K_{\text{B(OO)}}$, we began our DFT studies with high level calculations on isomers 5 and 8 (Fig. 1). Using the B3LYP/6-311++G(3df,3pd) method with PCM implicit solvent correction for geometry optimizations and zero point energies, followed by a single point calculation at the same level of theory with both SMD implicit solvent correction, the dioxaborirane 8 was found to be 0.2 kcal mol$^{-1}$ less stable than the peroxoborate anion 5. Including the calculated entropy term (at 298 K) gave an estimate for $\Delta G$ of 0.3 kcal mol$^{-1}$ in favour of 8, suggesting that $K_{\text{B(OO)}} \sim 1$. Calculations on the equilibrium between 9 and 10, using the same level of theory, indicated that 10 is energetically favoured over 9 by 6.9 kcal mol$^{-1}$, whilst including the entropy term gave $\Delta G$ of 14.0 kcal mol$^{-1}$ in favour of 10.

For the catalytic reactions, in our previous work, we used the B3LYP/6-31+G(d,p) level of theory, with gas phase optimizations on the equilibrating anions, such as iodide with dioxaborirane. In such cases, gas phase geometry optimization will not be adequate, since the solvent plays an important role in overcoming electrostatic repulsion. Therefore, we have retained the same level of theory, but geometry optimizations were carried out in implicit solvent using the PCM method, followed by a final single point calculation using the SMD implicit solvent method. Also, in view of the more complex nature of the reactants in the present study, we have not included any explicit water molecules. This model gave barriers for the reactions of Me$_2$S with H$_2$O$_2$ and dioxaborirane of 16.6 and 8.9 kcal mol$^{-1}$ respectively. These values are ~6.3 kcal mol$^{-1}$ higher than in our previous study, nevertheless the data set presented in this work is internally self-consistent.

We have carried out calculations on the reactions of H$_2$O$_2$ and dioxaborirane with all of the species in Table 2, except for the dianion S$_2$O$_3^{2-}$. The results are summarised in Table 3, and the calculated structures for Br$^-$, NH$_2$NH$_2$ and SCN$^-$ are shown in Fig. 5; structures for the other species are given in the ESI.† The calculations on SCN$^-$ were complicated by the possibility that this anion could react at either S or N; we have checked both alternatives, and find that the transition states for attack at S are 12.2 and 10.5 kcal mol$^{-1}$ lower in energy than the equivalent states for attack at N, for the uncatalysed and catalysed reactions respectively. The final product for attack at S, OSCN$^-$, is calculated to be 13.1 kcal mol$^{-1}$ more stable than the isomeric ONCS$^-$; hence, the data in Table 3 are for the reaction of SCN$^-$ at S.

The initial reaction products from all the other calculations are the expected oxo adducts, with the exception of hydrazine, for which a spontaneous proton rearrangement occurred during geometry optimization of the dioxaborirane system to give aminohydroxylamine, H$_2$NNHOH [see Fig. 5(b)]. Although the equivalent H$_2$O$_2$ system did not show such a spontaneous

