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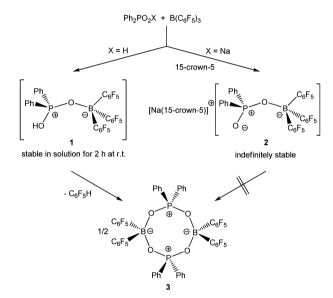
Increasing the Brønsted acidity of Ph₂PO₂H by the Lewis acid $B(C_6F_5)_3$. Formation of an eight-membered boraphosphinate ring $[Ph_2POB(C_6F_5)_2O]_2\dagger$

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Autoprotolysis of the metastable acid (C₆F₅)₃BOPPh₂OH, prepared in situ by the reaction of the rather weak Brønsted acid Ph2PO2H with the strong Lewis acid B(C₆F₅)₃, gave rise to the formation of the eight-membered ring $[Ph_2POB(C_6F_5)_2O]_2$ and C_6F_5H . The conjugate base was isolated as stable sodium crown ether salt [Na(15-crown-5)]- $[Ph_2PO_2B(C_6F_5)_3].$

Lewis acids can significantly increase the acidity of Brønsted acids. This principle is operative in the prototypical Lewis pair complex (C₆F₅)₃BOH₂, the adduct of the electron pair acceptor $B(C_6F_5)_3$ and the electron pair donor $H_2O.^2$ In MeCN, the acidity of $(C_6F_5)_3BOH_2$ (p $K_a = 8.4$) is very similar to that of HCl (p $K_a = 8.5$). Thus, (C₆F₅)₃BOH₂ is a strong acid that readily protonates basic organic⁴ and organometallic compounds.^{2,5} Diphenylphosphinic acid, Ph2PO2H, is a rather weak acid. As it is well-known that $B(C_6F_5)_3$ forms Lewis pair complexes with phosphine oxides, we were curious to study if B(C₆F₅)₃ will also increase the Brønsted

nuclear NMR spectroscopy indeed indicates the formation of a single product that was assigned to (C₆F₅)₃BOPPh₂OH (1) (Scheme 1). The ³¹P NMR spectrum (CDCl₃) of 1 shows signal at δ = 42.1 ppm that differs substantially from that of Ph₂PO₂H (33.9 ppm). The ¹¹B NMR spectrum (CDCl₃)₃ of 1 exhibits a broad signal at $\delta = -1.3$ ppm, which is significantly different



Scheme 1 Formation and reactivity of 1 and its stable sodium salt 2

from that of $B(C_6F_5)_3$ (59.0 ppm). Solutions in CDCl₃ show a limited stability and all attempts to isolate 1 by removal of the solvents failed. However, these solutions are stable at r.t. for 2 h; within this time NMR spectroscopy gave no evidence for the formation of other species. While the acid 1 could not be isolated, the reaction of Ph_2PO_2Na , $B(C_6F_5)_3$ and 15-crown-5 provided the indefinitely stable, conjugate base [Na(15-crown-5)]-[Ph₂PO₂B(C₆F₅)₃] (2), which was obtained as colourless crystals in 73% yield (Scheme 1). The 31 P and 11 B NMR spectra (THF- d_8) gave signals at δ = 22.3 and -2.7 ppm. The molecular structure of 2 reveals that the $[Na(15-crown-5)]^+$ ion and the $[Ph_2PO_2B(C_6F_5)_3]^$ ion are associated by a Na···O contact (Fig. 1). When a solution of 1 in CDCl3 was kept standing for a few hours or heated under reflux for a few minutes NMR spectroscopy indicates the formation of new species, which were identified as the eightmembered boraphosphinate ring [Ph2POB(C6F5)2O]2 (3) and

acidity of Ph2PO2H. Upon dissolving Ph₂PO₂H and B(C₆F₅)₃ in CDCl₃, multi-

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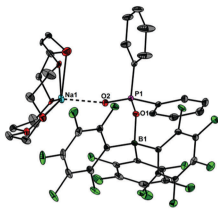


Fig. 1 Molecular structure of 2 showing 30% probability ellipsoids and the crystallographic numbering scheme. Selected bond parameters [Å, °]: B1-O1 1.508(2), P1-O1 1.544(1), P1-O2 1.482(1), Na1-O2 2.211(2), B1-O1-P1 134.5(1).

