Multiply bridged organodiantimony(v) compounds; crystal structures of $[(SbR_2)_2(\mu-O)_2(\mu-O_2AsMe_2)_2]$ (R = Ph or *p*-tolyl) and $[(SbR_3)_2(\mu-O)(\mu-O_2MR'_2)_2]$ (R = Ph, M = P, R' = Me; R = Ph, M = As, R' = Me or Ph; R = *p*-tolyl, M = P, R' = Me)

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A number of quadruply and triply bridged organodiantimony(v) compounds have been synthesized by reactions of $[(SbR_2BrO)_2]$ (R = Ph or *p*-tolyl) and $[(SbR_3X)_2O]$ (R = Ph or *p*-tolyl, X = Br or Cl) with Na(O₂AsMe₂), Na(O₂AsPh₂) or Na(O₂AsMe₂). The compounds have been characterised by a range of spectroscopic methods and crystal structures are reported for two quadruply bridged, $[(SbPh_2)_2(\mu-O)_2(\mu-O_2AsMe_2)_2]$ and $[\{Sb(p-MeC_6H_4)_2\}_2(\mu-O)_2(\mu-O_2AsMe_2)_2]$, and four triply bridged compounds, $[(SbPh_3)_2(\mu-O)(\mu-O_2PMe_2)_2]$, $[(SbPh_3)_2(\mu-O)(\mu-O_2AsMe_2)_2]$, $[(SbPh_3)_$

Until recently there were very few diantimony compounds known in which the atoms are spanned by either three or four bridging groups and when this work was started only two compounds containing quadruple bridges had been characterised. The first was the zwitterionic $[(SbPh_2)_2(\mu-O)_2(\mu-O_2SbPh_2)_2]$,¹ obtained as a by-product in the oxidation of triphenylantimony while the second was an anionic species, $[(SbPh_2)_2(\mu-O)_2(\mu-O_2P_2)_2]^{-1}$, containing two oxo and two molybdate bridges.^{2,3} More recently two further species, $[(SbPh_2)_2(\mu-O)_2(\mu-O_2PR_2)_2]$ where R = cyclohexyl or cyclooctyl, have been synthesized.⁴

There is a more extensive series of triply bridged compounds, synthesized by Schmidt and others, which contain in addition to dioxo chelating ligands both oxo and either hydroxo or alkoxo bridges. The general formula is $[(SbCl_3)_2(\mu-O)(\mu-OR)-(\mu-X)]$ with, for example, R = H, $X = O_2CMe$,⁵ O_2CCl_3 ,⁶ HSO_4^7 or ClO_4 ;⁸ R = Me, $X = SO_4Me$,⁹ SO_2Me ,¹⁰ SO_3Me^{10} or $O_2PMe(OMe)$;¹¹ and R = Et, $X = O_2PEt(OEt)^{11}$ or $O_2PMe(OMe)$;¹¹ Bridging in the tetranuclear compound, $[Sb_4Ph_8-O_6(MeCO_2H)_3]$,¹² is similar with pairs of antimony atoms spanned by hydroxo, oxo and acetate bridges. Pairs of antimony atoms can also be bridged by either three halogens or two halogens and one oxygen, as in $[NMe_4]_3[(SbCl_3)_2(\mu-Br)_3]^{13}$ and $[Hpy]_2[(SbCl_2)_2(\mu-O)(\mu-X)_2]$, where py = pyridine and X = Br or Cl,¹⁴ respectively.

One triply bridged compound, $[(SbCl_3)_2(\mu-O)(\mu-O_2PMe_2)_2]$, with an oxo and two dimethylphosphinate bridges has previously been synthesized ¹⁵ and, while this work was in progress, two further compounds of this type $[(SbPh_2Cl)_2(\mu-O)(\mu-O_2PR_2)_2]$,¹⁶ where R = cyclo-hexyl or -octyl, were isolated. Crystal structures for both $[(SbCl_3)_2(\mu-O)(\mu-O_2PMe_2)_2]$ and $[(SbPh_2Cl)_2(\mu-O){\mu-O_2P(C_6H_{11})_2}_2]$ show octahedral coordination at antimony with bidentate phosphinate groups; the geometry is similar in the related carboxylate-bridged compound $[(SbF_3)_2(\mu-O)(\mu-O_2CCF_3)_2]$.¹⁷

Phosphinates and arsinates can, however, behave as either unidentate or chelating ligands and the work described in this paper was carried out to define further the conditions under which the groups might bridge between pairs of antimony atoms already linked by either one or two oxygen atoms. The starting materials were the singly bridged dihalides $[(SbR_3X)_2O]$, where X = Br or Cl and R = phenyl, p-tolyl or methyl, and the double bridged analogues $[(SbR_2BrO)_2]$ where R = phenyl or *p*-tolyl. Related dithiophosphinates are also potentially good bridging ligands but complex redox reactions occur with these antimony(v) compounds and reduction products such as $[SbPh_2(S_2PR_2)]$ can be isolated.¹⁸

Discussion

Quadruply bridged compounds

Preparation. Quadruply bridged diantimony compounds $[(SbPh_2)_2(\mu-O)_2(\mu-O_2AsR_2)_2]$, where $R = Me \ 1$ or Ph 2, have been prepared by treating [(SbPh₂BrO)₂], which contains a fourmembered Sb₂O₂ ring, with 2 mol of either Na(O₂AsMe₂) or Na(O₂AsPh₂) in dichloromethane. They are high-melting stable compounds, which crystallise as chloroform solvates. An attempt to introduce a single arsinate bridge by treating [(SbPh₂BrO)₂] with 1 mol of Na(O₂AsMe₂) gave a complex reaction mixture and only a small amount of unidentified crystalline material was isolated. Similar reactions with related phosphinates, Na(O₂PMe₂) and Na(O₂PPh₂), however, failed to give the expected quadruply bridged analogues, and with dimethylphosphinate a triply bridged compound, [(SbPh₃)₂(µ- $O(\mu - O_2 PMe_2)_2$] 3, was obtained. This can only arise by an extensive rearrangement, involving loss of an oxo bridge and gain of one phenyl group per antimony, and currently no realistic mechanism can be advanced.

