Dalton Transactions

PAPER

Check for updates

Cite this: *Dalton Trans.*, 2024, **53**, 11762

Received 31st May 2024, Accepted 21st June 2024 DOI: 10.1039/d4dt01591f

rsc.li/dalton

Introduction

The early contributions of Gutmann, Olah and Gillespie on antimony-containing Lewis acids demonstrated their reaction potential for synthesis and formation of donor-acceptor systems.¹ It might seem obvious to integrate antimony-containing reagents as Lewis acidic functions into the chemistry of frustrated Lewis pairs (FLPs) and thus to extend the diversity of these intra- or intermolecular systems. The first known FLPs were mainly based on boron- or aluminium-containing functions as Lewis acidic parts and phosphorus- or nitrogen-containing functions as complementary Lewis basic parts. Owing to the suppressed acid/base adduct formation, FLPs can be used to activate C-H bonds, hydrogen and small molecules or for the reduction of CO₂.²⁻⁴ However, the range of reactive Lewis acid-base combinations has increased over the last 20 years.⁴⁻⁸ The application of weaker B Lewis acids and stronger N Lewis bases, the so called "inverse" FLP by Krempner et al.,^{3,9} provides a further impetus to the still expanding FLP chemistry.

Antimony, as a versatile Lewis acid function, exhibits a particularly high calculated fluoride ion affinity in both oxidation states V and III, depending on the substituents.¹⁰ Therefore it has a broader applicability as Lewis acid than other pnictogens.¹¹ In addition to the theoretical considerations men-

Oxidation-dependent Lewis acidity in chalcogen adducts of Sb/P frustrated Lewis pairs*

Jonas Krieft, Beate Neumann, Hans-Georg Stammler and Norbert W. Mitzel 吵 *

The reactions of the frustrated Lewis pair $(F_5C_2)_2SbCH_2P(tBu)_2$ with oxygen, sulphur, selenium and tellurium led to the mono-oxidation products $(F_5C_2)_2SbCH_2P(E)(tBu)_2$ (E = O, S, Se, Te). Further oxidation of these chalcogen adducts with tetrachloro-*ortho*-benzoquinone (o-chloranil) gave $(F_5C_2)_2Sb(CH_2)(\mu-E)P(tBu)_2\cdotCat^{Cl}$ ($Cat^{Cl} = o-O_2C_6Cl_4$) with a central four-membered ring heterocycle for E = O, S, and Se. For E = Te the elimination of elemental tellurium led to an oxidation product with two equivalents of *o*-chloranil, $(F_5C_2)_2SbCH_2P(tBu)_2\cdot2Cat^{Cl}$, which is also accessible by reaction of $(F_5C_2)_2SbCH_2P(tBu)_2$ with *o*-chloranil. The synthesised compounds were characterised by NMR spectroscopy and X-ray structure analyses, and the structural properties were analysed in the light of the altered Lewis acidity due to the oxidation of the antimony atoms.

> tioned above, Gabbaï et al. performed impressive experimental studies on e.g. perfluorinated stibonium cations and stiboranes, which are air stable and able to activate strong elementfluorine bonds or complex formaldehyde; both in aqueous media.^{12,13} In other contributions, Gabbaï, Matile and Cozzolino highlighted the strong binding and chelation ability of stibanes, distibanes and -stiboranes towards anions like halides.^{14,15} Our group has recently presented rigid systems, based on anthracene, substituted with strong Lewis acidic $Sb(C_2F_5)_2$ functions, which allow the chelate-binding of halide ions and various Lewis bases in the form of pnictogen bonds.¹⁶ In antimony-containing systems the Lewis acidity can be varied both by oxidation of the antimony atom from III to V and by the electron withdrawing substituents. The apparent influence of the oxidation state on the Lewis acidity of organoantimony compounds has been investigated in terms of the distinctness of the σ -holes.^{14,17} Gabbaï showed that both stiboranes and stibonium cations with sufficiently pronounced Lewis acidity can interact with oxygen atoms to form heterocycles (Scheme 1).18-20 Even with lower Lewis acidity of the antimony atom, the formation of such (four-membered) antimony- and chalcogen-containing heterocycles was demonstrated.²¹

> In previous work in our group, we gained insight into the interaction of tetrel-based Lewis acids, $(F_5C_2)_3ECH_2P(tBu)_2$ with E = Si, Ge, Sn, with oxygen. Depending on the Lewis acidity, a four-membered heterocycle is obtained while the backbone of the FLP is deformed (Scheme 1).^{8,22} In this work, we intended to investigate the intramolecular interaction of analogously structured, Sb-based Lewis acids in the oxidation states III and V with chalcogen atoms.