C₆F₅H. On a preparative scale, 3 was isolated in 76% yield when a solution of 1 prepared in situ from Ph_2PO_2H and $B(C_6F_5)_3$, in toluene was heated overnight under reflux (Scheme 1). This reactivity resembles the autoprotolysis of $(C_6F_5)_3BOH_2$ at elevated temperatures giving rise to the formation of [(C₆F₅)₂BOH]₃ and C₆F₅H.⁷ The eight-membered boraphosphinate ring 3 seems to be the first member of this compound class, however, we note the closely related series of cubic boraphosphonate cages in the literature comprising similar eight-membered ring subunits within the cage structure.8 The 31P and 11B NMR spectra (CDCl₃) of 3 revealed signals at δ = 37.8 and 6.3 ppm, but no coupling information. The molecular structure of 3 comprises a strongly puckered B₂P₂O₄ ring (puckering factor = 0.890), whereas isolobal eight-membered siloxane rings are usually almost planar (Fig. 2).9 The bond parameters of 3 are very similar to those of the cubic boraphosphonate cages.8 In a failed attempt to isolate 1 by crystallisation, a small crop of single crystals 4 was isolated, which turned out to be a hydrogen-bonded complex between

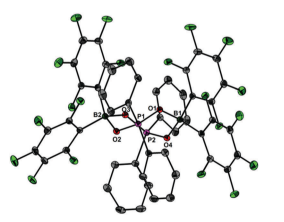


Fig. 2 Molecular structure of 3 showing 30% probability ellipsoids and the crystallographic numbering scheme. Selected bond parameters [Å, °]: B1-O1 1.501(3), B1-O4 1.507(3), B2-O2 1.506(3), B2-O3 1.513(3), P1-O1 1.538(2), P1-O2 1.537(2), P2-O3 1.534(2), P2-O4 1.539(2), B1-O1-P1 129.2(1), B1-O4-P2 129.1(1), B2-O2-P1 128.8(1), B2-O3-P2 134.2(1).

Scheme 2 Reactivity of 1 towards polymeric group 14 oxides (Me₂SiO)_n and (Me₂SnO)_n

two molecules of (C₆F₅)₃BOH₂ and the disiloxadiphosphinate [Ph₂P(O)OSiMe₂]₂O. The formation of 4 can be rationalized by the accidental cleavage of silicon grease used to seal the joints and stopcocks (Scheme 2).10 The facile cleavage of siloxanes is remarkable and points to the high Brønsted acidity of 1. Variation of the stoichiometric ratio of the reactants gave no other product than 4. The O···O donor acceptor distances (2.542(5), 2.684(4), 2.681(4), 2.559(4) Å) are indicative of medium strength hydrogen bonding. 11 The 31P, 29Si and 11B NMR spectra (THF- d_8) of 4 show signals at $\delta = 32.4, -23.9$ and 3.4 ppm. The molecular structure of 4 comprises a novel hydrogen bond motif featuring two BOH2 hydrogen bond donors and two PO hydrogen acceptors (Fig. 3). The hydrogen bond motif can be described as binary graph set $R_4^{4}(8)^{12}$ and is strongly reminiscent to that of (Ph₃SiOH)₄, ¹³ in which four silanol groups serve as donors and acceptors.

To provide a quantitative description of the Brønsted acidity increase upon going from Ph2PO2H to 1 and to reveal the corresponding electronic structure changes we carried out DFT calculations of these acids and the conjugate bases with use of the Gaussian09 package.14 The optimized molecular geometries agree well with the experimental data for 2 (Fig. 1), $[Ph_2PO_2]^-$ and $Ph_2PO_2H^{15}$ (Table S2, see ESI†). The difference in the dissociation enthalpies of Ph₂PO₂H and 1 (eqn (1) and (2)) $\Delta\Delta H = \Delta H_1 - \Delta H_2$ is estimated at the M052X/ 6-31+G** level of theory as 34.0 kcal mol⁻¹ (gas phase) and 14.1 kcal mol⁻¹ (MeCN solution). These values are indicative of much higher Brønsted acidity of 1 as compared to that of the Ph2PO2H. Our calculations of atomic charges show that the O-H bond becomes more polar upon going from Ph₂PO₂H to 1 (Table S3, see ESI†). Calculated deformation electron densities (DED) reveal a weakening of the O-H covalent bonding upon coordination of B(C₆F₅)₃ to Ph₂PO₂H (Fig. S35, see ESI†).

