Synthesis and reactivity of cyclo-tetra(stibinophosphonium) tetracations: redox and coordination chemistry of phosphine–antimony complexes†

Saurabh S. Chitnis, Alasdair P. M. Robertson, Neil Burford,* Jan J. Weigand* and Roland Fischer

Reductive elimination of \([R_3PPR_3]^{2+}\), \([11(R)]^{2+}\), from the highly electrophilic Sb\(^{III}\) centres in \([R_3PPR_3]^{3+}\), \([8(R)]^{3+}\), gives Sb\(^{1+}\) containing cations \([R_3PPR_3]^{2+}\), \([9(R)]^{2+}\), which assemble into frameworks identified as cyclo-tetra(stibinophosphonium) tetracations, \([R_3PPR_3]^{4+}\), \([10(R)]^{4+}\). A phosphine catalyzed mechanism is proposed for conversion of fluoroantimony complexes \([R_3PPR_3]^{2+}\), \([7(R)]^{2+}\), to \([10(R)]^{4+}\), and the characterization of key intermediates is presented. The results constitute evidence of a novel ligand activation pathway for phosphines in the coordination sphere of hard, electron deficient acceptors. Characterization of the associated reactants and products supports earlier, albeit less definitive, detection of analogous phosphine ligand activation in Cu\(^{II}\) and Tl\(^{III}\) complexes, demonstrating that these prototypical ligands can behave simultaneously as reducing agents and \(\sigma\) donors towards a variety of hard acceptors. The reactivity of the parent cyclo-tetra(stibinophosphonium) tetracation, \([10(\text{Me})]^{4+}\), is directed by high charge concentration and strong polarization of the P–Sb bonds. The former explains the observed facility for reductive elimination to yield elemental antimony and the latter enabled activation of P–Cl and P–H bonds to give phosphinophosphonium cations, \([\text{Me}_3PPR_3]^{4+}\), including the first example of an H-phosphinophosphonium, \([\text{Me}_3PP(H)R]^{2+}\), and 2-phosphino-1,3-diphosphonium cations, \([\text{Me}_2P(\text{PR})_3]^{2+}\). Exchange of a phosphine ligand in \([10(\text{Me})]^{4+}\) with \([\text{nacnac}]^{1-}\) gives \([\text{Me}_3PPR_3Sb]^{1+}\), \([\text{nacnac}]^{1-}\), and \([15(\text{Me})]^{3+}\), and with dmap gives \([\text{Me}_3PPSb]^{1+}\). The lability of P–Sb or Sb–Sb interactions in \([10(\text{Me})]^{4+}\) has also been illustrated by characterization of heteroleptically substituted derivatives featuring PMe\(_3\) and PET\(_3\) ligands.

Introduction

Phosphines are prototypical ligands in the coordination chemistry of d-block metals. While the chemistry of p-block elements is primarily defined by covalent bonding as typified by organic frameworks, an array of phosphine adducts has also been characterized for main group element acceptors.\(^1,\(^3\)\(^9\)\) Beyond their versatile ligand properties as neutral, two-electron donors (L-type),\(^6\) phosphines also exhibit redox reactivity within the coordination sphere of an acceptor. For example, reductive elimination of tetraorgano- or halotriorganophosphonium cations (Scheme 1a),\(^6\) and oxidative addition of PR–X bonds (Scheme 1b),\(^6\) or P–R bonds (Scheme 1c)\(^6\) are all known pathways of tertiary phosphine activation in transition metal chemistry. One report\(^5\) hints at the reductive elimination of a diphosphonium dication from a phosphine–metal complex (Scheme 1d). In this instance, spectroscopic studies indicate that the reaction of excess PMe\(_3\) with \([\text{Cu(MeCN)}_3]^{[PF_6]}\) or \([\text{Tl(MeCN)}_3]^{[UF_6]}\) yields \([\text{Me}_3PPMe_3]^{2+}\), and the reduced metal complexes \([\text{Cu(PMe}_3)_2]^{[PF_6]}\) and \([\text{Tl(PMe}_3)_2]^{[UF_6]}\), respectively.\(^6\) However, neither the high oxidation state reactants nor the reduced products have been structurally verified and three different \(^{31}\)P NMR chemical shifts were ascribed to \([\text{Me}_3PPMe_3]^{2+}\) (depending upon the counterion: +65.0 ppm, +46.3 ppm, or +27.8 ppm). As reductive elimination is observed for both a transition metal (Cu\(^{II}\)) and a main group metal (Tl\(^{III}\)) acceptor, phosphine activation may be broadly applicable to complexes exhibiting a mismatch between hard (high oxidation state/charge) acceptors and soft phosphine donors. Indeed phosphines are considered poor donors for hard acceptors and coordination to such centres generally requires enforcement by chelate or pincer ligands.\(^1,\(^11\)\)

† Electronic supplementary information (ESI) available. See DOI: 10.1039/c4sc03939d
As part of a systematic evolution of p-block element phosphine complexes, we have sought derivatives featuring multiply-charged, hard acceptors and now report evidence of a new phosphine ligand activation pathway in the coordination sphere of polycationic SbIII centres. Specifically, reductive elimination of diphosphonium dications (Scheme 1e) from trialkylphosphine complexes of SbIII has been demonstrated, comprehensively defining a fundamental P–P bond forming redox process. The reduction products are the unusual cyclo-tetra(stibinophosphonium) tetracations \([10(R)]^{4+}\), representing a new catena-homocyclic framework. Examples of cationic homocycles for p-block metalloids are limited to unsupported selenium and tellurium dications and heavily substituted silicon or germanium monocations. For antimony, a number of acyclic catenated monocations ([1]^{1+} and [2]^{1+}) and dications ([3]^{2+}, [4]^{2+}, [5]^{2+} and [6]^{2+}) have recently been isolated (Chart 1), but generally on small scales, precluding further reactivity studies of these interesting species. Enabled by a rational and large scale synthetic protocol for cations \([10(R)]^{4+}\), we now report the reaction chemistry of the prototypical derivative, \([10(Me)]^{4+}\), debuting the coordination chemistry of a new catena-element framework.

### Results and discussion

**Reactions of PR₃ with FSb(OTf)₂ and Sb(OTf)₃**

Combinations of FSb(OTf)₂ or Sb(OTf)₃ with PR₃ (R = Me, Et, Pr, or Bu) in MeCN solvent at the optimized stoichiometries given in Scheme 2 have been investigated. The \(^{31}\)P, \(^{13}\)C, \(^{19}\)F and \(^{1}\)H NMR spectra of reaction mixtures indicate quantitative formation of \([10(R)]^{4+}\), \([11(R)]^{2+}\), and \([12(R)]^{1+}\) as triflate salts in reaction mixtures containing trialkylphosphines and (a) Sb(OTf)₃ or (b) FSb(OTf)₂.

![Scheme 1](https://example.com/scheme1.png)

**Chart 1** Structurally confirmed cations featuring Sb–Sb bonds. See text for references.

As part of a systematic evolution of p-block element phosphine complexes, we have sought derivatives featuring multiply-charged, hard acceptors and now report evidence of a new phosphine ligand activation pathway in the coordination sphere of polycationic SbIII centres. Specifically, reductive elimination of diphosphonium dications (Scheme 1e) from trialkylphosphine complexes of SbIII has been demonstrated, comprehensively defining a fundamental P–P bond forming redox process. The reduction products are the unusual cyclo-tetra(stibinophosphonium) tetracations \([10(R)]^{4+}\), representing a new catena-homocyclic framework. Examples of cationic homocycles for p-block metalloids are limited to unsupported selenium and tellurium dications and heavily substituted silicon or germanium monocations. For antimony, a number of acyclic catenated monocations ([1]^{1+} and [2]^{1+}) and dications ([3]^{2+}, [4]^{2+}, [5]^{2+} and [6]^{2+}) have recently been isolated (Chart 1), but generally on small scales, precluding further reactivity studies of these interesting species. Enabled by a rational and large scale synthetic protocol for cations \([10(R)]^{4+}\), we now report the reaction chemistry of the prototypical derivative, \([10(Me)]^{4+}\), debuting the coordination chemistry of a new catena-element framework.