Unexpectedly, a third quadruply bridged compound, [{Sb(p-MeC₆H₄)₂}₂(μ -O)₂(μ -O₂AsMe₂)₂] **4**, was obtained in low yield by treating [{Sb(p-MeC₆H₄)₃Br}₂O] with 2 mol of Na(O₂AsMe₂) in dichloromethane. In this case, rearrangement with gain of an oxo bridge and loss of a p-tolyl group from each antimony may result from hydrolysis and elimination of toluene as shown in equation (1).

$$[\{\mathrm{Sb}(p\operatorname{-MeC}_{6}\mathrm{H}_{4})_{3}\mathrm{Br}\}_{2}\mathrm{O}] + 2\mathrm{Na}(\mathrm{O}_{2}\mathrm{AsMe}_{2}) + \mathrm{H}_{2}\mathrm{O} \longrightarrow$$
$$[\{\mathrm{Sb}(p\operatorname{-MeC}_{6}\mathrm{H}_{4})_{2}\}_{2}(\mu\operatorname{-O})_{2}(\mu\operatorname{-O}_{2}\mathrm{AsMe}_{2})_{2}] + 2\mathrm{NaBr} + 2\mathrm{MeC}_{e}\mathrm{H}_{e} \quad (1)$$

Spectroscopy. All three quadruply bridged compounds **1**, **2** and **4** give complex infrared spectra with many bands associated with aryl groups,¹⁹ but an intense broad band centred at *ca.* 800 cm⁻¹ can be assigned to AsO_2 stretching. The fact that the antisymmetric and symmetric modes are not resolved is taken to point to symmetrical bonding. Rather surprisingly, it is not



Fig. 1 Molecular structure and atom numbering scheme for $[(SbPh_2)_2 \cdot (\mu-O)_2(\mu-O_2AsMe_2)_2] \cdot 2CHCl_3$ 1

Table 1 Selected bond distances (Å) and angles (°) for $[(SbPh_2)_2(\mu-O)_2(\mu-O_2AsMe_2)_2]\cdot 2CHCl_3$ 1, with estimated standard deviations(e.s.d.s) in parentheses *

Sb(1)-O(1)	1.995(3)	Sb(1)–C(7)	2.156(4)
Sb(1)–O(1')	2.004(3)	As(1)–O(2)	1.709(3)
Sb(1)-O(2)	2.101(3)	As(1)–O(3')	1.711(3)
Sb(1)-O(3)	2.083(3)	As(1)–C(13)	1.909(4)
Sb(1)-C(1)	2.157(4)	As(1)-C(14)	1.911(4)
O(1)-Sb(1)-O(1')	79.5(1)	O(3)-Sb(1)-C(7)	89.6(1)
O(1)-Sb(1)-O(2)	89.9(1)	C(1)-Sb(1)-C(7)	97.4(1)
O(1')-Sb(1)-O(2)	85.5(1)	O(2)-Sb(1)-O(3)	174.3(1)
O(1)-Sb(1)-O(3)	85.3(1)	O(1')-Sb(1)-C(1)	170.0(1)
O(1')-Sb(1)-O(3)	90.7(1)	O(1)-Sb(1)-C(7)	169.3(1)
O(1)-Sb(1)-C(1)	92.5(1)	O(2)-As(1)-O(3')	114.0(1)
O(2)-Sb(1)-C(1)	88.6(1)	C(13)-As(1)-C(14)	110.0(2)
O(3)-Sb(1)-C(1)	94.6(1)	Sb(1)–O(1)–Sb(1')	100.5(1)
O(1')-Sb(1)-C(7)	91.1(1)	Sb(1)–O(2)–As(1)	119.0(1)
O(2)–Sb(1)–C(7)	94.7(1)	Sb(1)–O(3)–As(1')	119.5(1)
* Atoms carrying a $1 - v 2 - z$	prime are rela	ted by the symmetry operat	ion 1 – <i>x</i> ,
1 J, w Z.			

possible to assign any bands to the Sb_2O_2 ring modes, which occur for the starting material between 495 and 670 cm^{-1.20}

There were no unexpected features in the ¹H NMR spectra for the three compounds but $[(SbPh_2)_2(\mu-O)_2(\mu-O_2AsPh_2)_2]$ **2** showed a broad unresolved multiplet at δ 7.30–7.80. All three compounds show parent ions in their FAB mass spectra, although for **2** the highest mass ion is attributed to $[M + 2H]^+$. Subsequent fragmentation, as expected, involves loss of both aryl and arsinate groups and there are a number of prominent monoantimony species, including $[SbPh_2(O_2AsMe_2)]^+$ for **1**, $[SbPh_2(O_2AsPh_2) + H]^+$ for **2** and the rearrangement ion $[Sb-(p-MeC_6H_4)_3(O_2AsMe_2)]^+$ is the base peak for **4**.

Structures. Single-crystal structures have been determined for $[(SbPh_2)_2(\mu-O)_2(\mu-O_2AsMe_2)_2]$ **1**, as a bis(chloroform) solvate, and $[\{Sb(p-MeC_6H_4)_2\}_2(\mu-O)_2(\mu-O_2AsMe_2)_2]$ **4**. The molecular structure of **1**, which has crystallographic C_i symmetry, is shown in Fig. 1 and selected bond distances and angles are listed in Table 1. The determination confirms the presence of a quadruply bridged structure with octahedrally co-ordinated antimony atoms. The Sb–O distances in the four-membered ring [1.995(3) and 2.004(3) Å] are effectively equal, in contrast to those in the starting material, which are either 'short' (1.936, 1.918 Å) or 'long' (2.061, 2.030 Å) depending on their position in the trigonal-bipyramidal co-ordination about antimony.²⁰

Co-ordination of the dimethylarsinate groups, which consti-



Fig. 2 Molecular structure and atom numbering scheme for $[\{Sb(p-MeC_6H_4)_2\}_2(\mu\text{-}O)_2(\mu\text{-}O_2AsMe_2)_2]$ 4

Table 2 Selected bond distances (Å) and angles (°) for [{Sb(p-

$O_2AsMe_2)_2$] 4 , with e.s.d.s in parentheses	*
2.007(3)	Sb(1)-C(8)	2.161(4)
2.007(3)	As(1)–O(1A)	1.701(3)
2.087(3)	As(1)–O(2)	1.703(3)
2.084(3)	As(1)–C(15)	1.904(4)
2.148(4)	As(1)-C(16)	1.903(4)
119.5(1)	O(1)-Sb(1)-C(1)	93.2(1)
79.0(1)	C(1)-Sb(1)-C(8)	99.0(2)
85.4(1)	O(1)–Sb(1)–O(2)	173.0(1)
89.4(1)	O(3A)-Sb(1)-C(8)	167.2(1)
89.9(1)	O(3)–Sb(1)–C(1)	171.0(1)
84.6(1)	O(1A)-As(1)-O(2)	113.4(1)
89.3(1)	C(15)-As(1)-C(16)	110.4(2)
95.1(1)	Sb(1)-O(3)-Sb(1A)	101.0(1)
90.1(2)	Sb(1)–O(2)–As(1)	119.0(1)
92.9(1)	Sb(1)–O(1)–As(1A)	119.5(1)
90.6(1)		
	O ₂ AsMe ₂) ₂ 2.007(3) 2.087(3) 2.087(3) 2.084(3) 2.148(4) 119.5(1) 79.0(1) 85.4(1) 89.9(1) 84.6(1) 89.3(1) 95.1(1) 90.1(2) 92.9(1) 90.6(1)	$\begin{array}{llllllllllllllllllllllllllllllllllll$

*Atoms designated A are related by the symmetry operation -x, 1 - y, -z.