The C_2F_5 group as one of the most electron withdrawing organyl ligands, is particularly suitable to increase the Lewis

View Article Online

Chair of Inorganic and Structural Chemistry, Center for Molecular Materials CM₂, Faculty of Chemistry, Bielefeld University, Universitätsstraße 25, Bielefeld 33615, Germany. E-mail: mitzel@uni-bielefeld.de

[†]Electronic supplementary information (ESI) available. CCDC 2350426–2350434. For ESI and crystallographic data in CIF or other electronic format see DOI: https://doi.org/10.1039/d4dt01591f



Scheme 1 Different interaction motifs of a stiborane and a stibonium cation with oxygen atoms by Gabbaï^{19,20} and various structural motifs of FLP chalcogen adducts.^{8,22}

acidity of stibanes and stiboranes.¹¹ Naumann and Hoge, among others, introduced this substituent into pnictogenbased systems, resulting in highly reactive reagents.²³ We have used the C_2F_5 substituent to generate sufficient Lewis acidity in the aforementioned tetrel-based FLP.^{6–8} Especially the tincontaining FLP shows a broad spectrum of reactivity towards small molecules such as hydrogen, carbon dioxide or unsaturated systems.^{7,22,24}

Results and discussion

Recently we introduced the FLP $(F_5C_2)_2$ SbCH₂P $(tBu)_2$ (1).²⁵ Its antimony(m)-based Lewis acid function turned out to be relatively soft, in accordance with the qualitative predictions of the HSAB concept.²⁶

This is one of the reasons why we studied the reaction of the FLP with the different chalcogen elements (or substitutes) to explore the behaviour of **1** under oxidative conditions.

Because of the decomposition or over-oxidation that occurs on contact with molecular oxygen, N₂O was used instead. Under elimination of molecular nitrogen, the oxygen oxidation product $(F_5C_2)_2SbCH_2P(O)(tBu)_2$ (2) is formed in which only the phosphorus atom is oxidised (Fig. 1 and Scheme 2).

It appears that the strong electron-withdrawing pentafluoroethyl substituents make the antimony atom so electron deficient that only the phosphorus atom is oxidised. However, the X-ray diffraction experiments indicate no formation of a four-membered ring by Sb–O interactions in the solid state; we can conclude that the Lewis acidity of the antimony function is too weak to deform the conformation of the Sb–C–P–O backbone. Instead, the oxidation has no effect on the angle of the bridging Sb–C–P unit; it is identical in the crystal structures of 1 and 2.

It is therefore not surprising that the distance (2.784(3)/2.888(3) Å) between the Sb and O atom is less than the sum of their van der Waals radii ($\sum r_{vdW}(Sb-O) = 3.58 \text{ Å})^{27,28}$ but well



Fig. 1 Molecular structure of compound **2** in the solid state. The asymmetric unit contains two molecules, so two different values are given. Ellipsoids are set at 50% probability; hydrogen atoms are omitted for clarity. Selected bond lengths and atom distances [Å] and angles [°]: Sb(1)–C(1) 2.260(3), Sb(1)–C(3) 2.236(4), Sb(1)–C(5) 2.174(3), P(1)–C(5) 1.812(3), P(1)–C(6) 1.844(4), P(1)–C(10) 1.858(3), P(1)–O(1) 1.509(3)/1.504(3), Sb(1)····O(1) 2.784(3)/2.888(3); C(1)–Sb(1)–C(3) 94.8(1), C(1)–Sb(1)–C(5) 91.6(2), C(1)–Sb(1)–O(1) 153.9(1), C(5)–Sb(1)–C(3) 91.2(2), C(5)–P(1)–C(6) 107.7(2), C(5)–P(1)–C(10) 107.3(2), C(6)–P(1)–C(10) 114.4(2), O(1)–P(1)–C(5) 106.0(2), O(1)–P(1)–C(6) 111.6(2), O(1)–P(1)–C(10) 109.3(2), P(1)–C(5)–Sb(1) 100.7(2)/102.2(2), Sb(1)–C(5)–P(1)–O(1) 19.5(1)/22.9(1).



Scheme 2 Overview of the reactions of 1 with various chalcogens (or substitutes) and the subsequent oxidation of these products with o-chloranil, as well as the direct reaction of 1 with o-chloranil.

above the sum of their covalence radii ($\sum r_{cov}(Sb-O) = 2.03 \text{ Å}$).²⁹ Another indicator for a possible Sb…O interaction is the P–O bond length, which should be longer if the Lewis acid interacts with the oxygen atom. For comparison we examined trimethyl-phosphane oxide and tris(*tert*-butyl)phosphane oxide (as a similar reference without the acid function). In trimethyl-phosphane oxide the P–O bond is shorter than in 2. Due to the greater steric demand of the *tert*-butyl groups, the P–O bond in the *tert*-butyl phosphane oxide; actually, it is identical to 2 (Table 1).