**Results and discussion**

**Reactions of PR₃ with FSb(OTf)₂ and Sb(OTf)₃**

Combinations of FSb(OTf)₂ or Sb(OTf)₃ with PR₃ (R = Me, Et, Pr, or Bu) in MeCN solvent at the optimized stoichiometries given in Scheme 2 have been investigated. The \(^{31}\)P, \(^{13}\)C, \(^{19}\)F and \(^{1}\)H NMR spectra of reaction mixtures indicate quantitative formation of \([10(R)]^{4+}\), \([11(R)]^{2+}\), and \([12(R)]^{1+}\) as triflate salts in reaction mixtures containing trialkylphosphines and (a) Sb(OTf)₃ or (b) FSb(OTf)₂.

![Scheme 2](https://example.com/scheme2.png)

**Chart 1** Structurally confirmed cations featuring Sb–Sb bonds. See text for references.

As part of a systematic evolution of p-block element phosphine complexes, we have sought derivatives featuring multiply-charged, hard acceptors and now report evidence of a new phosphine ligand activation pathway in the coordination sphere of polycationic SbIII centres. Specifically, reductive elimination of diphosphonium dications (Scheme 1e) from trialkylphosphine complexes of SbIII has been demonstrated, comprehensively defining a fundamental P–P bond forming redox process. The reduction products are the unusual cyclo-tetra(stibinophosphonium) tetracations \([10(R)]^{4+}\), representing a new catena-homocyclic framework. Examples of cationic homocycles for p-block metalloids are limited to unsupported selenium and tellurium dications and heavily substituted silicon or germanium monocations. For antimony, a number of acyclic catenated monocations ([1]^{1+} and [2]^{1+}) and dications ([3]^{2+}, [4]^{2+}, [5]^{2+} and [6]^{2+}) have recently been isolated (Chart 1), but generally on small scales, precluding further reactivity studies of these interesting species. Enabled by a rational and large scale synthetic protocol for cations \([10(R)]^{4+}\), we now report the reaction chemistry of the prototypical derivative, \([10(Me)]^{4+}\), debuting the coordination chemistry of a new catena-element framework.

**Results and discussion**

**Reactions of PR₃ with FSb(OTf)₂ and Sb(OTf)₃**

Combinations of FSb(OTf)₂ or Sb(OTf)₃ with PR₃ (R = Me, Et, Pr, or Bu) in MeCN solvent at the optimized stoichiometries given in Scheme 2 have been investigated. The \(^{31}\)P, \(^{13}\)C, \(^{19}\)F and \(^{1}\)H NMR spectra of reaction mixtures indicate quantitative formation of \([10(R)]^{4+}\), \([11(R)]^{2+}\), and \([12(R)]^{1+}\) as triflate salts in reaction mixtures containing trialkylphosphines and (a) Sb(OTf)₃ or (b) FSb(OTf)₂.

![Scheme 2](https://example.com/scheme2.png)

**Chart 1** Structurally confirmed cations featuring Sb–Sb bonds. See text for references.

As part of a systematic evolution of p-block element phosphine complexes, we have sought derivatives featuring multiply-charged, hard acceptors and now report evidence of a new phosphine ligand activation pathway in the coordination sphere of polycationic SbIII centres. Specifically, reductive elimination of diphosphonium dications (Scheme 1e) from trialkylphosphine complexes of SbIII has been demonstrated, comprehensively defining a fundamental P–P bond forming redox process. The reduction products are the unusual cyclo-tetra(stibinophosphonium) tetracations \([10(R)]^{4+}\), representing a new catena-homocyclic framework. Examples of cationic homocycles for p-block metalloids are limited to unsupported selenium and tellurium dications and heavily substituted silicon or germanium monocations. For antimony, a number of acyclic catenated monocations ([1]^{1+} and [2]^{1+}) and dications ([3]^{2+}, [4]^{2+}, [5]^{2+} and [6]^{2+}) have recently been isolated (Chart 1), but generally on small scales, precluding further reactivity studies of these interesting species. Enabled by a rational and large scale synthetic protocol for cations \([10(R)]^{4+}\), we now report the reaction chemistry of the prototypical derivative, \([10(Me)]^{4+}\), debuting the coordination chemistry of a new catena-element framework.

**Results and discussion**

**Reactions of PR₃ with FSb(OTf)₂ and Sb(OTf)₃**

Combinations of FSb(OTf)₂ or Sb(OTf)₃ with PR₃ (R = Me, Et, Pr, or Bu) in MeCN solvent at the optimized stoichiometries given in Scheme 2 have been investigated. The \(^{31}\)P, \(^{13}\)C, \(^{19}\)F and \(^{1}\)H NMR spectra of reaction mixtures indicate quantitative formation of \([10(R)]^{4+}\), \([11(R)]^{2+}\), and \([12(R)]^{1+}\) as triflate salts in reaction mixtures containing trialkylphosphines and (a) Sb(OTf)₃ or (b) FSb(OTf)₂.

![Scheme 2](https://example.com/scheme2.png)

**Chart 1** Structurally confirmed cations featuring Sb–Sb bonds. See text for references.
observed in rare examples of catena-aniony cations [cf. \[1\] \[11\] = 2.8205(12) Å, \[12\] = 2.8278(3) Å, \[13\] = 2.8203(4) Å, \[14\] = 2.7624(11) Å and 2.7867(12) Å, \[15\] = 2.811(1) Å and 2.830(1) Å, \[16\] = 2.8484(12) Å and 2.8533(12) Å]. Consistent with the high Lewis acidity of the Sb center, the P-Sb distances \[2.552(2)-2.578(2) Å\] are similar to those observed in other trihalide salts such as \([\text{Me}_3\text{P}]_2\text{SbCl}_2\) and \([\text{Me}_3\text{P}]_2\text{SbBr}_2\). Given the high molecular charge, these values are expectedly smaller than the sum of the van der Waals radii (\(2.554(4)\) Å) and \([\text{Me}_3\text{P}]_2\text{SbF}_3\) (\(2.538(4)\) Å).

A number of Sb-O\(_{\text{OTf}}\) contacts are also observed, the shortest of which measure 3.210(4) Å for \([\text{10(Me)}]^{+}\) and 2.871(8) Å for \([\text{10(Et)}]^{+}\). Given the high molecular charge, these values are expectedly smaller than the sum of the van der Waals radii (\(3.61(1)\) Å) but nevertheless significantly longer than the sum of the single bond covalent radii (\(2.05(1)\) Å) for the two elements. For \([\text{10(Me)}]^{+}\), a gas-phase optimization of the cation at the MP2 level in the absence of the trihalide anions produced a geometry that is essentially identical to that observed experimentally, and we therefore infer that the anion contacts do not distort the structural features to a measurable extent.

The reactions in Scheme 2a represent a two electron reduction of each antimony center, engaging an electron reductive coupling of four antimony centers to form derivatives of \([\text{10(R)}]^{+}\). In Scheme 2a, eight of the twelve equivalents of phosphine are involved in the redox process, being oxidatively coupled to give four diphosphonium cations, \([\text{11(R)}]^{2+}\), and the remaining four equivalents represent ligands on the reduced antimony centers of \([\text{10(R)}]^{+}\).

Scheme 2b describes a similar redox process that involves formation of \([\text{11(R)}]^{3+}\) as transients, which are converted to the corresponding fluorphosphonium cations, \([\text{12(R)}]^{+}\), in the presence of the fluoride ion, as envisaged in the mechanism outlined in Scheme 3 (left). The key feature in both processes is reductive elimination of a diphosphonium unit from a hard, tricationic Sb\(^{10}\) center to give a soft, monocationic Sb\(^{11}\) center, representing a novel mode of phosphine ligand activation in the coordination sphere of metals (Scheme 1c).

\(^{31}\)P NMR spectra (Fig. 2) of reaction mixtures containing \(\text{PR}_3\) and \(\text{FSb(OTf)}_2\) in a 2:1 stoichiometry show a broad doublet in the +20 to +40 ppm range and the signal due to the free phosphine (−60 to −20 ppm) is not observed. The \(^{19}\)F NMR spectra of these mixtures show a broad triplet in the region −170 to −175 ppm and no evidence of \(\text{FSb(OTf)}_2\). The broadness of peaks in the \(^{31}\)P and \(^{19}\)F NMR spectra is consistent with the connectivity of these nuclides to a quadrupolar antimony center \([I = 5/2 \text{ for } ^{121}\text{Sb} (57%), 7/2 \text{ for } ^{123}\text{Sb} (43%)]\), and we assign these signals to the dicationic bis-phosphonium cations \([\text{7(R)}]^{2+}\), which are stable as MeCN solutions (Scheme 3a). Upon addition of ca. 5 mol % of phosphine to these solutions, the \(^{31}\)P NMR signals due to cations \([\text{7(R)}]^{2+}\) are replaced over 16 hours by doublets corresponding to \([\text{12(R)}]^{+}\) (\(d^{11}: +140 \text{ to } +150 \text{ ppm}, d^{19}: 950 \text{ to } 1000 \text{ Hz})\) and a singlet in the −25 to 0 ppm range, corresponding to \([\text{10(R)}]^{+}\). Addition of ca. 15 mol % of phosphine increases the rate of the reaction and effects complete conversion of \([\text{7(R)}]^{2+}\) to \([\text{12(R)}]^{+}\) and \([\text{10(R)}]^{+}\) within an hour.