Table 3 Comparison of Sb₂O₂ mean ring parameters

Compound	Sb-O/Å	O–Sb–O/°	Sb-O-Sb/°	Ref.
1	2.000	79.5	100.5	This work
4	2.007	79.0	101.0	This work
$[{SbPh_2(MoO_4)O}_2]^{2-}$	2.019	77.1	103.0	2
$[{SbPh_2O(O_2SbPh_2)}_2]$	2.089	72.7	107.3	1
[(SbPh ₃ O) ₂]	2.002	77.5	102.5	1
[(SbPh ₂ BrO) ₂]	1.986	78.6	101.5	21

tute the other two bridging groups, is highly symmetrical with identical As–O bond lengths, *i.e.* As(1)–O(2) 1.709(3) Å and As(1)–O(3') 1.711(3) Å, implying complete delocalisation of the π component of the As–O system. The arsenic atom is basically tetrahedral although the O(2)–As(1)–O(3') angle is opened to 114.0(1)°. This together with angles of *ca.* 119.5° at O(2) and O(3) probably represents the extent of the flexibility in this part of the system towards matching ligand bite with Sb · · · Sb separation.

Co-ordination about antimony is distorted octahedral with the O(1)–Sb(1)–O(1') angle associated with the four-membered ring constrained to 79.5(1)°. An important point is that the *trans* angle involving the bridging groups [O(2)–Sb(1)–O(3)174.3(1)°] is close to 180° implying that incorporation of the bridging arsinate groups imposes no great strain. This structure is clearly very similar to that of $[(SbPh_2)_2(\mu-O)_2(\mu-MOQ_4)_2]^{2-,2}$ where the significantly longer Mo–O separations (mean 1.83 Å; *cf.* As–O, mean 1.71 Å) imply a larger bite and even less strain.

Compound **4**, $[{Sb(p-MeC_6H_4)_2}_2(\mu-O)_2(\mu-O_2AsMe_2)_2]$ (see Fig. 2 and Table 2) has a closely related structure with again

crystallographically imposed C_i symmetry, symmetrically coordinated dimethylarsinate groups [As(1)-O1(A) and As(1)-O(2) 1.701(3) and 1.703(3) Å, respectively] and similar geometries about antimony and arsenic. An interesting feature in both compounds is the relative orientations of the aryl groups. The C(1)–C(6) group is almost coplanar with the Sb_2O_2 plane $(4.0^{\circ}$ between best-fit planes) while the C(8) to C(14) group is almost perpendicular (94.5°).

Table 3 summarises data for compounds containing Sb₂O₂ rings, showing there is little variation in either Sb-O distances (ca. 2.00 Å) or angles (ca. 78°) in **1** and **4** compared with those in related compounds. The only noticeable difference is with the zwitterionic $[{SbPh_2O(O_2SbPh_2)}_2]$ where bridging oxo groups are involved in additional bonding to the antimony centres of the bridging diphenylstibinate moieties. With the exception of [(SbPh₂BrO)₂], the rings are centrosymmetric and the compounds have near D_{2h} local symmetry.

Triply bridged compounds

Preparations. Triply bridged compounds, [(SbPh₃)₂(µ-O)(µ- $O_2MR_2)_2$] (M = P, R = Me **3**; M = As, R = Me **5** or Ph **6**), were prepared in good yield as stable crystalline solids by reactions of [(SbPh₃Br)₂O], with 2 mol of the appropriate Na(O₂MR₂) but only unidentified products were obtained in the reaction with Na(O_2PPh_2). Reaction between [{Sb(p-MeC_6H_4)_3Br}_2O] and 2 mol of Na(O₂PMe₂) gave the expected bridged compound, $[{Sb(p-MeC_6H_4)_3}_2(\mu-O)(\mu-O_2PMe_2)_2]$ 7, in good yield, but surprisingly the o-tolyl analogue could not be obtained from [{Sb(o-MeC₆H₄)₃Br}₂O] and 2 mol of Na(O₂PMe₂). An attempt to form an unsymmetrically substituted compound, [(SbPh₃)₂(µ-O)(µ-O₂AsMe₂)Br], from (SbPh₃Br)₂O and 1 mol of Na(O₂AsMe₂) gave only mixtures of 5 and unchanged (SbPh₃Br)₂O.

Related trimethylantimony(v) compounds, [(SbMe₃)₂(µ- $O((\mu - O_2 PMe_2)_2)$ 8 and $[(SbMe_3)_2(\mu - O)(\mu - O_2 AsPh_2)_2]$ 9, were prepared in good yield from [(SbMe₃Cl)₂O] and 2 mol of either Na(O₂PMe₂) or Na(O₂AsPh₂), but they are highly moisture sensitive and handling, even with stringent Schlenk and glovebox techniques, was extremely difficult. Conventional microanalysis and IR spectroscopy were not possible but satisfactory ¹H and ¹³C-{¹H} NMR spectra were obtained and these compounds are considered to have the same triply bridged structures as those described above.

It was noted above that reaction of $[{Sb(p-MeC_6H_4)_3Br}_2O]$ with 2 mol of $Na(O_2AsMe_2)$ gave, instead of the expected triply bridged product, a quadruply bridged rearrangement compound 4, formed presumably because of its greater stability. This is all the more surprising as the related phenylated starting material, [(SbPh₃Br)₂O], reacts with Na(O₂AsMe₂) to give $[(SbPh_3)_2(\mu-O)(\mu-O_2AsMe_2)_2]$ 5 in good yield. Detailed examination of the latter reaction by ¹H NMR spectroscopy, however, does show evidence for formation of 1, the analogous rearrangement product, but here rearrangement is of minor importance.

Spectroscopy. The IR spectrum of [(SbPh₃)₂(µ-O)(µ- $O_2PMe_2)_2$] 3 contains a strong, broad band at 1112 cm⁻¹ assigned to the $v_{asym}(\text{PO}_2)$ mode, and a doublet at 1046 and 1032 cm⁻¹, probably the in- and out-of-phase components of $v_{sym}(PO_2)$. For related tin dimethylphosphinates,²¹ bands between 1200 and 1000 cm⁻¹ are associated with either chelating or bridging ligands, supporting triply bridged structures here. Bands for $[{Sb(p-MeC_6H_4)_3}_2(\mu-O)(\mu-O_2PMe_2)_2]$ 7 are at similar positions, with $\nu_{\text{asym}}(\text{PO}_2)$ assigned to peaks at 1120 and 1111 cm⁻¹ and $v_{sym}(PO_2)$ at 1048 and 1033 cm⁻¹.