This underlines the independence of the Lewis acidic function and therefore excludes any strong Sb…O interaction. The C(1)–Sb(1)–O(1) angle of 153.9(1)/150.4(1)°, which should be closer to 180° for a σ -hole interaction, also supports this. In other molecules, such as $(F_5C_2)_2SbCH_2P(tBu)_2\cdot CS_2$, the CS₂

Table 1 Selected bond lengths and atom distances of the different chalcogen adducts of 1 and their oxidation adducts compared with the corresponding trimethylphosphane chalcogenides,^{30,31} tris(tert-butyl)phosphane chalcogenides³²⁻³⁵ and the oxygen and sulphur adducts of (C₂F₅)₃SnCH₂tBu₂²² with Cat^{Cl} = o-O₂C₆Cl₄. If two values are given, the asymmetric unit of the corresponding compound contains two molecules (or three for 7). (Please note, that $tBu_3P = 0$ is an adduct with H_2O_2 with a hydrogen bond between the P atom and one H atom of H_2O_2)

		<i>t</i> Bu₃P==E <i>d</i> (P==E) [Å]	$ \begin{array}{l} (C_2F_5)_2 \text{SbCH}_2 P(E) t B u_2 \ (2-5) \\ d(P = E) \left[\mathring{A} \right] \end{array} $	$(C_2F_5)_2Sb(\mu CH_2)(\mu - E)PtBu_2 \cdot Cat^{Cl} (6-8)$		$(C_2F_5)_3Sn(\mu-CH_2)(\mu-E)PtBu_2$	
Е	$Me_3P = E$ d(P = E) [Å]			d(Sb…E) [Å]	d(Р-Е) [Å]	d(Sb-Е) [Å]	d(Р-Е) [Å]
O S Se Te	$\begin{array}{c} 1.489(6)^{30} \\ 1.969(1)^{31} \\ 2.122(1)^{31} \\ - \end{array}$	$\begin{array}{c} 1.510(1)^{32} \\ 1.962(3)^{33} \\ 2.133(1)^{34} \\ 2.368(4)^{35} \end{array}$	$\begin{array}{c} 1.509(3)/1.504(3)\\ 1.982(2)/1.976(2)\\ 2.131(1)\\ 2.382(1) \end{array}$	2.784(3)/2.888(3) 3.145(2)/3.108(1) 3.171(1) 3.362(1)	1.567(1) 2.036(2)/2.031(2) 2.194(2)/2.187(2) —	2.158(1) 2.633(2)/2.619(2) 2.748(1)/2.703(1) —	$\begin{array}{c} 1.532(2)^{21} \\ 2.010(2)^{21} \\ \\ \\ \end{array}$

adduct of 1,²⁵ or anthracene-based poly Lewis acids with terminal $Sb(C_2F_5)_2$ functions,¹⁶ we have presented examples of this σ -hole interaction between a Lewis acidic antimony(III) function and Lewis bases.

The torsion angle of the Sb-C-P-E motif of 2 is 19.5(1)/22.9 (2)°, so the atoms involved are not in the same plane. In contrast to the previous synthesis of 2, the reactions of 1 with the elemental forms of sulphur, selenium and tellurium could be used to generate the corresponding oxidation products 3-5; $(F_5C_2)_2$ SbCH₂P(E)(tBu)₂ with E = S, Se, Te (Scheme 2 and Fig. 2). In all cases, only the phosphorus atom was oxidised. There was no formation of a four-membered ring, nor any significant deformation of the Sb-C-P backbone compared to 1, although the Sb-C-P angle in 2-5 increases slightly with increasing size of the chalcogen atom (Table 1).

The Sb---E distances are all well below the sum of their corresponding van der Waals radii ($\sum r_{vdW}(Sb-S) = 3.86$ Å, $\sum r_{vdW}(Sb-Se) = 3.96 \text{ Å}, \sum r_{vdW}(Sb-Te) = 4.12 \text{ Å})^{27,28}$ but also clearly above the sum of their respective covalence radii $(\sum r_{cov}(Sb-S) = 2.43 \text{ Å}, \sum r_{cov}(Sb-Se) = 2.56 \text{ Å}, \sum r_{cov}(Sb-Te) =$ 2.76 Å).²⁹ As mentioned above for the P-O bond length, the other P-E bond lengths in 3 and 4 are also slightly longer than in the trimethylphosphane chalcogenides.

Even shorter is the P-S bond length in tris(tert-butyl)phosphane sulphide. However, for 4 and 5 the P-E bond lengths are more similar to their corresponding tert-butyl analogues, respectively. In addition to the natural elongation of the P-E

bond with increasing atomic number of the chalcogen, the distance between the antimony and the chalcogen atom also increases (Table 1).

The oxygen oxidation products of the analogous tetrelbased FLPs, $(F_5C_2)_3ECH_2P(O)(tBu)_2$, form a four-membered heterocycle for E = Si and Sn. For E = Ge the same open-chain structural motif as described above for the Sb(III) compound 2 is obtained (Scheme 1).8 In the latter two cases the Lewis acidity is apparently insufficient to show a significantly strong interaction between the Lewis acid function and the chalcogen atom. The molecular structures in the solid-state are supposedly more informative than investigations in the liquid phase, because they are usually more close to the ground state structure of a molecule. As with the Si and Sn analogues, there is no additional splitting of the resonances of the C₂F₅ or tertbutyl substituents due to formation of the four-membered heterocycle.8,22

Since the oxidation of Sb(III) to Sb(v) generally increases the Lewis acidity, this should also increase the interaction strength between the Lewis acid function and the chalcogen atom in compounds 2-5. Therefore, we exposed the mono-oxidation products 2-5 to a different and stronger oxidant. Tetrachloroortho-benzoquinone (o-chloranil) is a well-established reagent for the oxidation of stibanes³⁶ and proved to be suitable for this purpose. The change in colour of the oxidant solution, which was deep red before it was added to the mono-oxidations products 2-5, allowed the reaction to be followed