We propose that the displacement of fluoride from \([\text{7(R)}]^{2+}\) by added phosphine yields the highly-charged trications \([\text{8(R)}]^{3+}\) (Scheme 3b), which undergo reductive elimination of \([\text{11(R)}]^{2+}\) and \([\text{9(R)}]^{+}\) (Scheme 3c). Subsequent tetramerization of the six-valence electron cations \([\text{9(R)}]^{+}\) to \([\text{10(R)}]^{4+}\) (Scheme 3d), and displacement of \(\text{PR}_3\) from \([\text{11(R)}]^{2+}\) by fluoride gives \([\text{12(R)}]^{+}\), regenerating the phosphine catalyst (Scheme 3e). Cyclization of transients \([\text{9(R)}]^{+}\) is analogous to the formation of tetrameric \((\text{Mes-E})_2(\text{MeCN})_2\) \(d = 2,4,6\)-trimethylphenyl, \(E = \text{As or Sb}) \) via catalytic oxidation of \(\text{MesAs}^{2+}\) from a zirconium complex or \(\text{MesSb}^{1+}\) from a hafnium complex. Nucleophilic displacement of \(\text{PMMe}_3\) has been reported in reaction mixtures of \([\text{11(Me)}][\text{ClO}_4]_2\) and \([\text{NEt}_4][\text{F}]\), and we have further confirmed that the equimolar reaction of \([\text{11(Me)}][\text{OTf}]_2\) with CsF (Fig. S1, ESIF) yields a 1:1 mixture of \(\text{PMMe}_3\) and \([\text{12(Me)}]^{+}\). Trications \([\text{8(R)}]^{3+}\) are also implicated in the formation of \([\text{10(R)}]^{4+}\) from \(\text{Sb(OTf)}_3\) (Scheme 3f) and we have previously reported the structure of the ternary salt \([\text{8(Me)}][\text{11(Me)}][\text{OTf}]_2\) from a 3:1 mixture of \(\text{Sb(OTf)}_3\) and \(\text{PMMe}_3\) at −30 °C (vide infra). Consistent with the role of \([\text{8(Me)}]^{+}\) as an intermediate, the same reaction stoichiometry yields only \([\text{10(Me)}][\text{OTf}]_2\) and \([\text{11(Me)}][\text{OTf}]_2\) at ambient temperature.

The \(^{31}\)P NMR spectra of reaction mixtures containing \([[\text{Me}_3\text{P}]_2\text{SbCl}]^{2+}\) and 20 mol% \(\text{PMMe}_3\) show only partial conversion to \([\text{10(Me)}][\text{OTf}]_2\) and \([\text{11(Me)}][\text{OTf}]_2\) after 48 hours. Additionally, a broad signal at +10.4 ppm is also observed (Fig. S2, ESIF), which is close to the average for the values in \([\text{Me}_3\text{P}]_2\text{SbCl}]^{2+}\) (±15.8 ppm) and \([\text{Me}_3\text{P}]_2\text{SbCl}]^{4+}\) (±6.2 ppm), suggesting that the free chloride ion is sequestered in an...
The equilibrium between the starting material and \([\text{Me}_3\text{P})_2\text{SbCl}_2]^{1+}\). Consequently, nucleophilic attack by chloride to liberate free phosphine from \([11(\text{Me})]^{2+}\) is precluded in these reaction mixtures and neither \([\text{Me}_3\text{PCl}]^{1+}\) nor free phosphine are detected by \(^{31}\text{P}\) NMR spectroscopy. Signifying the role of free phosphine as a catalyst, formation of \([10(\text{Me})]^{4+}\) does not occur catalytically in the chloride system because the reaction is arrested upon formation of \([11(\text{Me})]^{2+}\), which is the spectroscopically detected oxidation product. Generation of free phosphine from diphosphonium, the turnover limiting step, does not take place (Scheme 3, right). In contrast, no diphosphonium is detected in reactions involving the \(\text{F}^\text{–}\)-diphosphonium complexes \([7(\text{R})]^{2+}\) (Scheme 3, left), where, due to nucleophilic attack by fluoride anions on \([11(\text{R})]^{2+}\), only the fluorophosphoniums \([12(\text{R})]^{1+}\) are detected as the oxidation product and the formation of \([10(\text{R})]^{4+}\) occurs catalytically in the presence of free \(\text{PR}_3\). Differences in the reactivity of homologous \(\text{Sb}\text{–}X\) (\(X = \text{Cl, F}\)) complexes towards Lewis acids have been noted previously.\(^{33}\)

Solution NMR data for derivatives of \([7(\text{R})]^{2+}\), \([8(\text{R})]^{3+}\), \([9(\text{R})]^{1+}\), \([10(\text{R})]^{4+}\), \([11(\text{R})]^{2+}\), and \([12(\text{R})]^{1+}\) are summarized in Table 2, with evidence for the assignments discussed below. It has not been possible to detect or isolate derivatives of \([9(\text{R})]^{1+}\). Attempts to trap these cations, or radical intermediates arising from one-electron processes, in the presence of a twenty-fold excess of 2,3-dimethyl-1,3-butadiene were unsuccessful.

---

**Scheme 3**  (Left) Proposed catalytic mechanism for the formation of derivatives of cations \([7(\text{R})]^{2+}\), \([8(\text{R})]^{3+}\), \([9(\text{R})]^{1+}\), \([10(\text{R})]^{4+}\), \([11(\text{R})]^{2+}\), and \([12(\text{R})]^{1+}\). See text for descriptions of a–f. (Right) Non-catalytic formation of \([10(\text{Me})]^{4+}\) from the reaction of \([\text{Me}_3\text{P})_2\text{SbCl}_2]^{2+}\) with \(\text{PMe}_3\).
Table 2 Solution NMR data (CD$_3$CN, 298 K) for derivatives of [7(R)]$^{2+}$, [8(R)]$^{3+}$, [10(R)]$^{2+}$, [11(R)]$^{2+}$, and [13]$^{2+}$. Values in parentheses indicate literature values for chemical shifts of known compounds. Values in square brackets denote peak width at half-maximum where the expected $^3$J$_{PF}$ coupling was not observed (n.o.).

<table>
<thead>
<tr>
<th>Compound</th>
<th>$^3$P (ppm)</th>
<th>$^3$F (ppm)</th>
<th>$^3$J$_{PF}$ (Hz)</th>
</tr>
</thead>
<tbody>
<tr>
<td>[7(Me)]$^{2+}$</td>
<td>+15.9</td>
<td>−178.2</td>
<td>44</td>
</tr>
<tr>
<td>[7(Me)]$^{2+}$</td>
<td>+38.0 [23]</td>
<td>−174.2 [73]</td>
<td>n.o.</td>
</tr>
<tr>
<td>[7(Pr)]$^{2+}$</td>
<td>+29.3</td>
<td>−173.4</td>
<td>41</td>
</tr>
<tr>
<td>[7(Bu)]$^{2+}$</td>
<td>+32.4 [62]</td>
<td>−173.9 [32]</td>
<td>n.o.</td>
</tr>
<tr>
<td>[8(Me)]$^{3+}$</td>
<td>+1.3</td>
<td>−187.1</td>
<td>39</td>
</tr>
<tr>
<td>[8(Me)]$^{3+}$</td>
<td>+21.3</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[8(Pr)]$^{3+}$</td>
<td>+27.8</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[8(Bu)]$^{3+}$</td>
<td>+22.1</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[8(Bu)]$^{3+}$</td>
<td>+22.4</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[10(Me)]$^{3+}$</td>
<td>−24.4</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[10(Pr)]$^{3+}$</td>
<td>+9.3</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[10(Bu)]$^{3+}$</td>
<td>+0.6</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[11(Me)]$^{3+}$</td>
<td>+28.5 (28.4)$^{27}$</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[11(Pr)]$^{3+}$</td>
<td>+31.4 (32)$^{11}$</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[11(Bu)]$^{3+}$</td>
<td>+31.0 (32)$^{11}$</td>
<td>—</td>
<td>—</td>
</tr>
<tr>
<td>[12(Me)]$^{3+}$</td>
<td>+148.0</td>
<td>−138.0</td>
<td>948</td>
</tr>
<tr>
<td>[12(Pr)]$^{3+}$</td>
<td>+150.2</td>
<td>−159.3</td>
<td>973</td>
</tr>
<tr>
<td>[12(Bu)]$^{3+}$</td>
<td>+145.6</td>
<td>−154.7</td>
<td>971</td>
</tr>
</tbody>
</table>

Scheme 4 Synthesis of [11(ET)]$^{3+}$ via oxidative coupling of PEt$_3$ with [Ph$_3$Sb][OTf]$_2$.