<table>
<thead>
<tr>
<th>Substrate</th>
<th>H$_2$O$<em>2$ $\Delta E</em>{\text{TS}}$/kcal mol$^{-1}$</th>
<th>O–O/Å</th>
<th>H$_2$O$_2$ $\Delta E_p$/kcal mol$^{-1}$</th>
<th>BH$_2$O$<em>4^-$ $\Delta E</em>{\text{TS}}$/kcal mol$^{-1}$</th>
<th>O–O/Å</th>
<th>BH$_2$O$_4^-$ $\Delta E_p$/kcal mol$^{-1}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>CH$_3$CH$_3$</td>
<td>+16.6</td>
<td>1.928</td>
<td>−6.3</td>
<td>+8.9</td>
<td>1.875</td>
<td>−3.5</td>
</tr>
<tr>
<td>HS$^-$</td>
<td>+13.8</td>
<td>1.815</td>
<td>−38.8</td>
<td>+9.0</td>
<td>1.844</td>
<td>−38.2</td>
</tr>
<tr>
<td>I$^-$</td>
<td>+18.9</td>
<td>2.028</td>
<td>−8.7</td>
<td>+11.6</td>
<td>2.036</td>
<td>−5.1</td>
</tr>
<tr>
<td>Br$^-$</td>
<td>+22.7$^a$</td>
<td>2.083</td>
<td>−3.0$^b$</td>
<td>+12.9$^d$</td>
<td>2.050</td>
<td>−0.1$^a$</td>
</tr>
<tr>
<td>NH$_2$NH$_2$</td>
<td>+22.3</td>
<td>1.989</td>
<td>−33.7$^b$</td>
<td>+14.4</td>
<td>1.909</td>
<td>−42.1$^c$</td>
</tr>
<tr>
<td>pNH$_2$C$_6$H$_4$N(CH$_3$)$_2$</td>
<td>+19.0</td>
<td>1.989</td>
<td>−26.2</td>
<td>+12.5</td>
<td>1.938</td>
<td>−24.0</td>
</tr>
<tr>
<td>p-CH$_2$C$_6$H$_4$N(CH$_3$)$_2$</td>
<td>+20.4</td>
<td>2.003</td>
<td>−25.0</td>
<td>+13.5</td>
<td>1.978</td>
<td>−22.2</td>
</tr>
<tr>
<td>C$_6$H$_5$N(CH$_3$)$_2$</td>
<td>+20.8</td>
<td>2.012</td>
<td>−24.3</td>
<td>+14.2</td>
<td>1.983</td>
<td>−21.5</td>
</tr>
<tr>
<td>p-BrC$_6$H$_4$N(CH$_3$)$_2$</td>
<td>+21.7</td>
<td>2.020</td>
<td>−23.5</td>
<td>+14.5</td>
<td>1.995</td>
<td>−20.0</td>
</tr>
<tr>
<td>p-CH$_2$C$_6$H$_4$SO(CH$_3$)</td>
<td>+22.9</td>
<td>1.957</td>
<td>−59.8</td>
<td>+13.2</td>
<td>1.885</td>
<td>−54.9</td>
</tr>
<tr>
<td>SCN$^-$</td>
<td>+21.9</td>
<td>1.947</td>
<td>−25.5</td>
<td>+13.8</td>
<td>1.918</td>
<td>−23.7</td>
</tr>
</tbody>
</table>

$^a$Energies calculated using an intrinsic Coulomb radius of 2.095 Å for Br. $^b$Product assumed to be H$_2$NNHOH rather than H$_2$NNH$_2$O. $^c$Product rearranges to H$_2$NNHOH during geometry optimization.
rearrangement, this product is again lower in energy. For all the reactions, the calculated reactant structures involve a hydrogen bond between the substrate heteroatom and an H atom of H₂O₂ or dioxaborirane, whilst the products generally show hydrogen bonds between the added oxo atom and an H atom of water or BO(OH)₂; in the case of hydrazine, hydrogen bonding in the product H₂NNHOH involves the N atoms.

The dioxaborirane systems all show lower transition state barriers than the uncatalysed systems; the reduction in barrier height varies from 4.9 kcal mol⁻¹ for HS⁻ to 9.8 kcal mol⁻¹ for p-CH₂C₆H₄S(O)CH₃. The four dimethyl anilines show good linear correlations between the values of ln(k) obtained from Table 2 and the calculated transition state energies in Table 3, with correlation coefficients R² of 0.986 and 0.997 for H₂O₂ and dioxaborirane respectively. Indeed, there is a reasonable qualitative correlation between these parameters for all the species in Table 3, as shown in Fig. 6. The only obvious exception to this trend came from our initial calculations on Br⁻. In this case, the calculated barriers for the uncatalysed reactions were +16.8 and +9.3 kcal mol⁻¹ respectively; these values are much lower than would be expected from the experimentally observed reaction rates. In order to probe the reason for this anomaly, we examined the reactions of H₂O₂ with halides in more detail. The experimental order of reactivity is known to be I⁻ > Br⁻ > Cl⁻,²¹ but using the method described above to calculate the transition state energies gave a predicted order of Br⁻ > I⁻ > Cl⁻. Using MP4 or alternative basis sets failed to remedy the situation, but a PCM solvent correction with the Gaussian03 defaults rather than the SMD solvent correction gave the correct order. Therefore, we suspect that the current implementation of the SMD method is the source of our difficulties. SMD was parameterized using a range of organobromine compounds,²² and uses an intrinsic Coulomb radius of 3.06 Å for Br. Comparison with the PCM radius of 2.095 Å shows that this value has changed markedly in SMD. We have therefore used the PCM radius with the SMD solvent model for Br in calculating the transition state energies for bromide, recorded in Table 3; this modification gave a much better agreement with experiment (Fig. 6).