Fig. 2 Molecular structures of compounds 3-5 in the solid state. If two values are given, the asymmetric unit contains two molecules. Ellipsoids are set at 50% probability; hydrogen atoms are omitted for clarity. Selected bond lengths and atom distances [Å] and angles [°]; negative value for torsion angle means rotation in opposite direction: 3: P-S 1.982(2)/1.976(2), Sb.-S 3.145(2)/3.108(1), P-C-Sb 106.8(2)/106.0(2), Sb-C-P-S 25.0(3); 4: P-Se 2.131(1), Sb…Se 3.171(1), P-C-Sb 107.6(1), Sb-C-P-Se -22.2(1); 5: P-Te 2.382(1), Sb…Te 3.362(1), P-C-Sb 109.1(1), Sb-C-P-Te 27.7(1).

macroscopically. As expected, the antimony atom was oxidised. The significant increase in acidity of the antimony site led to the formation of a four-membered SbCPE ring with strong deformation of the Sb–C–P backbone.

Dalton Transactions

Fig. 3 shows the molecular structure of compound **6** $((Cl_4C_6O_2)(F_5C_2)_2SbCH_2P(\mu-O)(tBu)_2)$ in the solid state; analogous structural motifs were obtained for compounds 7 (S) and **8** (Se).

Due to the ring closure, the Sb–C–P angles in the molecules **6–8** are significantly more acute than in their respective openchain precursors **2–5**. Along with this deformation and the for-



Fig. 3 Molecular structure of compound 6 in the solid state. Ellipsoids are set at 50% probability; hydrogen atoms are omitted for clarity. Selected bond lengths [Å] and angles [°] of 6: b(1)-C(1) 2.240(2), b(1)-C(3) 2.279(2), b(1)-C(5) 2.194(2), b(1)-O(1) 2.158(1), b(1)-O(2) 2.009(1), b(1)-O(3) 2.044(1), P(1)-C(5) 1.809(2), P(1)-C(6) 1.853(2), P(1)-C(10) 1.844(2), P(1)-O(1) 1.567(1); C(1)-Sb(1)-C(3) 93.2(1), C(1)-Sb(1)-C(3) 103.5(1), O(1)-Sb(1)-C(3) 93.2(1), C(1)-Sb(1)-C(3) 103.5(1), O(1)-Sb(1)-C(5) 71.2(1), O(1)-Sb(1)-O(2) 173.3(1), O(1)-Sb(1)-O(3) 93.1(1), O(2)-Sb(1)-C(1) 98.6(1), O(2)-Sb(1)-C(3) 82.6(1), O(2)-Sb(1)-C(3) 163.0(1), O(3)-Sb(1)-C(5) 92.1(1), C(5)-P(1)-C(6) 113.6(1), C(5)-P(1)-C(10) 111.3(1), C(6)-P(1)-C(10) 114.2(1), O(1)-P(1)-C(5) 97.0(1), O(1)-P(1)-C(6) 109.5(1), O(1)-P(1)-C(10) 109.9(1), P(1)-C(5)-Sb(1) 91.5(1), P(1)-O(1)-Sb(1) 100.0(1).

mation of the four-membered ring, the P–E bond lengths and the Sb–E distances have also undergone drastic changes (Fig. 4). The former are all longer than in the corresponding precursors 2–4 and also longer than in the tris(*tert*-butyl)phosphane chalcogenides. For 2 and 3, a comparison with the oxygen and sulphur adducts of the aforementioned Sn/P-FLP, which contain the same central structural motif, is possible: The P–E bond lengths are in a similar range in both cases (Table 1). The distances between the Lewis acidic Sb(v) atom and the chalcogen atom in compounds **6–8** are significantly shorter than in the corresponding precursors 2–4 and only slightly above the sum of the covalence radii of the atoms involved (the differences are: $\sum r_{vdW}(Sb,E) - d(Sb-E) = 0.13$ Å for **6**, 0.20 Å for 7, 0.19 Å for **8**).

An additional test, whether the ring formation could be precluded by donor solvents *e.g.* pyridine, demonstrated, that even in the presence of an excess of donor solvent, the fourmembered ring is formed by oxidation with *o*-chloranil.