Fig. 3 Molecular structure of the cation in [11(ET)][OTf]$_2$ in the solid state. Hydrogen atoms and triflate anions have been omitted for clarity. Thermal ellipsoids are drawn at 30% probability level. Bond lengths (Å) and angles (°) are as follows: P1–P2 = 2.2209(8), P1–C3 = 1.8001(2), P1–C5 = 1.8021(2), P1–C7 = 1.806(2), P2–C9 = 1.796(2), P2–C11 = 1.809(2), P2–C12 = 1.798(2), C3–P1–C2–C9 = 28.7 (1).

Derivatives of [7(R)]$^{2+}$ represent the first examples of phosphine complexes of fluoroantimony acceptors although numerous fluoroantimony complexes with hard, oxidatively-resistant donors such as pyridines, ethers, and nitrile oxide have been reported. The $^3$J$_{PF}$ couplings for [7(Me)]$^{2+}$ and [7(Pr)]$^{2+}$ are resolved as a doublet in the $^3$P NMR spectra and as a triplet in the $^3$F NMR spectra, consistent with an AX$_2$ spin system. Fine structure could not be resolved for [7(Et)]$^{2+}$ and [7(Bu)]$^{2+}$ even under the dilute conditions and low temperature (−30 °C) employed to mitigate broadening due to exchange.

Although [7(Me)][OTf]$_2$ and [7(ET)][OTf]$_2$ have both been isolated as analytically pure substances and spectroscopically characterized, we were unable to obtain X-ray quality crystals. Moreover, to the best of our knowledge, there are no known examples of $^3$J$_{PF}$ coupling constants through an antimony centre for direct comparison with our assigned NMR data. For this reason, we prepared and isolated the analogous [(dmpe) SbF][OTf]$_2$, [13][OTf]$_2$, from an equimolar mixture of 1,2-bis (dimethylphosphino)ethane (dmpe) and FSB[OTf]$_2$ in MeCN. The solid state structure of [13][OTf]$_2$, as determined by X-ray crystallography, shows a dimeric arrangement with the cations bridged by O–S–O contacts from the triflate anions, and additional interactions with two non-bridging triflate anions, as shown in Fig. S3 (ESI). The pyramidal geometry at Sb in the cation is retained in solution, as demonstrated by the two non-equivalent methyl group resonances in the $^3$C (6.1 and 7.2 ppm) and $^1$H NMR (1.86 and 2.10 ppm) spectra. Crucially, the expected $^3$J$_{PF}$ coupling was unambiguously observed (Fig. S4, ESI) in signals due to [13]$^{2+}$, and the chemical shift and coupling constants are comparable to those assigned to derivatives of [7(R)]$^{2+}$ (Table 2).

It was not possible to isolate salts of [8(R)]$^{3+}$ due to their high reactivity, consistent with their disproportionation to [11(R)]$^{2+}$ and [10(R)]$^{3+}$ in solution as proposed above. The $^3$P NMR signals assigned to derivatives of [8(R)]$^{3+}$ are singlets and broadened ($\Delta$P$_{1/2}$ = 90–500 Hz), presumably due to a combination of the quadrupolar antimony nuclides$^{40}$ and dynamic ligand exchange. Nevertheless, [8(Me)][OTf]$_5$ has been detected as a co-crystallate with [11(Ph)][OTf]$_2$ in a 3 : 1 reaction mixture of PMe$_3$ and Sb(OTf)$_3$ at −30 °C.$^{26}$ The molecular structure of [8(Me)]$^{3+}$ (Fig. S5, ESI) in the salt [8(Ph)][11(Me)][OTf]$_2$ shows a pyramidal arrangement around the Sb atom with three P–S–P bond lengths in the range 2.597(4)–2.611(5) Å and P–Sb–P angles in the range 101.3(3)–102.4(2)$^{26}$°. In addition, three interion Sb–O contacts are observed in the 2.791(2)–2.960(2) Å range (cf. $\sum r_{Sb-O} = 3.61$ Å)$^{44}$ with each contact appearing trans to a P–Sb bond, illustrating a triple displacement of triflate anions from Sb(OTf)$_3$ by three PMe$_3$ ligands.

Signals attributed to derivatives of [11(R)]$^{2+}$ are assigned by comparison with previously reported $^3$P chemical shifts for their triflate or perchlorate salts in MeCN for [11(Me)]$^{3+}$, [11(Pr)]$^{3+}$, and [11(Bu)]$^{2+}$.$^{27,31}$ Isolation of [11(Pr)][OTf]$_2$ enabled comprehensive characterization, including X-ray structural determination and we have reported this data elsewhere.$^{28}$ The salt [11(ET)][OTf]$_2$ has been prepared independently from a 2 : 1 reaction of PEt$_3$ with in situ generated Ph$_3$Sb[OTf]$_2$, according to Scheme 4, and the structure of the cation is shown in Fig. 3. The P–P bond length [2.2209(8) Å] is comparable to that in rare examples of acyclic diphenophonium dications such as [11(Ph)]$^{2+}$ [2.198(2) Å]$^{27}$ or [Me$_4$PPEt$_3$]$^{2+}$ [2.216(1) Å]$^{27}$ and a...
Phosphines such as PiPr$_3$ were unsuccessful. A $^{31}$P NMR assay of contact is 177.34(8) ppm for the proposed assignments (Fig. S6, ESI†).

Notably, the chemical shift of the phosphorus atom, which is marginally shorter than 106.38(6), 105.2(1), 106.38(6). Ellipsoids are drawn at 30% probability level. Bond lengths (Å) and angles (°) are as follows: P1–F2 = 1.551 (1), P1–C = 1.757(2), 1.755 (2), 1.755 (2). P1–O3 = 3.301(2), F2–P1–O3 = 177.34(8), P2–P1–C = 106.38(6), 105.2(1), 106.38(6).

![Fig. 4 Molecular structure of [12(Me)][OTf] in the solid state. Thermal ellipsoids are drawn at 30% probability level. Bond lengths (Å) and angles (°) are as follows: P1–F2 = 1.551 (1), P1–C = 1.757(2), 1.755 (2), 1.755 (2), P1–O3 = 3.301(2), F2–P1–O3 = 177.34(8), P2–P1–C = 106.38(6), 105.2(1), 106.38(6).](image)

**Scheme 5** Thermolysis of [10(Me)][OTf]$_4$ to yield [11(Me)][OTf]$_2$ and elemental antimony.

Partially eclipsed conformation is observed between the six ethyl groups. In contrast to a previously assigned $^{31}$P NMR chemical shift (+21 ppm)$^{29}$ for [11(ET)][ClO$_4$]$_2$, which was not structurally authenticated, [11(ET)][OTf]$_2$ exhibits a $^{31}$P NMR chemical shift of +38.5 ppm.

The $^{31}$P and $^{19}$F NMR resonances attributed to fluorophosphonium cations [12(R)]$^{11}$ were confirmed by comparison to literature values or independent synthesis of triflate salts from small-scale equimolar mixtures of the appropriate phosphine with XeF$_2$ followed by treatment with one equivalent of TMSOTf. To the best of our knowledge, the solid-state structure of [12(Me)][OTf] (Fig. 4) represents the first structural characterization of a trialkyll fluorophosphonium salt, and involves three hydrogen bonds with the triflate anion in addition to one weak contact [3.301(2) Å] between a triflate oxygen atom and the phosphorus atom, which is marginally shorter than $\Sigma_{\text{rad}}$ for the two elements (3.320 Å).$^{24}$ The O–P–F angle generated by this contact is 177.34(8)$^c$, representing adjustment of the $D_{th}$ structure of Me$_2$PF$_2$.$^{43}$

The $^{31}$P NMR chemical shifts for species in Table 2 over a 150 ppm range but within each class of cations, generally decrease in the order $\delta$(R = Et) > $\delta$(R = Pr) > $\delta$(R = Bu) > $\delta$(R = Me). Notably, the chemical shifts of the free phosphines (range of 30 ppm) also show the same order, providing additional support for the proposed assignments (Fig. S6, ESI†).