Spectra of the arsinates, [(SbPh₃)₂(µ-O)(µ-O₂AsMe₂)₂] 5 and $[(SbPh_3)_2(\mu-O)(\mu-O_2AsPh_2)_2]$ 6, contain intense broad bands at 852 and 831 cm⁻¹, respectively, consistent with the presence of bridging (O₂AsR₂) groups; these are at significantly higher

energy than corresponding bands (ca. 800 cm⁻¹) for related quadruply bridged compounds. The IR spectra for [(SbMe₃)₂(µ- $O((\mu - O_2 PMe_2)_2]$ 8 and $[(SbMe_3)_2(\mu - O)(\mu - O_2 AsPh_2)_2]$ 9 may be unreliable, as hydrolysis appears to take place, but for 8 the broad bands at 1172 and 1020 cm⁻¹ may be assigned tentatively to P-O stretches of bidentate O2PMe2 groups, while the broad 795 cm^{-1} band for **9** can probably be assigned to the related As-O stretches.

Proton NMR spectra for the arylantimony compounds point to complex solution behaviour and it is clear that, unless stringent precautions to exclude moisture are taken, there is either complete or partial hydrolysis to hydroxo-species, such as SbR₃(O₂MR'₂)(OH). Using CDCl₃ as received and with no precautions against moisture during solution preparation, i.e. 'moist' CDCl₃, the spectrum of [(SbPh₃)₂(µ-O)(µ-O₂PMe₂)₂] **3** can be assigned to a single species with the expected 3:1 ratio of phenyl and dimethylphosphinate groups. Following related studies on $[(SbPh_3X)_2O]$ solutions, where X = Br or $Cl_{,22}$ this is assigned to the hydroxo-species [SbPh₃(O₂PMe₂)(OH)] formed as a result of hydrolysis by adventitious water. As with the halide systems, recrystallisation of the NMR sample gave quantitative recovery of 3 and no evidence for the hydroxo-species in the solid.

Under strictly anhydrous conditions, *i.e.* with dried CDCl₃, a sealed NMR tube and manipulations carried out in a nitrogenfilled glove-box, there were signals for two species, neither of which was the hydrolysis product, and the oxo bridge is presumed to be intact in each. The O2PMe2 group of the first species **A** showed a doublet at $\delta 0.72 [^2 J(HP) = 14 Hz]$ and two triplets and a doublet $[{}^{3}J(HH) = 7 Hz]$ in the phenyl region. For the second species **B** the O₂PMe₂ signal collapsed to a broad featureless signal with two second-order multiplets, centred at δ 7.38 and 7.68, in the phenyl region. The **A** : **B** ratio was ca. 2:3. It is difficult to assign structures to these species with any certainty but it is likely that one has the solid-state structure with bridging dimethylphosphinate groups (see later), while in the second the dimethylphosphinate groups could, perhaps, be chelating or even unidentate.

The *p*-tolyl derivative **7** behaved similarly with a single hydrolysed species in 'moist' CDCl₃ and two species with chemical shifts closely related to those discussed above in anhydrous solutions. Species A is dominant with an A: B ratio of ca. 5:1. Exposure of the anhydrous sample to moisture led to a spectrum showing signals for all three species. Two species were also present for [(SbPh₃)₂(µ-O)(µ-O₂AsMe₂)₂] 5 in dry CDCl₃ and, although both dimethylarsinate signals (δ 0.71 and 1.89) were broad and featureless, the phenyl region clearly showed two distinct sets of signals. The first was a triplet, triplet, doublet pattern at 87.17, 7.27 (masked by a multiplet from a second species) and 7.61 for the meta, para and ortho protons, respectively, while the second showed multiplets at δ 7.27 and 7.71. The ratio of the two species was ca. 2:1.

In contrast, solution spectra for $[(SbMe_3)_2(\mu-O)(\mu-O_2PMe_3)_2]$ 8 and $[(SbMe_3)_2(\mu-O)(\mu-O_2AsPh_2)_2]$ 9 were more straightforward and only one species was observed in each case. The SbMe₃ signals were at higher field (δ 1.66) compared with those for related compounds, e.g. & 1.96 for [(SbMe₃Cl)₂O] and 2.07 for [{SbMe₃[S₂P(OMe)₂]}₂O].²³ The ¹³C shifts for the methyl groups at antimony were similar, δ 16.3 and 16.8 respectively, again to higher field than those for $[{SbMe_3[S_2P(OMe)_2]}_2O](\delta)$ 18.6).

The FAB mass spectra of the triply bridged triarylantimony compounds, in contrast to those of the quadruply bridged compounds, do not show the parent ion. The spectra of $[(SbPh_3)_2(\mu-O)(\mu-O_2PMe_2)_2]$ 3, $[(SbPh_3)_2(\mu-O)(\mu-O_2AsMe_2)_2]$ 5 and $[{Sb(p-MeC_6H_4)_3}_2(\mu-O)(\mu-O_2PMe_2)_2]$ 7 all show peaks resulting from loss of one ligand and are assigned to the appropriate $[(SbR_3)_2(\mu-O)(\mu-O_2MMe_2)]^+$ (R = aryl) ions. Major species in all the spectra are monometallic [SbR₃(O₂MMe₂)]⁺ and simple arylantimony ions.



Fig. 3 Molecular structure and atom numbering scheme for $[(SbPh_3)_2(\mu\text{-O})(\mu\text{-O}_2PMe_2)_2]\cdot 0.5C_6H_{14}$ 3, structures for compounds 5 and 7 are similar

Structures. Single-crystal structure determinations were carried out for four of these triply bridged compounds, three of which (**3**, **4** and **7**) were hemihexane solvates, while the fourth (**6**) was a chloroform solvate. The overall structures of all four compounds, $[(SbPh_3)_2(\mu-O)(\mu-O_2PMe_2)_2] \cdot 0.5C_6H_{14}$ **3**, $[(SbPh_3)_2(\mu-O)(\mu-O_2AsMe_2)_2] \cdot 0.5C_6H_{14}$ **5**, $[(SbPh_3)_2(\mu-O)(\mu-O_2AsPh_2)_2] \cdot CHCl_3$ **6** and $[\{Sb(p-MeC_6H_4)_3\}_2(\mu-O)(\mu-O_2PMe_2)_2] \cdot 0.5C_6H_{14}$ **7**, are very similar; **3** and **5** are isostructural. A diagram showing the structure of **3**, with the skeletal numbering scheme, which is appropriate for all four compounds, is shown in Fig. 3. Selected bond distances and angles for compounds **3**, **4** and **7** are listed in Table 4 while those for **6** with diphenylarsinate bridges are in Table 5.