In contrast, the Te oxidation product 5 behaves differently. Unlike reactants 2-4, the oxidation of 5 with o-chloranil does not lead to a four-membered ring, but to the elimination of tellurium. The result is compound 9, which contains a second o-chloranil unit that occupies the binding position between the Sb and P atoms of the FLP, resulting in a seven-membered ring (Fig. 5). The antimony atom is at the center of an octahedral coordination environment. The Sb-C-P angle, now part of the seven-membered ring, is smaller than in both reactant 5 and FLP 1; it has an angle of 118.4(2)°. As an alternative synthetic route to the unexpectedly formed compound 9, we found the reaction of 1 with two equivalents of o-chloranil. Interestingly, compound 9 is selectively formed at room temperature with both excess and deficient oxidant. To understand how the reaction proceeds, a solution of 1 was mixed with a dilute solution of o-chloranil under kinetic control at -60 °C. Theoretically, three different products are conceivable when one molecule of 1 reacts with one molecule of the oxidant. Surprisingly, neither the stiborane (product of the oxidation of the Sb atom) nor the phosphorane (product of the oxidation of



Fig. 4 Molecular structures of compounds 7 and 8 in the solid state. The asymmetric units contain more than one molecule, respectively. Ellipsoids are set at 50% probability; hydrogen atoms are omitted for clarity. Selected bond lengths and atom distances [Å] and angles [°]: 7: P–S 2.036(2)/2.031(2), Sb–S 2.633(2)/2.619(2), P–C–Sb 101.2(3), P–S–Sb 81.9(1); 8: P–Se 2.194(2)/2.187(2), Sb–Se 2.748(1)/2.703(1), P–C–Sb 102.5(3), P–Se–Sb 78.2(1)/79.4(1).





Fig. 5 Molecular structure of compound 9 in the solid state. Ellipsoids are set at 50% probability; hydrogen atoms are omitted for clarity. Only the non-disordered of the two molecules per asymmetric unit is depicted. Selected bond lengths [Å] and angles [°]: Sb(1)–C(1) 2.233(4), Sb(1)–C(3) 2.247(4), Sb(1)–C(5) 2.181(3), Sb(1)–O(1) 2.023(2), Sb(1)–O(2) 2.027(3), Sb(1)–O(3) 2.044(2), P(1)–C(5) 1.794(4), P(1)–C(6) 1.846(4), P(1)–C(10) 1.833(3), P(1)–O(4) 1.587(2); C(1)–Sb(1)–C(3) 93.7(2), C(1)–Sb(1)–C(5) 98.7(2), C(5)–Sb(1)–C(3) 89.7(2), O(1)–Sb(1)–C(1) 88.0(1), O(1)–Sb(1)–C(3) 94.5(2), O(1)–Sb(1)–C(5) 171.9(2), O(1)–Sb(1)–O(2) 81.2(1), O(1)–Sb(1)–O(3) 89.4(1), O(2)–Sb(1)–C(1) 169.1(1), O(2)–Sb(1)–C(3) 88.4(2), O(2)–Sb(1)–C(5) 92.0(2), O(2)–Sb(1)–O(3) 89.6(1), O(3)–Sb(1)–C(1) 89.1(1), O(3)–Sb(1)–C(3) 175.2(1), O(3)–Sb(1)–C(5) 86.1(2), C(5)–P(1)–C(6) 107.5(2), C(5)–P(1)–C(10) 110.6(2), C(6)–P(1)–C(10) 116.5(2), O(4)–P(1)–C(5) 113.8(2), O(4)–P(1)–C(6) 99.2(2), O(4)–P(1)–C(10) 109.0(2), P(1)–C(5)–Sb(1) 118.4(2).

the P atom) are formed, but rather compound **10**, in which a molecule of the oxidant is found in the binding pocket of **1**, forming the seven-membered heterocycle mentioned above (Fig. 6).

This is different from the behaviour described by Gabbaï *et al.* for the oxidation of a related Sb/P system, o-C₆H₄(PPh₂) (SbAr₂) with Ar = C₆H₅ and C₆F₅, in which only the antimony atom is oxidized and bears a chelating o-C₆Cl₄O₂ ligand after the reaction.¹³

The obvious reason for this difference is the extreme electronegativity of the C_2F_5 ligands (even if compared to C_6F_5), which makes the antimony(m) function in **1** so electron deficient, that the phosphorus atom becomes the site more easily oxidised. **10** is an adduct of FLP **1** with an *o*-chloranil molecule and has thus lost its reactive potential to bind and activate other molecules. Like **9**, **10** contains a seven-membered ring as a structural motif. The Sb–C–P angle of 113.6(1)/115.1(1)° is between that of free FLP **1** (110.6(1)°) and that of compound **9** (118.4(2)°). The same applies to the Sb-··P distance (**1**: 3.306(1) Å, **9**: 3.420(1) Å, **10**: 3.327(1)/3.352(1) Å). In contrast to the tetrahedrally surrounded phosphorus atom, the

Fig. 6 Molecular structure of compound 10 in the solid state. Ellipsoids are set at 50% probability; hydrogen atoms are omitted for clarity. Only one of the two molecules per asymmetric unit is depicted. Selected bond lengths [Å] and angles [°]: Sb(1)-C(1) 2.242(2), Sb(1)-C(3) 2.316(2), Sb(1)-C(5) 2.186(2), Sb(1)-O(1) 2.316(1), P(1)-C(5) 1.782(2), P(1)-C(6) 1.847(2), P(1)-C(10) 1.848(2), P(1)-O(2) 1.593(1); C(1)-Sb(1)-C(3) 90.2(1), C(1)-Sb(1)-C(5) 92.4(1), C(5)-Sb(1)-C(3) 83.5(1), O(1)-Sb(1)-C(3) 90.2(1), O(1)-Sb(1)-C(3) 156.8(1), O(1)-Sb(1)-C(5) 77.0(1), C(5)-P(1)-C(6) 108.2(1), C(5)-P(1)-C(10) 111.0(1), C(6)-P(1)-C(10) 116.3(1), O(2)-P(1)-C(5) 111.4(1), O(2)-P(1)-C(6) 99.2(1), O(2)-P(1)-C(10) 110.2(1), P(1)-C(5)-Sb(1) 113.6(1).