Attempts to isolate cations [7(R)]$^{+}$ or [10(R)]$^{+}$ with bulky phosphines such as PPr$_3$ were unsuccessful. A $^{31}$P NMR assay of the reaction mixture containing PPr$_3$ and FSb(OTf)$_2$ in a 2 : 1 ratio displayed numerous fluorine containing products as indicated by the observation of spin system with F–P couplings but no pure compounds could be isolated. A 3 : 1 mixture of PPr$_3$ with Sb(OTf)$_3$ also gave a complex mixture of products at room temperature which could not be separated. Deprotonation of MeCN solvent was observed upon refluxing the reaction mixture for short periods or stirring at room temperature for 16 hours. We conclude that steric bulk at the α-carbon of the phosphine hinders the coordination required for clean transformation of bis-phosphine cations [7(R)]$^{+}$ to tris-phosphine cations [8(R)]$^{+}$.

While the initial isolation of [10(Me)][OTf]$_4$ as a pure substance was achieved on a 150 mg scale (ca. 0.1 mmol), making it unamenable to reactivity studies, reaction conditions have now been optimized for a one-pot, three-step reaction (ESI) to give reproducible yields of analytically pure [10(Me)][OTf]$_4$ and [10(ET)][OTf]$_4$ on a scale up to 10 g. Consistent with the exquisite sensitivity of these compounds towards hydrolysis and oxidation, particularly in solution, the key determinant of purity and reactions yields is the rigorous drying and deoxygenation of the solvent and careful application of dynamic vacuum (ca. 10$^{-1}$ mbar) in the latter stages of the reaction to avoid free phosphine-catalyzed decomposition (vide infra).

**Thermolysis and photoysis of [10(Me)][OTf]$_4$**

The four-membered ring of [10(Me)]$^{14+}$ contains four of the six Sb–Sb bonds required to make neutral, tetrahedral Sb$_4$, which is directly analogous to P$_4$ and As$_4$. Moreover, [10(Me)]$^{14+}$ also contains four phosphine ligands which may be susceptible to further reductive elimination of two diphosphonium dications, [11(ET)][OTf]$_2$, to yield neutral Sb$_4$. While P$_4$ and As$_4$ are well characterized, Sb$_4$ has not been isolated as a bulk solid, and only one solid-state structural determination has been made using a scanning tunnelling microscope to characterize a thin film of Sb$_4$ under ultrahigh-vacuum conditions.$^{44}$ In this context, we envisioned the thermal or photochemical decomposition of [10(Me)][OTf]$_4$ as a route to bulk solid Sb$_4$.

A sample of solid [10(Me)][OTf]$_4$ (yellow-colored) heated under argon at 120 °C for 16 hours turned black, consistent with the formation of elemental antimony (Scheme 5). A CD$_3$CN extract of the black product showed $^{31}$P, $^1$H and $^{13}$C NMR signals corresponding exclusively to [11(ET)][OTf]$_2$ as the sole oxidation product. A Raman spectrum of the black solid (Fig. S7, ESI†) matched that of the amorphous α-phase (110 cm$^{-1}$, 150 cm$^{-1}$)$^{45}$ of antimony rather than the reported Raman spectrum of tetrahedral Sb$_4$ in argon matrix (138 cm$^{-1}$, 179 cm$^{-1}$, 242 cm$^{-1}$).$^{46}$ Identical results were obtained when heating was carried out in the dark, under vacuum, or in solution (toluene). Irradiating solid [10(Me)][OTf]$_4$ or as a solution in MeCN at 256 nm for 3 hours at room temperature had no measurable effect. It should be noted that in the gas phase tetrahedral Sb$_4$ is the preferred allotrope of the element up to 1050 K.$^{44}$ It is possible that despite its gaseous stability, tetrahedral Sb$_4$ is thermodynamically unstable with respect to...
its amorphous phases in the condensed state, preventing its isolation as a solid and is, in this context, analogous to tetrahedral As₄ (yellow arsenic) which spontaneously decomposes to a hexagonal allotrope (grey arsenic, α-As) at room temperature.⁸⁸

The thermolysis described above must be carried out in rigorously dried glassware, the surface of which has been treated with Me₃SiCl to silanize terminal –OH groups. Samples heated without prior passivation of glassware produced elemental antimony and [11(Me)][OTf]₄, but also showed resonances due to [Me₃PH]⁺ and a singlet at +115.6 ppm in the ³¹P [¹H] NMR spectrum (CD₃CN) of the reaction mixture, consistent with formation of [Me₃POPMe₃]²⁺. This assignment is supported by an independent synthesis from a 2 : 1 mixture of Me₃PO and triflic anhydride, using a well-established protocol for these reagents.⁹⁹ We interpret the formation of these by-products as being due to the reaction of the extremely moisture sensitive [10(Me)][OTf]₄ with surface hydroxyl groups in non-silanized glassware.

Reactions of [10(Me)][OTf]₄ with RₓPX(3–n); X = H, Cl; n = 1, 2

Addition of a solution of RₓPH (R = C₆H₁₁ (Cy), Ph) to a clear yellow-colored solution of [10(Me)][OTf]₄ in MeCN results in immediate deposition of a fine black precipitate and loss of the yellow coloration. The ³¹P [¹H] NMR spectra of reaction supernatants show a singlet due to [Me₃PH]⁺ and two doublets characteristic of phosphonophosphonium cations [Me₃PPR₂]⁺ (R = Cy, Ph) with typical ¹J_P_P values in the 300–350 Hz range (Scheme 6).⁹⁹ Cation [Me₃PPH₂]⁺ is known²⁰ and the assignment of [Me₃PPCy₂]⁺ was confirmed by comparison of chemical shifts and coupling constants with literature values²⁰ for phosphonophosphonium salts and by elemental analysis.

Analogously, addition of a solution of RPH₂ (R = Cy, butyl) to a solution of [10(Me)][OTf]₄ results in immediate precipitation of elemental antimony. The ³¹P NMR spectra of these reaction mixtures show complete consumption of RPH₂ and [10(Me)][OTf]₄, and formation of a singlet due to [Me₃PH]⁺ and a pair of doublets assigned to [Me₃PP(H)R]⁺ (Scheme 7). Consistent with this formulation, the ³¹P–¹H coupled NMR spectrum of the reaction involving CyPH₂ shows (Fig. 5a) both ¹J_P_P and ²J_P_H couplings for the phosphinic signal centered at −83.6 ppm. The Pₓ–Pₓ(Hₓ)Cy connectivity is also confirmed in the ¹H NMR spectrum of the reaction mixture (Fig. S8, ESI†), where H₀ resonates at +3.65 ppm exhibiting ¹J_H_Pₓ, ²J_H_Pₓ, and ³J_H_Hₓ couplings, the last of these arising from coupling to the ipso proton (H₁) of the cyclohexyl ring. The methyl protons (H₂) around Pₓ also show the expected ²J_H_Pₓ and ³J_H_Pₓ couplings, indicating a P–P bond. Finally, a two-dimensional ³¹P/¹H HSQC (Fig. 5b) spectrum, which was optimized to show one-bond couplings, shows coupling between H₀ and Pₓ but no coupling involving H₂ and Pₓ. The corollary two-dimensional HMBC experiment (Fig. 5c), optimized to exclude one-bond couplings, shows coupling between H₀ and Pₓ but no coupling involving H₂ and Pₓ. Despite numerous attempts, it was not possible to separate [Me₃PP(H)Cy][OTf] from [Me₃PH][OTf], precluding elemental analysis or structural determination by X-ray diffraction. Nevertheless, to the best of our knowledge this is the first spectroscopic detection of an H-phosphinophosphonium cation.