Discussion

Previous work on the oxidation of substituted phenyl methyl sulfides was carried out at low total boron concentrations, with an excess of hydrogen peroxide, so that the predominant species were the monoperoxoborate and diperoxoborate anions.¹ In addition kinetic runs carried out in our recent study on dimethyl sulfide used low concentrations of peroxide and high total boron concentrations, where, in addition to monoperoxoborate, there was a significant concentration of monoperoxodiborate.³ The formation constant of this species, obtained from the kinetic data, was consistent with it being unreactive, in good agreement with data previously obtained from a study of the effect of borate/boric acid on the photochemical decomposition of hydrogen peroxide.² In contrast, the oxidation of the dimethyl anilines was carried out at high concentrations of peroxide and extended to higher total boron concentrations, where formation of diperoxodiborate diamions has been observed using ¹¹B NMR.³⁰ The formation constant of this species, 61 ± 16 M⁻¹ obtained from the kinetic data, which is consistent with it being unreactive, is the first that has been published.

Fig. 7 shows that for each family of substrates the ratio of reactivity with peroxoborate (and reactive species in equilibrium with it) to that with hydrogen peroxide increases as the reactivity toward hydrogen peroxide decreases. This reflects the lower sensitivity of the reaction of peroxoborate to the nature...
of the substrate compared to that of hydrogen peroxide, as described previously for the substituted phenyl methyl sulfides.\(^1\) This could potentially provide a synthetic strategy using these types of substrate and peroxoborate at higher pH that could be used when the products are unstable at the low pH required for activation of hydrogen peroxide as H\(_3\)O\(_2\)\(^+\).

It can also be seen from Fig. 7 that the result for dimethyl sulfide obtained in the preceding paper\(^3\) lies just above the line for the substituted phenyl methyl sulfides. Hence the same factors, that have been fully discussed for the latter,\(^1\) apply equally for dimethyl sulfide, where the additional methyl group confers extra nucleophilicity compared to the substituted phenyl groups. The reactivity of peroxoborate with dimethyl sulfide lies between that reported for peroxyacetic acids and hydrogen peroxide,\(^33\) just as in the case of the substituted phenyl methyl sulfides.\(^1\)

An important feature of our previous study of organic sulfide substrates is that peroxoborates, compared with other hydroperoxides, are very much less selective, consistent, under the Hammond Postulate, with the highly reactive dioxaborane being the reactive species.\(^3\) The present work shows that the oxidation of substituted dimethyl anilines, which results in N-oxide formation, also exhibits the same feature, explained as follows. Hammett plots based on \(\sigma_p\) values,\(^34\) shown in Fig. 8, give the \(\rho\) (selectivity) and log \(k_0\) (reactivity) values shown in Table 4, where they are compared with the values for peroxomonomosulfate obtained from the literature.\(^35\) The \(\rho\) value for peroxoborate is considerably less than the value, 2.2, expected if a linear reactivity selectivity relationship were to hold. This is strong evidence against the possibility that the unusual selectivity–reactivity relationship reported for the sulfides\(^1\) is due to the sulfur atom exhibiting some kind of electrophilic character specifically with peroxoborate, since the dimethyl anilines show the same behavior although the tertiary amine nitrogen atom is incapable of acting as an electrophile because of its filled-shell electronic configuration.

The respective rate constants for the reaction of hydrogen peroxide and HS\(^-\), S\(_2\)O\(_3\)^\(-2\), I\(^-\) and Br\(^-\) are similar to the accepted literature values.\(^29,36,37\) These nucleophiles, and also thiocyanate,\(^38,39\) and hydrazine (whose rate constant in Table 2 is about five times lower than the previously reported value)\(^40\) are generally considered to react with hydrogen peroxide by mechanisms in which the initial, rate-limiting, step involves heterolytic cleavage of the peroxide bond,\(^18,19,29,41\) essentially similar to the organic sulfides and dimethyl anilines. This is supported by the reasonable correlation between theory and experiment seen in Fig. 6. The stoichiometric ratios of the overall reactions are high and under the present experimental conditions the halides catalyse the decomposition of hydrogen peroxide. It is notable that bromide, the pseudohalide, SCN\(^-\), and iodide appear to fall on the same line. SCN\(^-\), S\(_2\)O\(_3\)^\(-2\), and HS\(^-\) may also be considered as a group, and, as seen in Fig. 7, thiosulfate is expected to be less reactive toward the dioxaborirane anion compared to hydrogen peroxide because of its doubly negative charge. A similar, but inverse, effect is seen where thiosulfate is more reactive toward H\(_3\)O\(_2\)\(^+\) than hydrogen peroxide, compared with singly charged nucleophiles.\(^42\)

![Fig. 7](image1.png) Comparison of the rate constants for peroxoborate with those of hydrogen peroxide. The data for the substituted phenyl methyl sulfides is from ref. 1 and the point for thiocyanate from ref. 20. The long dashed line represents log \(k_{\text{HBOH}} = \log k_1\) so that points above the line represent catalysis while those below the line represent inhibition. The short dashed line is the best fit for the data for Br\(^-\), SCN\(^-\) and I\(^-\).