antimony atom is bisphenoidally surrounded due to the lone pair on the antimony atom in oxidation state III, as is also typical for other adducts of FLP $1.^{24}$

When *o*-chloranil is added to **10**, compound **9** is selectively formed. This is the second step in the reaction sequence of FLP **1** to **9** at room temperature with two equivalents of oxidant.

As with the bond lengths and angles of the molecular structures in the solid state of compounds **1–10**, some trends can be seen in the NMR spectroscopic chemical shifts of the substances in solution. Both the methylene protons and the resonance of the phosphorus nucleus are good indicators of the influence of different oxidants in these systems. The resonance of the methylene protons becomes increasingly low-field shifted as the atomic number of the chalcogen increases (Table 2). By oxidising the antimony atom, these proton reso-

Table 2	Selected NMR chemical shifts of the compounds 1-10
---------	--

Compound	¹ Η δ(C H ₂) [ppm]	³¹ Ρ δ [ppm]	Compound	¹ Η δ(C H ₂) [ppm]	³¹ Ρ δ [ppm]
1	2.17	15.9	6	4.38	88.4
2	2.66	63.7	7	4.68	78.9
3	2.79	76.6	8	4.92	73.2
4	2.93	73.0	9	3.09/2.89	110.0
5	3.18	45.4	10	2.90	111.4

nances undergo a significant shift of about 2 ppm towards low field. An opposite tendency, *i.e.* a shift towards the high field, is observed in the resonances of the phosphorus nuclei of **3–5**.

Compound 2 behaves contrary to this trend. The change in molecular symmetry caused by the oxidation with *o*-chloranil leads to the splitting of the pentafluoroethyl groups into two signal sets in ¹⁹F NMR for **7–9** and for **8** and **9** also the splitting into two signal sets for the *tert*-butyl groups in ¹H NMR.

Conclusions

We have found that the phosphorus atom in the FLP $(F_5C_2)_2$ SbCH₂P(tBu)₂ (1) is selectively oxidised by reaction with N₂O or the elemental chalcogens sulphur, selenium and tellurium. The solid state molecular structure data of the products $(F_5C_2)_2$ SbCH₂P(E)(*t*Bu)₂ (E = O, S, Se, Te; 2–5) do not indicate any interaction between the antimony(m) Lewis acid and the Lewis basic chalcogen atom; the different size and electrostatics of the bound chalcogen atoms do not seem to affect the nature of product formed. Increasing the Lewis acidity by oxidation of Sb(m) to Sb(v) strengthens the interaction between the Lewis acid function and the Lewis basic chalcogen atom, resulting in the formation of four-membered heterocycles and, in some cases, strong deformation of the Sb-C-P backbone. Selective oxidation of the antimony atom with o-chloranil is not possible due to the electron deficiency caused by the C₂F₅ groups; it is rather the phosphane function of 1 which is oxidised first. Only in a second oxidation step is the Sb(m) function oxidised.

Author contributions

J. Krieft: investigation, methodology, validation, visualisation, writing (original draft), B. Neumann: investigation (SCXRD), H.-G. Stammler: investigation (SCXRD), N. W. Mitzel: funding acquisition, project administration, supervision, reviewing and editing.

Data availability

The data published in this contribution are available as ESI,† submitted with the manuscript. Crystallographic data have been deposited with the Cambridge Crystal Structure Database (CCDC).

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

The authors thank Jan-Niklas Bollnow for lab assistance, Marco Wißbrock for recording NMR spectra and Barbara Teichner for performing elemental analyses. This work was founded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation, grant MI 477/44-1, project number 461833739).