The formation of [Me₃PH]⁺ and the phosphinophosphonium salts is understood in broad terms as a metathesis step followed by a reductive elimination step as outlined in Scheme 6. We speculate that coordination of Cy₂PH to one of the antimony centres in [10(Me)][OTf]₄ is followed by intramolecular deprotonation by PMe₃ to yield the observed [Me₃PH]⁺ cation and a tricationic intermediate, [A]³⁺. This trication can undergo rapid intramolecular reductive elimination of the first equivalent of the phosphinophosphonium cation to give dication [B]²⁺. A second round of coordination, deprotonation and reductive elimination completes the reduction of antimony to its elemental form and furnishes the observed distribution of products. Unfortunately, the partially reduced species were not observed and appear to be fleeting intermediates. Nevertheless, formation of [Me₃PP(H)R]⁺ from reactions involving primary phosphines (Scheme 7) is consistent with the proposed mechanism, although it is unclear why the second deprotonation does not occur to yield the corresponding dication [(Me₃P)₂PR]²⁺. As before, Raman analysis of the black

Scheme 6 Formation of [Me₃PH]⁺ and [Me₃PPCy₂]⁺ from the reaction of [10(Me)][OTf]₄ with Cy₂PH.
precipitate matches the amorphous α-phase of metallic antimony rather than pyramidal Sb₄.

The observation that [10(Me)][OTf]₄ serves as a source of PMe₃, which deprotonates added primary and secondary phosphines, implies a labile and polarized P–Sb bond that undergoes facile heterolytic cleavage. Consistently, addition of Cy₂PCl or CyPCl₂ to a solution of [10(Me)][OTf]₄ results in quantitative formation of [Me₃PPCy₂]⁺ or [(Me₃P)₂PCy]⁺, respectively, concomitant with deposition of elemental antimony (Scheme 8). In these cases, [10(Me)]⁺ behaves overall as a chloride abstractor and phosphine donor. We tentatively propose formation of chloroantimony species as transients that undergo loss of chlorine gas to yield elemental antimony as there is no evidence of Sb–Cl bond stretching modes in the Raman spectra of the insoluble black solid isolated from these reactions. However, since no products expected from reactions of dissolved Cl₂ could be detected, the fate of the chlorine atoms cannot yet be definitively described. When intermediate stoichiometries of CyPCl₂ are employed, formation of the known [Me₃PP(Cl)Cy]⁺ cation is also observed, indicating a single chloride abstraction event, that is analogous to the formation of [Me₃PP(H)Cy]⁺ in reactions with CyPH₂. ³¹P NMR data for Scheme 7 Reaction of CyPH₂ with [10(Me)][OTf]₄.

Fig. 5 (a) ³¹P NMR spectrum of the crude reaction mixture containing CyPH₂ and [10(Me)][OTf]₄ in a 2 : 1 ratio. Insets show detailed views of the ³¹P(¹H) and ³¹P NMR resonances assigned to the Me₃P⁻ (left) and -P(H)Cy (right) fragments in [Me₃PP(H)Cy]⁺. A list of coupling constants deduced from the combination of ³¹P and ¹H NMR is also given. (b) Sections of the ³¹P/¹H HSQC spectrum showing a ¹JPH coupling between P bè and H bè. (c) Sections of the ³¹P/¹H HMBC spectrum showing a ²JPH coupling between P α and H bè.
The reaction of $[10(Me)]^{4+}$ with PMe$_3$ shows slow disappearance of the signal due to the latter and evolution of broadened signals due to $[11(Me)]^{2+}$ and free PMe$_3$. Concomitantly, a mirror of antimony is deposited in the reaction vessel. Within 12 hours at 298 K, there is no evidence of $[10(Me)]^{4+}$, while signals due to $[11(Me)]^{2+}$ and free PMe$_3$ persist, consistent with complete decomposition of the tetracation, catalyzed by PMe$_3$. The proposed mechanisms (Scheme 9) involve nucleophilic attack by the added phosphine at either the antimony or the phosphorus center.

Scheme 8  Formation of $[Me_3PPCy_2]^{1+}$, $[Me_3PP(Cl)Cy]^{1+}$, and $[[Me_3P]_2PCy]^{2+}$ from the reaction of $[10(Me)][OTf]_4$ with Cy$_2$PCl or CyPCl$_2$.

Table 3  $^{31}$P NMR (CD$_3$CN, 298 K) chemical shifts and coupling constants for products obtained from the reaction of Cy$_2$PH, CyPH$_2$, Cy$_2$PCL, and CyPCl$_2$ with $[10(Me)][OTf]_4$

<table>
<thead>
<tr>
<th></th>
<th>$^{31}$P (ppm)</th>
<th>$J_{PP}$ (Hz)</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>$[Me_3PPCy_2][OTf]$</td>
<td>$+12.8$, $-5.1$</td>
<td>327</td>
<td>This work</td>
</tr>
<tr>
<td>$[Me_3PP(H)Cy][OTf]$</td>
<td>$+14.8$, $-83.6$</td>
<td>252</td>
<td>This work</td>
</tr>
<tr>
<td>$[[Me_3P]_2PCy][OTf]_2$</td>
<td>$+22.7$, $-30.8$</td>
<td>307, 326</td>
<td>52</td>
</tr>
<tr>
<td>$[Me_3PP(Cl)Cy][OTf]$</td>
<td>$+23.0$, $+78.4$</td>
<td>326</td>
<td>53</td>
</tr>
</tbody>
</table>

$^a$ $J_{PH} = 214$ Hz.

Scheme 9  Catalytic decomposition of $[10(Me)][OTf]_4$ by PMe$_3$ via nucleophilic attack at Sb (upper half) or P (lower half).

Reaction of $[10(Me)][OTf]_4$ with PMe$_3$

The $^{31}$P NMR spectrum of a reaction mixture containing 15 mol% of PMe$_3$ and $[10(Me)][OTf]_4$ shows slow disappearance of the signal due to the latter and evolution of broadened signals due to $[11(Me)]^{2+}$ and free PMe$_3$. Concomitantly, a mirror of antimony is deposited in the reaction vessel. Within 12 hours at 298 K, there is no evidence of $[10(Me)]^{4+}$, while signals due to $[11(Me)]^{2+}$ and free PMe$_3$ persist, consistent with complete decomposition of the tetracation, catalyzed by PMe$_3$. The proposed mechanisms involve nucleophilic attack by the added phosphine at either the antimony or the phosphorus center.
phosphorus centres. Attack at a stibine should yield intermediate [A]$^{3+}$ (Scheme 9), featuring a hypercoordinate antimony centre. Several examples of such hypervalent P-Sb complexes have been reported. Due to its high charge concentration, this complex is predicted to be strongly oxidizing, and, in a process analogous to reductive elimination from $[8\text{Me}]^{3+}$, an equivalent each of $[11\text{Me}]^{2+}$ and intermediate $[B]^{2+}$ (Scheme 9) can be generated, enabling dissociation of PMe$_3$. Alternatively, attack at one of the phosphorus centres of $[10\text{Me}]^{4+}$ directly generates intermediate $[B]^{2+}$ together with $[11\text{Me}]^{2+}$ and PMe$_3$. The liberated phosphine can further reduce $[B]^{2+}$ by a second nucleophilic attack either at Sb or P to evolve the second equivalent of $[11\text{Me}]^{2+}$ and yield fully reduced antimony. Nucleophilic attack by a neutral two-electron ligand at tetracoordinate trimethylchlorophosphonium, trimethylphosphonium and dimethylthiophosphonium cations has been demonstrated previously. The broadness of signals for $[11\text{Me}]^{2+}$ and PMe$_3$ in these reaction mixtures is attributed to an exchange process that is also detected when free PMe$_3$ is added to a solution of $[11\text{Me}]^{[\text{OTf}]_3}$. The catalytic decomposition of $[10\text{Me}]^{[\text{OTf}]_3}$ in the presence of PMe$_3$ explains the difficulties encountered during synthesis of this salt. For instance, if the addition rate of PMe$_3$ to FSB(OTf)$_2$ is too high, a dark orange solution is obtained which rapidly deposits elemental antimony (see note in Experimental section). However, if a dynamic vacuum is applied to the dark orange solution to remove the volatile PMe$_3$ (b.p. $= 38^\circ$C), the solution maintains a yellow colour, leading to the formation of $[10\text{Me}]^{[\text{OTf}]_4}$. Moreover reactions with Lewis bases that displace PMe$_3$ must be carried out with explicit steps to remove the liberated phosphine in order to avoid decomposition (vide infra).