![Fig. 8](image2.png) Hammett plots for the reactions of monoperoxoborate or hydrogen peroxide and \(\rho\)-substituted dimethyl anilines. Hammett \(\rho\) values calculated from the plots are shown in Table 4.

<table>
<thead>
<tr>
<th>Peroxide</th>
<th>Hammett (\log k_0)</th>
<th>Hammett (\rho)</th>
</tr>
</thead>
<tbody>
<tr>
<td>HOOSO(_3)^-(^a)</td>
<td>+3.69</td>
<td>-0.46</td>
</tr>
<tr>
<td>HOOB(OH)(_3)^-(^b)</td>
<td>-2.94</td>
<td>-1.00 ± 0.06</td>
</tr>
<tr>
<td>HOOH</td>
<td>-4.39</td>
<td>-2.53 ± 0.15</td>
</tr>
</tbody>
</table>

\(^a\) Calculated from data in ref. 35. \(^b\) Including reactive species in equilibrium with this species.

Table 4 Reactivity–selectivity parameters for the reaction of substituted dimethyl anilines (±1 standard deviation unit)
Conclusions

This study has examined the reactivity of a wide range of inorganic and organic nucleophiles with peroxoborates, confirming that peroxoborate, and those reactive species in equilibrium with it, is indeed an electrophile. Moreover, the lower than expected selectivity observed for the reaction of peroxoborate with a series of substituted dimethyl anilines, as was also observed for phenylmethyl sulfides, further supports the proposed involvement of dioxaborirane as the reactive species. At higher ratios of boron to peroxide there is kinetic evidence for unreactive peroxodiborate species, known in the literature from $^{11}$B NMR and Raman studies, and quantified in the current paper, and partly, in our previous paper on the photochemistry of peroxoborates. Knowledge of the lack of reactivity of the peroxodiborates will have a bearing on future synthetic strategies for borate catalysed hydrogen peroxide reactions. Finally, we note that dioxiranes such as dimethyl-dioxirane have proved to be valuable reagents for oxidation, particularly of hydrocarbons, whilst the wider potential of heteroatom dioxiranes as oxidation reagents has been noted, but not yet fully developed. Hence, dioxaboriranes are likely to have considerable potential as oxidants.

Experimental section

The general procedure has been described previously. The substituted dimethyl anilines, methyl-p-tolyl sulfoxide, dimethyl sulfoxide, sodium thiosulfate, sodium sulfide, potassium bromide, potassium iodide and hydrazine were obtained from Aldrich. Reactions with the dimethyl anilines, the sulfoxide, and thiosulfate were carried out with a large excess of hydrogen peroxide, monitoring the loss of the substrate absorbance spectrophotometrically at a suitable wavelength. When the absorbance of the reaction solution was so large that it was outside the dynamic range of the spectrophotometer, due to the presence of hydrogen peroxide, it was removed, after adjusting an aliquot of the reaction solution to neutral pH, using bovine catalase (BDH). Observed pseudo-first-order rate constants, $k_{obs}$, were obtained using nonlinear regression of the (in most cases) monoexponential loss of absorbance with time. Reactions with HS⁻, obtained by adjusting the oxygenated reaction solution containing sodium sulfide to pH 10.0, were carried out with excess sulfide, monitoring the absorbance of the yellow polysulfides, HSₙ⁻, produced. Reactions with bromide, iodide and hydrazine were carried out with an excess of the substrate, and the reaction followed by removing aliquots of solution and determining the hydrogen peroxide spectrophotometrically as the absorbance, A, of the titanium(iv) complex. Values of $k_{obs}$ were obtained from linear plots of ln(A−A₀) versus time. All reactions were carried out at 25 °C, and in the pH region 9.5–10.0, unless stated otherwise. All curve fitting was carried out using Grafit, version 7.

All DFT calculations were carried out within Gaussian09W. The oxidation reactions were modelled using the B3LYP functional and 6-31+G(d,p) basis set for all atoms except iodine, for which the LanL2DZ basis set was used. A trial calculation on the H₂O₂−Br⁻ system, using LanL2DZ on bromine, gave a barrier of +24.6 kcal mol⁻¹ which is comparable to the value in Table 3. Geometry optimizations were run using PCM implicit solvent corrections with water as solvent. Output geometries were verified as true minima or first order saddle points by the appropriate frequency calculations. SMD implicit solvent corrections were then obtained for single point jobs using the optimized geometries and water as solvent. The default Coulomb radii were used for all atoms except Br, for which the PCM radius of 2.095 Å was used.

Acknowledgements

We are grateful to Borax Europe for providing the initial funding for this research and to Dr K. Quill for his support for the project.

References