The structure determinations confirm that all four compounds are triply bridged diantimony species with one oxygen and two phosphinate or arsinate bridges and the structures are comparable with those of $[(SbCl_3)_2(\mu-O)(\mu-O_2PMe_2)_2]^{15}$ and $[(SbPh_2Cl)_2(\mu-O)\{\mu-O_2P(C_6H_{11})_2\}_2]^{.16}$ Antimony atoms are in distorted octahedral co-ordination to three oxygen and three carbon atoms in the fac isomeric form. The Sb-O-Sb bridge bonds are equal in **3** and **5** while in **6** and **7** the differences (*ca.* 0.02 Å) are on the limit of significance. As shown in Table 6, mean bridging distances in the present compounds are longer than those in the three compounds previously investigated, which have either chlorine or fluorine substituents on antimony, and the expected increased Lewis acidity would probably account for the bond shortening. The Sb-O-Sb angles in three of the compounds reported here are comparable with that in $[(SbPh_2Cl)_2(\mu\text{-}O)\{\mu\text{-}O_2P(C_6H_{11})_2\}_2]^{16}$ but again there are consequences if antimony carries three electronegative substituents when the angle closes substantially. The bridge angle in 6, on the other hand, is increased to 151.7° and is probably a consequence of the change from methyl substituents at arsenic in 5 to phenyl groups in 6. We have previously commented ¹⁸ on the fact that, in a series of compounds, substantial structural changes sometimes result from substituent changes at a 'remote' atom and this seems to be another such example.

There is extensive delocalisation of the π component of the co-ordinated diorgano-phosphinate or -arsinate groups, giving almost symmetrical bridges. An interesting feature in the structures of compounds **5** and **6** is the shortness of the As–O bonds (mean 1.681 and 1.686 Å, respectively) compared with those in the quadruply bridged compounds **1** (mean 1.710 Å) and **4** (mean 1.702 Å) discussed above. This was to some extent expected as the As–O stretching vibrations in the IR spectra of **5** and **6** were at higher energy.

Owing to bridging, the tetrahedral geometry at phosphorus or arsenic is distorted with O–M–O angles increased to *ca.* 115° in the methyl-substituted products and to *ca.* 119° in the diphe-

Table 4 Selected bond distances (Å) and angles (°) for $[(SbPh_3)_2(\mu - O)(\mu - O_2MMe_2)_2] \cdot 0.5C_6H_{14}$ where M = P 3 or As 5 and for $[\{Sb(p-MeC_6H_4)_3\}_2(\mu - O)(\mu - O_2MMe_2)_2] \cdot 0.5C_6H_{14}$ where M = P 7, with e.s.d.s in parentheses

	3	5	7
Sb(1)-O(1)	1.958(3)	1.963(4)	1.972(5)
Sb(1) - O(3)	2.153(3)	2.122(4)	2.139(5)
Sb(1) - O(5)	2.171(3)	2.137(4)	2.170(5)
Sb(1)-C(11)	2.164(5)	2.177(6)	2.158(8)
Sb(1) - C(21)	2.152(5)	2.164(6)	2.163(8)
Sb(1) - C(31)	2,155(5)	2.164(6)	2,159(5)
Sb(2) - O(1)	1 952(3)	1 963(4)	1.954(5)
Sb(2) - O(2)	2,194(3)	2,154(4)	2,215(5)
Sb(2) - O(4)	2.154(3)	2.100(4)	2.213(5)
Sb(2)-C(41)	2.153(5)	2.173(7)	2.144(7)
Sb(2)-C(51)	2.163(5)	2.167(6)	2.144(7)
Sb(2) - C(61)	2.151(5)	2.154(7)	2.156(5)
M(1)-O(2)	1.519(4)	1.671(5)	1.521(5)
M(1)-O(3)	1.528(4)	1.685(5)	1.533(5)
M(1)–C(1)	1.798(5)	1.913(7)	1.783(8)
M(1)–C(2)	1.789(6)	1.911(7)	1.800(8)
M(2)–O(4)	1.514(4)	1.680(4)	1.524(5)
M(2)–O(5)	1.525(4)	1.687(5)	1.526(5)
M(2)–C(3)	1.787(6)	1.904(7)	1.788(8)
M(2)–C(4)	1.795(6)	1.916(7)	1.792(8)
O(1)–Sb(1)–O(3)	83.8(1)	85.2(2)	84.3(2)
O(1)–Sb(1)–O(5)	85.8(1)	87.7(2)	85.8(2)
O(3)–Sb(1)–O(5)	83.7(1)	83.1(2)	82.4(2)
O(3)-Sb(1)-C(11)	83.7(2)	85.3(2)	85.5(2)
O(5)-Sb(1)-C(11)	83.7(2)	83.4(2)	83.5(2)
O(1)-Sb(1)-C(21)	91.5(2)	90.8(2)	91.9(2)
O(3)-Sb(1)-C(21)	88.9(2)	88.4(2)	88.4(3)
C(11) - Sb(1) - C(21)	97.4(2)	96.7(2)	97.2(3)
O(1) - SD(1) - C(31)	91.5(2)	91.2(2)	92.4(2)
O(5) - SD(1) - C(31)	88.3(Z)	90.6(2) 07.4(9)	89.3(Z)
C(11) = SD(1) = C(31) C(21) = SD(1) = C(21)	99.0(<i>L</i>)	97.4(2) 07.8(2)	90.3(3)
O(1) = SD(1) = O(31)	90.7(2)	97.0(2)	99.0(3) 166 1(9)
O(1) = SD(1) = C(11) O(5) = SD(1) = C(21)	104.3(2) 179 $A(9)$	107.7(2) 171 5(9)	170 8(2)
O(3) - Sb(1) - C(21)	172.4(2)	172 8(2)	171 3(2)
O(1) - Sb(2) - O(2)	84 9(1)	85.4(2)	84 6(2)
O(1) - Sb(2) - O(4)	84.3(1)	86.0(2)	82.2(2)
O(2)-Sb(2)-O(4)	82.0(1)	82.7(2)	85.8(2)
O(2)-Sb(2)-C(41)	84.1(2)	84.6(2)	81.6(2)
O(4)-Sb(2)-C(41)	84.0(2)	85.0(2)	82.4(2)
O(1)-Sb(2)-C(51)	91.8(2)	92.8(2)	94.8(3)
O(4)-Sb(2)-C(51)	88.4(2)	88.7(2)	87.4(3)
C(41)-Sb(2)-C(51)	97.3(2)	95.9(3)	97.2(3)
O(1)–Sb(2)–C(61)	91.8(2)	91.1(2)	94.1(3)
O(2)–Sb(2)–C(61)	89.7(2)	90.5(2)	89.0(2)
C(41)-Sb(2)-C(61)	98.4(2)	96.7(3)	100.0(3)
C(51)-Sb(2)-C(61)	99.7(2)	98.0(3)	97.8(3)
O(1)-Sb(2)-C(41)	165.0(2)	167.3(2)	160.0(2)
O(2)-Sb(2)-C(51)	170.2(2)	171.4(2)	173.2(2)
O(4) - Sb(2) - C(61)	171.2(2)	172.8(2)	174.0(2)
O(2) - M(1) - O(3)	115.5(2)	110.0(2)	115.1(3) 107.1(4)
C(1) = W(1) = C(2) O(4) = M(2) = O(5)	107.7(3)	110.0(3) 112.0(2)	107.1(4) 114.9(2)
$O(4)^{-1}VI(2)^{-}O(3)$ $O(3)^{-}M(2)^{-}O(3)$	113.0(2)	113.3(2)	106 4(4)
Sh(1) = O(1) = Sh(2)	144 2(2)	145 5(2)	143 6(3)
Sb(2) - O(2) - M(1)	133.6(2)	130.2(2)	134.1(3)
Sb(1) - O(3) - M(1)	128.4(2)	123.6(2)	129.2(3)
Sb(2) - O(4) - M(2)	130.6(2)	126.2(2)	125.8(3)
Sb(1)-O(5)-M(2)	133.5(2)	129.6(2)	133.1(3)