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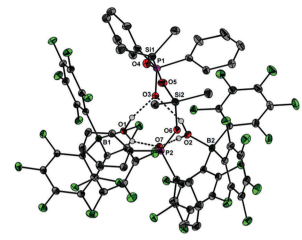


Fig. 3 Molecular structure of **4** showing 30% probability ellipsoids and the crystallographic numbering scheme. Selected bond parameters [Å, °]: B1–O1 1.562(5), B2–O2 1.556(5), P1–O3 1.492(3), P1–O4 1.558(4), P2–O6 1.567(3), P2–O7 1.502(3), Si1–O4 1.650(4), Si1–O5 1.613(4), Si2–O5 1.611(4), Si2–O6 1.668(3), P1–O4–Si1 149.7(2), P2–O6–Si2 143.8(2), Si1–O5–Si2 159.1(3), O1···O3 2.542(5), O1···O7 2.684(4), O2···O3 2.681(4), O2···O7 2.559(4).

These changes in the electronic structures explain the increased Brønsted acidity of **1**. The $\Delta H_1 - \Delta H_2$ enthalpy change is equal to the $\Delta H_3 - \Delta H_4$ difference in the B–O bond dissociation energies in the $\lceil \text{Ph}_2\text{PO}_2\text{B}(\text{C}_6\text{F}_5)_3 \rceil^-$ anion and **1** (eqn (3) and (4)).

$$Ph_2PO_2H \rightleftharpoons [Ph_2PO_2]^- + H^+ \Delta H_1$$
 (1)

$$(C_6F_5)_3BOPPh_2OH \rightleftharpoons [Ph_2PO_2B(C_6F_5)_3]^- + H^+ \Delta H_2$$
 (2)

$$[Ph_2PO_2B(C_6F_5)_3]^- \rightleftharpoons [Ph_2PO_2]^- + B(C_6F_5)_3 \Delta H_3$$
 (3)

$$(C_6F_5)_3BOPPh_2OH \rightleftharpoons Ph_2PO_2H + B(C_6F_5)_3 \Delta H_4$$
 (4)

The B-O bond in $[Ph_2PO_2B(C_6F_5)_3]^-$ is expected, therefore, to be stronger than that in 1. Indeed, the DED maps (Fig. S36, see ESI†) demonstrate a higher B-O deformation density in the anion. This stabilization of [Ph₂PO₂B(C₆F₅)₃]⁻ also contributes to the higher Brønsted acidity of 1. To compare the acidities of Ph₂PO₂H and 1 with those of other acids we calculated 16 the pK_a values in the gas phase and MeCN solution for a series of 15 compounds with tabulated experimental data in the ranges of $pK_{a(gas)} = 209-251$ and $pK_{a(MeCN)} = 0-30$ (Tables S4 and S5, see ESI†). On the basis of the linear regressions between experimental and calculated pK_a values (Fig. S37 and S38, see ESI†) the expected pK_a values for Ph₂PO₂H and 1 were found to be, respectively, 239.2 and 214.4 in the gas phase and 20.5 and 9.4 in MeCN solution. The gas-phase acidity of 1 appears to be stronger than that of CF_3SO_3H $(pK_{a(gas)}$ 219.6)¹⁷ while in MeCN solution 1 is comparable with HCl and tosylic acid (p $K_{a(MeCN)}$ 8.5³ and 8.6, ¹⁸ respectively).

In light of the remarkable siloxane bond cleavage, we have started to elaborate the reactivity of **1** towards other element oxides. Indeed, the reaction of polymeric $(Me_2SnO)_n$ with **1** rapidly occurred at r.t. and produced the eight-membered $Sn_2P_2O_4$ heterocycle $[Me_2Sn(OPPh_2O)_2SnMe_2][HOB(C_6F_5)_3]_2$ (5) in 83% yield (Scheme 2). The ¹¹⁹Sn and ³¹P MAS NMR spectra

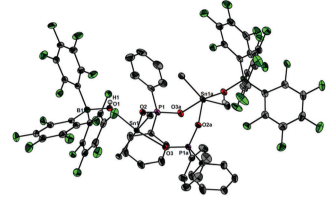


Fig. 4 Molecular structure of **5** showing 30% probability ellipsoids and the crystallographic numbering scheme. Selected bond parameters $[\mathring{A}, °]$: B1–O1 1.512(3), P1–O2 1.538(2), P1–O3a 1.515(2), Sn1–O1 2.231(2), Sn1–O2 2.040(2), Sn1–O3a 2.156(2), B1–O1–Sn1 133.5(2), P1–O2–Sn1 139.5(1), P1–O3a–Sn1a 138.1(1).