Reaction of $[10\text{Me}]^{[\text{OTf}]_3}$ with $[\text{Li}]^{[\text{nacnac}](dipp)}$

In contrast to the sterically unhindered and neutral base PMe$_3$, a bulky and anionic base is expected to yield products arising from ligand substitution rather than from addition. Consistently, the $^{31}\text{P}^\text{[H]}$ NMR spectra of equimolar reaction mixtures of $[10\text{Me}]^{[\text{OTf}]_4}$ and $[\text{Li}]^{[\text{nacnac}](dipp)}$ (dipp $= 2,6$-disoproplyphenyl), indicate quantitative formation of $[15\text{Me}]^{[\text{OTf}]_3}$ (Scheme 10). The 1,3-diketiminate anion $[\text{nacnac}(dipp)]^{1-}$, abbreviated as nacnac, displaces one PMe$_3$ ligand from $[10\text{Me}]^{3+}$ to give $[[\text{Me}_3\text{P}]_3\text{Sb}_4[\text{nacnac}]^{3+}$ (Scheme 9), which is an analogue of $[[\text{Me}_3\text{P}]_3\text{Sb}_4[\text{PCy}_3^{3+}$ (Scheme 10, intermediate [A]$^{1+}$ in Scheme 6). The $^{31}\text{P}^\text{[H]}$ NMR spectrum (Fig. 6) of $[15\text{Me}]^{[\text{OTf}]_3}$, shows the expected AX$_2$ spin system $[26.6 \text{ ppm (triplet)}, -33.6 \text{ ppm (doublet)}, J_{PP} = 32 \text{ Hz}]$ and a corresponding AX$_2$ spin system $[6.3 \text{ ppm (triplet)}, -2.5 \text{ ppm (doublet)}, J_{PP} = 23 \text{ Hz}]$ is also observed for $[15\text{Et}]^{3+}$, prepared from the reaction of $[10\text{Et}]^{4+}$ with $[\text{Li}]^{[\text{nacnac}](dipp)}$. Isolation of $[15\text{Me}]^{[\text{OTf}]_3}$ is only possible when the reaction is performed under a mild dynamic vacuum to remove the displaced phosphine, which effects redox decomposition at high concentrations, presumably via similar mechanisms as described above for $[10\text{Me}]^{4+}$.

The solid-state structure of the cation in $[15\text{Me}]^{[\text{OTf}]_3}$, MeCN (Fig. 7) shows three phosphine ligands and the rare $\gamma$-coordination mode for the nacnac substituent, which, to the best of our knowledge, has not been observed for haloantimony centres bound to this substituent. Heteroleptic substitution is very rare in antimony homocycles and examples for cationic systems have not been reported. The range of Sb–Sb [2.8209(5)–2.8612(5) Å] and Sb–P [2.538(5)–2.604(9) Å] distances are similar to those in...
indicating minimal distortion of the Sb₄ ring upon displacement of PMe₃ with nacnac. While [15(Me)][OTf]₃ is stable in the solid state under inert atmosphere, ³¹P{¹H} NMR spectra of MeCN solutions show decomposition over five days at 20 °C to elemental antimony, [10(Me)][OTf]₄ and [11(Me)][OTf]₂ (Fig. S9, ESI†).

To assess whether or not bonding via the γ carbon of nacnac is a general feature of antimony compounds and because nacnac functionalized antimony centers are rare in the literature, we also prepared (nacnac)Sb(OTf)₂ by salt metathesis between an equimolar mixture of in situ generated Sb(OTf)₃ and [Li][nacnac(dipp)]. Upon removal of LiOTf, the compound was isolated as a pure substance and comprehensively characterized. The molecular structure of (nacnac)Sb(OTf)₂, determined by X-ray diffraction, shows a see-saw geometry around antimony with two strongly-interacting triflate anions in axial positions (Fig. S10, ESI†). In contrast to γ-coordination observed for [15(Me)]³⁺, N,N'-chelation is observed for (nacnac) Sb(OTf)₂, and we attribute the difference in bonding modes to the different steric environments around antimony in the two compounds, rather than intrinsic features of the nacnac-Sb interaction.

Interestingly, the ¹⁹F resonances for the two triflate CF₃ groups in (nacnac)Sb(OTf)₂ are different (−78.3 and −78.4
ppm), implying a rigid ring system with non-equivalent positions above and below the plane of the ring. Consistently, the isopropyl substituents show two unique resonances for the C\textsubscript{ipso} protons. Furthermore, there is restricted rotation around the C\textsubscript{ipso}-C phenyl bond giving rise to four unique signals for the methyl groups in the 1H NMR spectrum of the compound. We speculate that this is due to solution-phase persistence of the weak hydrogen bonding interactions between the nitrogen atoms and the isopropyl C\textsubscript{ipso} protons, detected as short contacts in the solid state molecular structure (Fig. S10, ESI†).

Reaction of [10(Me)]\textsubscript{4+} with dmap
The reaction of [10(Me)]\textsubscript{4+} with 4-dimethylaminopyridine (dmap) has been examined by 31P NMR (Fig. 8) and shows displacement of one phosphine ligand by dmap (Scheme 11). It was not possible to isolate the resulting products. Following filtration of the reaction mixture (black suspension), the yellow-green filtrate shows the expected AX\textsubscript{2} spin system (triplet at +66.9 ppm, doublet at +42.5 ppm, \(J_{PP} = 24\) Hz), tentatively assigned to [Me\textsubscript{3}P\textsubscript{3}Sb\textsubscript{4}(dmap)]\textsuperscript{4+} ([16]\textsuperscript{4+}), and broad signals due to PMe\textsubscript{3} (+54 ppm), [Me\textsubscript{3}P(dmap)]\textsuperscript{2+} ([17]\textsuperscript{2+}, +89.0 ppm) and [11(Me)]\textsuperscript{2+}. Within hours, signals due to [Me\textsubscript{3}PCH\textsubscript{2}PMe\textsubscript{2}\textsuperscript{1+} ([18]\textsuperscript{1+}, doublet at +26.0 ppm, \(J_{PP} = 58\) Hz) appear in the 31P NMR spectrum and a significant amount of dmap\textsuperscript{H}\textsuperscript{+} is observed by 1H NMR spectroscopy. We propose that the latter two species arise from deprotonation of the slightly acidic protons of [11(Me)]\textsuperscript{2+} by dmap and the subsequent rearrangement of [Me\textsubscript{3}PPMe\textsubscript{2}CH\textsubscript{3}]\textsuperscript{1+} (Scheme 11). Consistently, a 1 : 1 control reaction of dmap and [11(Me)]\textsuperscript{2+} initially shows broad signals for [17]\textsuperscript{2+} and free PMe\textsubscript{3} as the kinetic products, but within 4 hours

![Scheme 11](image-url)

**Scheme 11** Proposed pathways to formation of [16]\textsuperscript{4+} (a), [17]\textsuperscript{2+}, [11(Me)]\textsuperscript{2+}, elemental antimony (b and d), and [18]\textsuperscript{1+} (c) in reaction mixtures containing [10(Me)]\textsubscript{4+} and dmap in a 1 : 1 stoichiometry.

![Scheme 12](image-url)

**Scheme 12** Kinetic and thermodynamic pathways in the reaction between [11(Me)]\textsuperscript{2+} and dmap.

**Table 4** Comparison of 31P NMR chemical shifts for some phosphorus containing main-group cations stabilized by PMe\textsubscript{3} or dmap. Values for tetracoordinate phosphorus centers are given in parentheses, where applicable.

<table>
<thead>
<tr>
<th>31P NMR Reference</th>
<th>31P NMR</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>[Me\textsubscript{3}PMe\textsubscript{3}]\textsuperscript{1+}</td>
<td>+18 (−59)</td>
<td>27</td>
</tr>
<tr>
<td>[Me\textsubscript{3}P(dmap)]\textsuperscript{1+}</td>
<td>+91</td>
<td>54</td>
</tr>
<tr>
<td>[Ph\textsubscript{3}PMe\textsubscript{3}]\textsuperscript{2+}</td>
<td>−23 (+15)</td>
<td>51</td>
</tr>
<tr>
<td>[Ph\textsubscript{3}P(dmap)]\textsuperscript{2+}</td>
<td>+88</td>
<td>55</td>
</tr>
<tr>
<td>[Me\textsubscript{3}SPPMe\textsubscript{3}]\textsuperscript{1+}</td>
<td>+16 (+38)</td>
<td>56</td>
</tr>
<tr>
<td>[Me\textsubscript{3}SPP(dmap)]\textsuperscript{1+}</td>
<td>+88</td>
<td>56</td>
</tr>
<tr>
<td>[11(Me)]\textsuperscript{2+}</td>
<td>(+28.4)</td>
<td>27</td>
</tr>
<tr>
<td>[17]\textsuperscript{2+}</td>
<td>(+89.0)</td>
<td>54</td>
</tr>
<tr>
<td>[10(Me)]\textsuperscript{3+}</td>
<td>(−24.5)</td>
<td>This work</td>
</tr>
<tr>
<td>[15(Me)]\textsuperscript{3+}</td>
<td>(−26.6), (−33.6)</td>
<td>This work</td>
</tr>
<tr>
<td>[13(Et)]\textsuperscript{3+}</td>
<td>(−2.5), (−6.3)</td>
<td>This work</td>
</tr>
<tr>
<td>[16]\textsuperscript{4+}</td>
<td>(+66.9), (+42.5)</td>
<td>This work</td>
</tr>
<tr>
<td>[18]\textsuperscript{1+}</td>
<td>(+26.0), −53.9</td>
<td>60</td>
</tr>
</tbody>
</table>