nylarsinate **6**. The Sb–O–M angles are also increased and are larger in the phosphinates (mean 131.0°), as a consequence of the shorter P–O bond length, than in the corresponding arsinates (mean 127.6°). For comparison, these angles in the quadruply bridged arsinates (**1** and **2**) are further reduced to a mean of 119.2° presumably as the double oxygen bridge reduces the Sb · · · Sb separation.

It was not possible to assign unequivocally similar triply bridged structures to the trimethylantimony compounds, **8** and **9**, although such structures are likely. Decreased antimony

Sb(1)-O(1)	1.967(5)	Sb(2)-C(51)	2.166(7)
Sb(1)-O(3)	2.148(5)	Sb(2)-C(61)	2.148(7)
Sb(1)-O(5)	2.152(5)	As(1)–O(2)	1.699(5)
Sb(1)-C(11)	2.135(7)	As(1)–O(3)	1.680(4)
Sb(1)-C(21)	2.156(7)	As(1)-C(71)	1.926(7)
Sb(1)-C(31)	2.158(7)	As(1)-C(81)	1.921(7)
Sb(2)-O(1)	1.947(4)	As(2)–O(4)	1.686(5)
Sb(2)–O(2)	2.169(4)	As(2)–O(5)	1.677(5)
Sb(2)-O(4)	2.141(5)	As(2)–C(91)	1.919(7)
Sb(2)-C(41)	2.167(7)	As(2)–C(101)	1.923(7)
O(1)-Sb(1)-O(3)	86.6(2)	O(1)-Sb(2)-C(51)	91.9(2)
O(1)-Sb(1)-O(5)	86.4(2)	O(4)-Sb(2)-C(51)	90.4(2)
O(3)–Sb(1)–O(5)	80.2(2)	C(41)-Sb(2)-C(51)	97.4(3)
O(3)–Sb(1)–C(11)	86.3(2)	O(1)-Sb(2)-C(61)	92.6(2)
O(5)-Sb(1)-C(11)	83.4(2)	O(2)-Sb(2)-C(61)	86.7(2)
O(1)-Sb(1)-C(21)	90.9(2)	C(41)-Sb(2)-C(61)	95.7(3)
O(3)-Sb(1)-C(21)	89.5(2)	C(51)-Sb(2)-C(61)	100.1(3)
C(11)-Sb(1)-C(21)	98.1(3)	O(1)-Sb(2)-C(41)	166.2(3)
O(1)-Sb(1)-C(31)	87.7(2)	O(2)–Sb(2)–C(51)	173.1(2)
O(5)–Sb(1)–C(31)	89.7(2)	O(4) - Sb(2) - C(61)	169.4(2)
C(11)-Sb(1)-C(31)	97.7(3)	O(2) - As(1) - O(3)	118.8(2)
C(21)-Sb(1)-C(31)	100.4(3)	C(71)-As(1)-C(81)	109.3(3)
O(1)-Sb(1)-C(11)	168.4(2)	O(4) - As(2) - O(5)	118.6(2)
O(5)–Sb(1)–C(21)	169.4(2)	C(91)–As(2)–C(101)	106.8(3)
O(3)–Sb(1)–C(31)	168.7(2)	Sb(1)-O(1)-Sb(2)	151.7(3)
O(1)-Sb(2)-O(2)	86.3(2)	Sb(2)-O(2)-As(1)	125.9(2)
O(1)-Sb(2)-O(4)	85.9(2)	Sb(1)–O(3)–As(1)	129.1(3)
O(2)-Sb(2)-O(4)	82.8(2)	Sb(2)–O(4)–As(2)	128.6(2)
O(2)–Sb(2)–C(41)	83.3(2)	Sb(1)–O(5)–As(2)	127.4(2)
O(4)-Sb(2)-C(41)	83.9(2)		

Table 6 Comparison of Sb–O mean bridge parameters

Compound	Sb-O/Å	Sb-O-Sb/°	Ref.
3	1.955	144.2	This work
5	1.963	145.5	This work
6	1.957	151.7	This work
7	1.963	143.6	This work
$[{SbCl_3(O_2PMe_2)}_2O]$	1.942	136.0	15
$[{SbPh_2Cl[O_2P(C_6H_{11})_2]}_2O]$	1.937	144.7	16
$[{SbF_3(O_2CCF_3)}_2O]$	1.893	131.1	17

Lewis acidity as a consequence of the electron-donating methyl groups could, however, influence ligand secondary bonding leading to changes in co-ordination mode.

Experimental

Diphenylantimony bromide oxide was prepared by oxidising diphenylantimony(III) bromide with *tert*-butyl hydroperoxide²⁰ and μ -oxo-bis[bromotriphenylantimony(v)], μ -oxo-bis[chloro-triphenylantimony(v)] and μ -oxo-bis[bromotris(*p*-tolyl)-antimony(v)] by partial hydrolysis of the appropriate triaryl-antimony(v) dihalide.²⁴ μ -Oxo-bis[chlorotrimethylantimony(v)] was obtained by treating trimethylantimony dichloride with the corresponding dihydroxide.²⁵ Dimethylphosphinic acid was obtained by the method of Reinhardt *et al.*²⁶ and the sodium salts Na(O₂AsMe₂), Na(O₂AsPh₂), Na(O₂PMe₂) and Na-(O₂PPh₂) by treating stoichiometric amounts of the corresponding acid with sodium ethoxide in ethanol.