References

- (a) M. Baaz, V. Gutmann and O. Kunze, *Monatsh. Chem.*, 1962, **93**, 1142–1161; (b) G. A. Olah and R. H. Schlosberg, *J. Am. Chem. Soc.*, 1968, **90**, 2726–2727; (c) G. A. Olah, *J. Org. Chem.*, 2005, **70**, 2413–2429; (d) J. Bacon and R. J. Gillespie, *J. Am. Chem. Soc.*, 1971, **93**, 6914–6919; (e) J. Y. Calves and R. J. Gillespie, *J. Am. Chem. Soc.*, 1977, **99**, 1788–1792.
- 2 (a) D. W. Stephan and G. Erker, Angew. Chem., Int. Ed., 2010, 49, 46-76; (b) C. M. Mömming, E. Otten, G. Kehr, R. Fröhlich, S. Grimme, D. W. Stephan and G. Erker, Angew. Chem., Int. Ed., 2009, 48, 6643-6646; (c) D. W. Stephan, Dalton Trans., 2009, 17, 3129-3136; (d) A. E. Ashley, A. L. Thompson and D. O'Hare, Angew. Chem., Int. Ed., 2009, 48, 9839-9843; (e) O. J. Metters, S. J. K. Forrest, H. A. Sparkes, I. Manners and D. F. Wass, J. Am. Chem. Soc., 2016, 138, 1994-2003.
- 3 C. Manankandayalage, D. K. Unruh and C. Krempner, *Chem. Eur. J.*, 2021, **27**, 6263–6273.
- 4 D. W. Stephan and G. Erker, Angew. Chem., Int. Ed., 2015, 54, 6400-6441.
- 5 (a) G. C. Welch, R. R. San Juan, J. D. Masuda and Stephan, Science, 2006, 314, 1124–1126; D. W. (b) G. C. Welch and D. W. Stephan, J. Am. Chem. Soc., 2007, 129, 1880–1881; (c) C. Appelt, H. Westenberg, F. Bertini, A. W. Ehlers, J. C. Slootweg, K. Lammertsma and W. Uhl, Angew. Chem., Int. Ed., 2011, 50, 3925-3928; (d) C. Appelt, J. C. Slootweg, K. Lammertsma and W. Uhl, Angew. Chem., Int. Ed., 2013, 52, 4256-4259; (e) D. W. Stephan, J. Am. Chem. Soc., 2015, 137, 10018-10032; (f) D. W. Stephan, Science, 2016, 354, aaf7229; (g) W. Uhl, J. Possart, A. Hepp and M. Layh, Z. Anorg. Allg. Chem., 2017, 643, 1016-1029; (h) L. Wickemeyer, N. Aders, A. Mix, B. Neumann, H.-G. Stammler, J. J. Cabrera-Trujillo, I. Fernández and N. W. Mitzel, Chem. Sci., 2022, 13, 8088-8094.
- 6 B. Waerder, M. Pieper, L. A. Körte, T. A. Kinder, A. Mix,
 B. Neumann, H.-G. Stammler and N. W. Mitzel, *Angew. Chem., Int. Ed.*, 2015, 54, 13416–13419.
- 7 P. Holtkamp, F. Friedrich, E. Stratmann, A. Mix,
 B. Neumann, H.-G. Stammler and N. W. Mitzel, *Angew. Chem., Int. Ed.*, 2019, 58, 5114–5118.
- 8 T. A. Kinder, R. Pior, S. Blomeyer, B. Neumann, H.-G. Stammler and N. W. Mitzel, *Chem. – Eur. J.*, 2019, **25**, 5899–5903.
- 9 (a) H. Li, A. J. A. Aquino, D. B. Cordes, F. Hung-Low,
 W. L. Hase and C. Krempner, *J. Am. Chem. Soc.*, 2013, 135, 16066–16069; (b) S. Mummadi, D. K. Unruh, J. Zhao, S. Li and C. Krempner, *J. Am. Chem. Soc.*, 2016, 138, 3286–3289; (c) S. Mummadi, A. Brar, G. Wang, D. Kenefake, R. Diaz,

D. K. Unruh, S. Li and C. Krempner, *Chem. – Eur. J.*, 2018, **24**, 16526–16531.

- 10 P. Erdmann, J. Leitner, J. Schwarz and L. Greb, *ChemPhysChem*, 2020, **21**, 987–994.
- (a) M. Schorpp, R. Yadav, D. Roth and L. Greb, Angew. Chem., Int. Ed., 2022, 61, e202207963; (b) M. Á. García-Monforte, M. Baya, D. Joven-Sancho, I. Ara, A. Martín and B. Menjón, J. Organomet. Chem., 2019, 897, 185–191; (c) D. Sharma, S. Balasubramaniam, S. Kumar, E. D. Jemmis and A. Venugopal, Chem. Commun., 2021, 57, 8889–8892; (d) B. L. Murphy and F. P. Gabbaï, J. Am. Chem. Soc., 2023, 145, 19458–19477.
- 12 B. Pan and F. P. Gabbaï, J. Am. Chem. Soc., 2014, **136**, 9564–9567.
- 13 D. Tofan and F. P. Gabbaï, Chem. Sci., 2016, 7, 6768-6778.
- 14 S. Benz, A. I. Poblador-Bahamonde, N. Low-Ders and S. Matile, *Angew. Chem., Int. Ed.*, 2018, 57, 5408–5412.
- (a) L. M. Lee, M. Tsemperouli, A. I. Poblador-Bahamonde, S. Benz, N. Sakai, K. Sugihara and S. Matile, *J. Am. Chem. Soc.*, 2019, **141**, 810–814; (b) J. Qiu, B. Song, X. Li and A. F. Cozzolino, *Phys. Chem. Chem. Phys.*, 2017, **20**, 46–50; (c) Di You, B. Zhou, M. Hirai and F. P. Gabbaï, *Org. Biomol. Chem.*, 2021, **19**, 4949–4957.
- 16 (a) J. L. Beckmann, J. Krieft, Y. V. Vishnevskiy, B. Neumann, H.-G. Stammler and N. W. Mitzel, Angew. Chem., Int. Ed., 2023, 62, e202310439; (b) J. L. Beckmann, J. Krieft, Y. V. Vishnevskiy, B. Neumann, H.-G. Stammler and N. W. Mitzel, Chem. Sci., 2023, 14, 13551–13559.
- 17 (a) M. Yang, D. Tofan, C.-H. Chen, K. M. Jack and F. P. Gabbaï, *Angew. Chem., Int. Ed.*, 2018, 57, 13868–13872;
 (b) A. M. Christianson and F. P. Gabbaï, *Organometallics*, 2017, 36, 3013–3015.
- 18 J. E. Smith, H. Yang and F. P. Gabbaï, *Organometallics*, 2021, **40**, 3886–3892.
- 19 V. M. Gonzalez, G. Park, M. Yang and F. P. Gabbaï, *Dalton Trans.*, 2021, **50**, 17897–17900.
- 20 C.-H. Chen and F. P. Gabbaï, *Dalton Trans.*, 2018, 47, 12075–12078.
- 21 (a) B. Eberwein and J. Weidlein, Z. Anorg. Allg. Chem., 1976, 420, 229–239; (b) M. J. Begley, D. B. Sowerby, D. M. Wesolek, C. Silvestru and I. Haiduc, J. Organomet. Chem., 1986, 316, 281–289; (c) J. M. Mercero, X. Lopez, J. E. Fowler and J. Ugalde, J. Phys. Chem. A, 1997, 101, 5574–5579; (d) I. Ghesner, L. Opris, G. Balazs, H. J. Breunig, J. E. Drake, A. Silvestru and C. Silvestru,