**Fig. 9** 31P{1H} NMR spectra (CD\textsubscript{3}CN, 298 K) of (a) [10(Me)]\textsuperscript{4+}, (b) [10(Et)]\textsuperscript{4+}, (c) 1 : 1 mixture of [10(Me)]\textsuperscript{4+} and [10(Et)]\textsuperscript{4+}, and (d) 1 : 1 mixture of [10(Me)]\textsuperscript{4+} and PEt\textsubscript{3}. Symbols denote tentative assignments for [10(Me)]\textsubscript{2}(Et\textsuperscript{3})\textsuperscript{2+} (x), cis-[10(Me)]\textsubscript{2}(Et\textsubscript{2})\textsuperscript{4+} (O), trans-[10(Me)]\textsubscript{2}(Et\textsubscript{2})\textsuperscript{4+} ( ), and [10(Me) (Et)]\textsuperscript{3+} ( •).
signals due to $^{18}1^+$ and $[\text{dmap}]^{1+}$ are observed, revealing them to be the thermodynamic products (Scheme 12, Fig. S11 in ESI†). In addition to $^{18}1^+$, four unique and mutually coupled phosphorus environments (by $^{31}P$ NMR spectroscopy) are also observed which could not been assigned definitively.

The $^3J_{PP}$ coupling constant for the signal assigned to $^{16}4^+$ (24 Hz) is comparable to the values in $^{15}(\text{Me})^3^+$ (32 Hz) and $^{15}(\text{Et})^3^+$ (23 Hz). However, the $^{31}P$ NMR chemical shifts observed for $^{16}4^+$ (A: +66.9, X₂: +42.5) are significantly downfield from those of $^{15}(\text{Me})^3^+$ and $^{15}(\text{Et})^3^+$ (Table 4), and this cannot be attributed solely to the different formal charges in the species as the PMe₃ groups in tetracationic $^{10}(\text{Me})^4^+$ resonate at −24.5 ppm. We propose that dmap-stabilized main-group cations generally show $^{31}P$ NMR chemical shifts that are substantially downfield from their PMe₃-stabilized homologues (Table 4) due to the greater electronegativity of nitrogen relative to phosphorus, supporting the assignment for $^{16}4^+$.

**Reaction of $[10(\text{Me})][\text{OTf}]_4$ with $[10(\text{Et})][\text{OTf}]_4$**

Neutral catena-antimony rings are known to participate in ring-ring equilibria unless bulky substituents or dilute solutions are employed. For instance, solutions of hexaphenylcyclohexastibine (Ph₆Sb₆) equilibrate to give a mixture of four-, five-, and six-membered rings suggesting labile Sb–Sb bonds.¹¹

To assess the possibility of preparing heteroleptic derivatives of $[10(R)][\text{OTf}]_4$, pure samples of $[10(\text{Me})][\text{OTf}]_4$ and $[10(\text{Et})][\text{OTf}]_4$ were combined in a 1:1 stoichiometry. The $^{31}P$ NMR spectrum (Fig. 9c) of the resulting mixture suggests formation of multiple constitutional isomers of $[(\text{PMe}_3)_x(\text{PET}_3)(4/C_0^x[Sb]_4)]^4^+$, implicating a scrambling process in the two ring systems via Sb–Sb or P–Sb bond cleavage. A scrambling process involving Sb–Sb cleavage has been described previously for distibines.⁶² However, a control experiment, where free PET₃ was added to $[10(\text{Me})][\text{OTf}]_4$, also showed (Fig. 9d) formation of these isomers. Therefore a nucleophilic displacement pathway, where a bound PR₃ ligand is displaced by an added PR₃ ligand, cannot be precluded. However

**Scheme 13** Proposed formation of constitutional isomers from the equimolar reaction of $[10(\text{Me})]^4^+$ and $[10(\text{Et})]^4^+$.

**Scheme 14** Reactivity of a prototypical cyclo-tetra(stibinophosphonium) tetracation, $[10(\text{Me})]^4^+$. See text for descriptions of a–i.
this displacement route to heteroleptically substituted derivatives also yields significant amounts of [11(Me)]2+ and elemental antimony, presumably due to free PMe3 catalyzed decomposition of [10(Me)][OTf]4 as described earlier. Although, it has not yet been possible to purify these reaction mixtures and isolate the first examples of heteroleptically substituted catena-antimony rings, signal multiplicities consistent with AmSx, Asx3, and AA'XX' spin systems are observed, as expected from a mixture of [10(Me)]2[Et]4+, cis/trans-[10(Me)]2[Et]3+, and [10(Me)]2[Et]2+ (Scheme 13). Moreover, the coupling constants lie in the 21–26 Hz range and are comparable to J_{Pn} coupling constants detected in [15(Me)]4+ (32 Hz), [15(Ph)]4+ (23 Hz), and [16]4+ (24 Hz). Collectively, these data enable a tentative assignment of the spectral features observed in Fig. 9.

Conclusions

The reductive elimination of diphosphonium dications [11(R)]2+ from trialkylphosphine complexes of highly electrophilic antimony(III) centres is reported. The reduced antimony(i) fragments cyclize into frameworks identified as cyclo-tetra(stibinophosphonium) tetracations, [10(R)]4+. As outlined in Scheme 3, a phosphine catalyzed mechanism is proposed for fluoroantimony complexes, and isolation or spectroscopic characterization of key mechanistic intermediates is presented. The scope of this reductive assembly is dependent upon the reactivity studies that are summarized in Scheme 14. The reductive elimination of diphosphonium dications [11(Me)]2+ is directed by two features: (i) high charge concentration, and (ii) the presence of strongly polarized P–Sb bonds. The former explains the electrophility of cation [10(Me)]4+, its thermolysis to extrude [11(Me)]3+, and the observed facility for reductive elimination to yield elemental antimony (Scheme 14, reactions a–f). The significant polarization of the P–Sb bonds enables activation of a wide spectrum of bonds with the unusual outcome of yielding the same products via reaction with oppositely polarized substrates (e.g. P–Cl and P–H containing reagents) (Scheme 14, reactions c–f). This unique feature has led to the spectroscopic detection of the an H-phosphino-phosphonium cation, [Me3PP(H)Cy]2+, examples of which have not been reported previously. The high P–Sb bond polarization also supports a coordinate bonding model, consistent with ligand displacement reactivity demonstrated for cation [10(Me)]4+ (Scheme 14, reactions g–i). Ligand displacement has permitted functionalization of the four-membered Sb ring with substituents such as [nacnac][2]− or dmap (transiently). A heteroleptic phosphine substitution pattern around the Sb4 is feasible, but multiple isomers are observed on a relatively shallow potential energy surface hindering the isolation of a single derivative.

Within the broader context of phosphines as ubiquitous ligands in coordination chemistry, evidence of a novel ligand activation pathway has been presented and the associated reactants and products characterized. Taken together with previous, albeit less definitive, detection of such reactivity,10,43 the observation of this reductive elimination pathway confirms that these prototypical ligands can behave simultaneously as reducing agents and stabilizing ligands, a feature that may be generally applicable for phosphine complexes of highly electrophilic acceptors across the periodic table. Diversification of this synthetic protocol may therefore provide access to more extensively catenated systems for antimony as well as other elements. As demonstrated for [10(Me)]4+, a unique and rich reaction chemistry can be expected, in addition to the potential for valuable emergent properties such as σ-bond conjugation and cooperative catalysis due to metal catenation.

Acknowledgements

We thank the Natural Sciences and Engineering Research Council (NSERC) of Canada and the Vanier Canada Graduate Scholarships Program for funding. We gratefully acknowledge financial support from the ERC (SynPhos 307616) for a six month research stipend for S.S.C. at the TU Dresden. We also thank Dipl.-Chem. Kai Schwedtmann for experimental assistance and Prof. Lisa Rosenberg for valuable discussion.

References