Preparations

 $[(SbPh_2)_2(\mu-O)_2(\mu-O_2AsMe_2)_2]$ **1.** A solution of diphenylantimony bromide oxide (4.00 g, 5.4 mmol) in dichloromethane (50 cm³) was added to a stirring suspension of sodium dimethylarsinate (1.90 g, 11.9 mmol) and dichloromethane (10 cm³) and the mixture refluxed for 24 h. After filtration and removal of the solvent, the remaining solid was recrystallised from chloroform

overlayered with twice the volume of hexane to give the product as colourless crystals. Yield 3.98 g (86%), m.p. 324–327 °C (Found: C, 39.4; H, 3.8. Calc. for $C_{28}H_{32}As_2O_6Sb_2$: C, 39.2; H, 3.8%); δ_H [250 MHz, CDCl₃, room temperature (r.t.)] 2.00 (12 H, s, AsMe), 7.33 [12 H, m, *m*- and *p*-H of Ph] and 7.83 (8 H, m, *o*-H of Ph); \tilde{v}_{max} /cm⁻¹ (Nujol) 1430s, 1072m, 817vs (sh), 796vs, 738s, 696s, 652m, 608m, 518s, 487s and 466m; FAB mass spectrum *m*/*z* 858 (100, *M*⁺), 721 (30, [*M* – O₂AsMe₂]⁺), 704 (28, [*M* – 2Ph]⁺), 567 (43, [*M* – 2Ph – O₂AsMe₂]⁺), 413 {26%, [(SbO)₂O₂AsMe₂]⁺}.

[(SbPh₂)₂(μ-O)₂(μ-O₂AsPh₂)₂] 2. This was prepared in a similar manner from [(SbPh₂BrO)₂] (3.2 g, 4 mmol) and sodium diphenylarsinate (2.60 g, 9.2 mmol) in dichloromethane (10 cm³) after refluxing for 40 h. A small amount of product was obtained after repeated crystallisations from chloroform-hexane. Yield 0.14 g (3%), m.p. 260–266 °C (Found: C, 51.2; H, 3.4. Calc. for C₄₈H₄₀As₂O₆Sb₂: C, 52.1; H, 3.6%); δ_H(250 MHz, CDCl₃, r.t.) 7.30–7.80 (40 H, br m, AsPh and SbPh); \tilde{v}_{max} /cm⁻¹ (Nujol) 1087w, 823vs (sh), 812vs, 735s, 691s, 490m and 457m; FAB mass spectrum *m*/*z* 1108 (73, [*M* + 2H]⁺), 953 (16, [*M* – 2Ph + H]⁺), 845 (38, [*M* – O₂AsPh₂]⁺), 691 {100, [(SbPh₂O)₂O₂As]⁺}, 537 (75, [SbPh₂O₂AsPh₂]⁺) and 459 (36%, [SbPhO₂AsPh₂]⁺).

[(SbPh₃)₂(µ-O)(µ-O₂PMe₂)₂] 3. (a) From [(SbPh₃Br)₂O]. A solution of µ-oxo-bis[bromotriphenylantimony(v)] (0.53 g, 0.60 mmol) in dichloromethane (20 cm³) was added to a stirring suspension of sodium dimethylphosphinate (0.16 g, 1.34 mmol) and dichloromethane (10 cm³) and the mixture refluxed for 24 h. After filtration and removal of the solvent, the product was obtained by overlayering a chloroform solution with twice the volume of hexane. Yield 0.48 g (88%), m.p. 188-192 °C (Found: C, 53.1; H, 5.1. Calc. for C₄₀H₄₂O₅P₂Sb₂: C, 52.9; H, 4.7%); δ_H(250 MHz, CDCl₃, r.t.) moist solvent, 1.00 [6 H, d, ²J(HP) 14, PMe], 7.57 (9 H, m, *m*- and *p*-H of Ph) and 8.21 (6 H, m, *o*-H of Ph); anhydrous solvent, species A, 0.72 [12 H, d, ²J(HP) 14, MeP], 7.16 [12 H, t, ³J(HH) 7, m-H of Ph], 7.45 [6 H, t, ³J(HH) 7, p-H of Ph] and 7.56 [12 H, d, 3J(HH) 7 Hz, o-H of Ph]; species B, 1.03 (12 H, br, PMe), 7.38 (18 H, m, m- and p-H of Ph) and 7.68 (12 H, m, *o*-H of Ph); \tilde{v}_{max}/cm^{-1} 1430m, 1301m, 1112vs, 1062m, 1046vs, 1032vs, 998w, 871m, 863m, 749s, 735s, 693s, 659w, 512w, 487w, 464m and 437w; FAB mass spectrum m/z 831 (4, $[M - Ph]^+$), 815 (9, $[M - O_2PMe_2]^+$), 661 {5, $[(SbPh_2)_2O(O_2PMe_2)]^+$ and 445 {100%, $[SbPh_3(O_2PMe_2)]^+$ }.

(b) From $[(SbPh_2BrO)_2]$. A mixture of $[(SbPh_2BrO)_2]$ (0.53 g, 0.7 mmol), sodium dimethylphosphinate (0.17 g, 1.4 mmol) and toluene (30 cm³) was stirred at reflux for 24 h. The mixture was filtered and the solvent removed in a vacuum to give a residue which was recrystallised from chloroform-hexane mixture. Yield 77 mg (12%).

[{Sb(p-MeC₆H₄)₂}₂(μ-O)₂(μ-O₂AsMe₂)₂] 4. A mixture of μoxo-bis[bromotris(p-tolyl)antimony(v)] (0.42 g, 0.43 mmol), sodium dimethylarsinate (0.17 g, 1.06 mmol) and dichloromethane (20 cm³) was stirred under reflux for 16 h. After filtration of the precipitated sodium bromide and evaporation of the solvent, the remaining white solid was recrystallised from a chloroform-hexane mixture to give the required product. Yield 0.10 g (26%), m.p. 313-320 °C (Found: C, 41.7; H, 4.45. Calc. for C32H40As2O6Sb2: C, 42.05; H, 4.4. Calc. for C46H54As2O5Sb2: C, 51.1; H, 5.0%); δH(250 MHz, CDCl3, r.t.) 1.96 (12 H, s, AsMe), 2.32 (12 H, s, MeC₆H₄), 7.15 [8 H, d, ³J(HH) 8, m-H of aryl] and 7.73 [8 H, d, ³J(HH) 8 Hz, o-H of aryl]; \tilde{v}_{max} /cm⁻¹ 1492m, 1307w, 1270w, 1183m, 1070m, 1018w, 901m, 869m, 824vs (sh), 805vs, 793s (sh), 722w, 652m, 609m, 578m, 516s, 488s and 430w; FAB mass spectrum m/z 915 (24, $[M + H]^+$), 577 {13, $[Sb(p-MeC_6H_4)_2(O_2AsMe_2)_2]^+$ }, 531 {100, $[Sb(p-MeC_{6}H_{4})_{3}(O_{2}AsMe_{2})]^{+}$ and 349 {12%, $[Sb(p-MeC_{6}H_{4}) (O_2AsMe_2)]^+$.