show broad signals at δ = -180.5 ppm and 31.2 ppm. Freshly prepared solutions of phase-pure 5 (checked by powder diffraction) in CDCl₃ shows four ¹¹⁹Sn NMR signals and three ³¹P NMR signals, which point to a reversible dynamic process that is not yet understood in full detail (see ESI†). Similar solution behaviour was observed for the related heterocycles [R2Sn(OPPh2O)2- $SnR_2[O_3SCF_3]_2$ (R = Ph, t-Bu), which were obtained by the reaction of $(Ph_2SnO)_n$ or $(t-Bu_2SnO)_3$ with Ph_2PO_2H and triflic acid.¹⁹ On a longer time scale (several weeks) 5 shows signs of irreversible decomposition in solution and in the solid-state. In both states, the same unassigned decomposition product with a ¹¹⁹Sn chemical shift of δ = 71.1 ppm slowly forms. The molecular structure of 5 contains a strongly puckered Sn₂P₂O₄ ring (puckering factor = 0.888)⁹ that resembles that of the slightly less puckered [t-Bu₂Sn(OPPh₂O)₂Snt-Bu₂][O₃SCF₃]₂ (puckering factor = 0.921) (Fig. 4).¹⁹ The spatial arrangement of the Sn atoms is distorted trigonal bipyramidal (geometrical goodness = 89.7°)²⁰ and defined by a C₂O₃ donor set. The Sn-O bond lengths within the ring (2.040(2) and 2.156(2) Å) are shorter than that of the exocyclic HOB(C_6F_5)₃ moiety (2.231(2) Å). The same trend was observed for [t-Bu₂Sn(OPPh₂O)₂Snt-Bu₂]-[O₃SCF₃]₂, ¹⁹ in which the endocyclic Sn-O bonds (2.045(3) and 2.173(4) Å) are shorter than the Sn-O bond length related with the triflate moiety (2.303(1) Å). It might be speculated that the longer Sn-O bonds are subject to electrolytic dissociation, which could explain the dynamic behaviour in solution. We finally studied the reactivity of 1 towards Ph₄Sn, which proceeded with facile phenyl group cleavage providing Ph₃SnOPPh₂OB(C₆F₅)₃ in 86% yield (Scheme 3). This reaction closely resembles the quantitative reaction of Ph₄Sn with triflic acid giving rise

Scheme 3 Phenyl group cleavage in Ph₄Sn using **1**.

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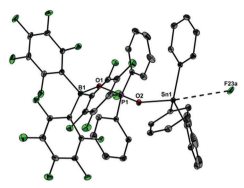


Fig. 5 Molecular structure of 6 showing 30% probability ellipsoids and the crystallographic numbering scheme. Selected bond parameters [Å. °]: B1-O1 1.527(2), P1-O1 1.521(1), P1-O2 1.527(1), Sn1-O2 2.058(1), Sn1-F23a 3.392(3), B1-O1-P1 139.9(1), P1-O2-Sn1 142.29(7)

to the formation $Ph_3SnO_3SCF_3.^{21}$ The $^{119}Sn\ NMR$ spectrum (CDCl₃) of 6 shows a doublet centred at $\delta = -59.6$ ppm with a ²/(¹¹⁹Sn-O-³¹P) coupling of 146 Hz, which suggests that the Sn atoms are tetracoordinate in solution (Fig. 5). In the solid-state, 6 comprises a 1D coordination polymer with distorted trigonal bipyramidal Sn atoms (geometrical goodness = 51.6°)²⁰ defined by a C₃OF donor set.

The Brønsted acidity of Ph₂PO₂H was significantly increased upon addition of the Lewis acid B(C₆F₅)₃ giving rise to (C₆F₅)₃BOPPh₂OH (1) in solution. Unlike its conjugate base $[Na(15\text{-crown-5})][Ph_2PO_2B(C_6F_5)_3]$ (2), the acid 1 is thermally unstable and undergoes autoprotolysis and formation of the boraphosphinate ring [Ph₂POB(C₆F₅)₂O]₂ (3) and C₆F₅H. Despite its limited life span, 1 can be used for synthetic purposes, as was demonstrated for two examples from organotin chemistry. The stable water adduct (C₆F₅)₃BOH₂ is known to bind up to two additional water molecules via hydrogen bonding, e.g. (C₆F₅)₃BOH₂·2H₂O,²² which adversely affects the stoichiometric control of protonation reactions. Moreover, the various related anions, e.g. $[HOB(C_6F_5)_3]^-$, $[HO\{B(C_6F_5)_3\}_2]^$ and $[O(B(C_6F_5)_3)_2]^{2-,2,4}$ suggest that hydroxide and oxide ions may be also transferred upon protonation. These adverse properties have not been observed for 1. We are currently investigating if the acidity of other Brønsted acids, such sulfinic and sulfonic acids, may be also increased by applying the same concept.

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