J. Organomet. Chem., 2002, 642, 113-119; (e) O. Kysliak and J. Beck, Z. Anorg. Allg. Chem., 2013, 639, 2860-2867.

- 22 P. Holtkamp, T. Glodde, D. Poier, B. Neumann, H.-G. Stammler and N. W. Mitzel, *Angew. Chem., Int. Ed.*, 2020, **59**, 17388–17392.
- 23 (a) D. Naumann, G. Nowicki and K.-J. Sassen, Z. Anorg. Allg. Chem., 1997, 623, 1183–1189; (b) M. B. Murphy-Jolly, L. C. Lewis and A. J. M. Caffyn, Chem. Commun., 2005, 4479–4480; (c) S. Solyntjes, J. Bader, B. Neumann, H.-G. Stammler, N. Ignat'ev and B. Hoge, Chem. – Eur. J., 2017, 23, 1557–1567.
- 24 (a) P. Holtkamp, J. Schwabedissen, B. Neumann, H.-G. Stammler, I. V. Koptyug, V. V. Zhivonitko and N. W. Mitzel, *Chem. Eur. J.*, 2020, 26, 17381–17385;
 (b) P. Holtkamp, D. Poier, B. Neumann, H.-G. Stammler and N. W. Mitzel, *Chem. Eur. J.*, 2021, 27, 3793–3798.
- 25 J. Krieft, P. C. Trapp, Y. V. Vishnevskiy, B. Neumann, H.-G. Stammler, J.-H. Lamm and N. W. Mitzel, *Chem. Sci.*, 2024, in revision.
- 26 R. G. Pearson, J. Am. Chem. Soc., 1963, 85, 3533-3539.
- 27 A. Bondi, J. Phys. Chem., 1964, 68, 441-451.
- 28 M. Mantina, A. C. Chamberlin, R. Valero, C. J. Cramer and D. G. Truhlar, *J. Phys. Chem. A*, 2009, **113**, 5806–5812.
- 29 P. Pyykkö and M. Atsumi, Chem. Eur. J., 2009, 15, 186-197.
- 30 M. Engelhardt, C. L. Raston, C. R. Whitaker and A. H. White, *Aust. J. Chem.*, 1986, **39**, 2151–2154.
- 31 J. Ramler and C. Lichtenberg, *Chem. Eur. J.*, 2020, 26, 10250–10258.
- 32 S. H. Ahn, K. J. Cluff, N. Bhuvanesh and J. Blümel, *Angew. Chem., Int. Ed.*, 2015, **54**, 13341–13345.
- 33 H. U. Steinberger, B. Ziemer and M. Meisel, Acta Crystallogr., Sect. C: Cryst. Struct. Commun., 2001, 57, 835– 837.
- 34 H. U. Steinberger, B. Ziemer and M. Meisel, Acta Crystallogr., Sect. C: Cryst. Struct. Commun., 2001, 57, 323– 324.
- 35 N. Kuhn, H. Schumann and G. Wolmershäuser, Z. Naturforsch., B: J. Chem. Sci., 1987, 42, 674–678.
- 36 (a) M. M. Sidky, M. R. Mahran and W. M. Abdou, *Phosphorus Sulfur Relat. Elem.*, 1983, 15, 129–135;
 (b) R. R. Holmes, R. O. Day, V. Chandrasekhar and J. M. Holmes, *Inorg. Chem.*, 1987, 26, 157–163.I.-S. Ke and F. P. Gabbaï, *Inorg. Chem.*, 2013, 52, 7145–7151.