Table 7 Crystallographic data ^a for quadruply bridged compounds 1 and 4

$\begin{array}{cccc} \mbox{Chemical formula} & \mbox{C}_{30}\mbox{H}_{34}\mbox{As}_2\mbox{C}_{6}\mbox{Sb}_2 & \mbox{C}_{32}\mbox{H}_{40}\mbox{As}_2\mbox{O}_6\mbox{Sb}_2 & \mbox{M} & 1096.61 & 913.98 & \mbox{Crystal size/mm} & \mbox{0.30}\times 0.25\times 0.15 & \mbox{0.35}\times 0.24\times 0.06 & \mbox{Space group} & \mbox{P_2_1/a} & \mbox{P_2_1/c} & \mbox{as}_2\mbox{As}_2\mbox{O}_6\mbox{Space group} & \mbox{P_2_1/a} & \mbox{P_2_1/c} & \mbox{as}_2\mbox{As}_2\mbox{O}_6\mbox{Space group} & \mbox{P_2_1/a} & \mbox{P_2_1/c} & \mbox{as}_2\mbox{As}_2\mbox{O}_6\mbox{As}_2\mbox{As}_$		1	4
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Chemical formula	CarHarAsaCleOaSba	CasHasAsoOsSba
$\begin{array}{cccccccccccccccccccccccccccccccccccc$	Μ	1096.61	913.98
$\begin{array}{c ccccccccccccccccccccccccccccccccccc$	Crystal size/mm	$0.30 \times 0.25 \times 0.15$	$0.35 \times 0.24 \times 0.06$
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	Space group	$P2_1/a$	$P2_1/c$
b/Å10.2590(3)11.4790(10) $c/Å$ 14.5744(12)11.2265(10) $\beta/^\circ$ 113.13(2)99.411(5) $U/Å^3$ 1901.5(7)1701.3(3) $D_c/g \ cm^{-3}$ 1.9151.784 μ/cm^{-1} 36.0935.58 $F(000)$ 1064896Index ranges (for unique data) $-16 < h < 14, 0 < k < 12, 0 < l < 17$ $-10 < h < 16, -12 < k < 13, -11 < l < 13$ Total data collected80517176Unique data29322960 R_{int} 0.07660.0502	a/Å	13.830(3)	13.3821(13)
$c/Å$ 14.5744(12)11.2265(10) $\beta/^{\circ}$ 113.13(2)99.411(5) $U/Å^3$ 1901.5(7)1701.3(3) $D_c/g {\rm cm}^{-3}$ 1.9151.784 $\mu/{\rm cm}^{-1}$ 36.0935.58 $F(000)$ 1064896Index ranges (for unique data) $-16 < h < 14, 0 < k < 12, 0 < l < 17$ $-10 < h < 16, -12 < k < 13, -11 < l < 13$ Total data collected80517176Unique data29322960 $R_{\rm int}$ 0.07660.0502	b/Å	10.2590(3)	11.4790(10)
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	c/Å	14.5744(12)	11.2265(10)
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	β/°	113.13(2)	99.411(5)
$ \begin{array}{cccccccccccccccccccccccccccccccccccc$	$U/Å^3$	1901.5(7)	1701.3(3)
$ \begin{array}{cccc} \mu/cm^{-1} & 36.09 & 35.58 \\ F(000) & 1064 & 896 \\ Index ranges (for unique data) & -16 < h < 14, 0 < k < 12, 0 < l < 17 \\ Total data collected & 8051 & 7176 \\ Unique data & 2932 & 2960 \\ R_{int} & 0.0766 & 0.0502 \end{array} $	$D/g \text{ cm}^{-3}$	1.915	1.784
$F(000)$ 1064896Index ranges (for unique data) $-16 < h < 14, 0 < k < 12, 0 < l < 17$ $-10 < h < 16, -12 < k < 13, -11 < l < 13$ Total data collected80517176Unique data29322960 R_{int} 0.07660.0502	μ/cm^{-1}	36.09	35.58
Index ranges (for unique data) $-16 < h < 14, 0 < k < 12, 0 < l < 17$ $-10 < h < 16, -12 < k < 13, -11 < l < 13$ Total data collected80517176Unique data29322960 $R_{\rm int}$ 0.07660.0502	F(000)	1064	896
Total data collected 8051 7176 Unique data 2932 2960 R_{int} 0.0766 0.0502	Index ranges (for unique data)	-16 < h < 14, 0 < k < 12, 0 < l < 17	-10 < h < 16, -12 < k < 13, -11 < l < 13
Unique data 2932 2960 R _{int} 0.0766 0.0502	Total data collected	8051	7176
R _{int} 0.0766 0.0502	Unique data	2932	2960
in the second	R _{int}	0.0766	0.0502
Observed data $[I > 3\sigma(I)]$ 2172	Observed data $[I > 3\sigma(I)]$	2172	
Absorption correction DIFABS ^b Gaussian	Absorption correction	DIFABS ^b	Gaussian
minimum 0.961 0.394	minimum	0.961	0.394
maximum 1.044 0.810	maximum	1.044	0.810
Structure solution Direct (SIR 92) Direct (SHELXS 86)	Structure solution	Direct (SIR 92)	Direct (SHELXS 86)
Refinement Full-matrix least squares on F Full-matrix least squares on F^{2}	Refinement	Full-matrix least squares on F	Full-matrix least squares on F^2
Data, variables 2172, 209 2933, 195	Data, variables	2172, 209	2933, 195
Goodness of fit (<i>S</i>) 1.309 1.065	Goodness of fit (S)	1.309	1.065
Final difference map features/e Å ⁻³ +1.18, -0.70 +0.604, -1.120	Final difference map features/e Å ⁻³	+1.18, -0.70	+0.604, -1.120
R 0.0276	R	0.0276	
<i>R</i> ′ 0.0306	R'	0.0306	
<i>R</i> observed data $[I > 2\sigma(I)]$ (all data) 0.0315 (0.0355)	<i>R</i> observed data $[I > 2\sigma(I)]$ (all data)		0.0315 (0.0355)
<i>R</i> ' observed data (all data) 0.0794 (0.0873)	R' observed data (all data)		0.0794 (0.0873)

^{*a*} Details in common: monoclinic; Z = 2; Mo-K α radiation ($\lambda 0.710$ 69 Å); θ limits 3–25°; 150 K. ^{*b*} Ref. 27.

Table 8 Crystallographic data * for triply bridged compounds 3 and 5-7

	3	5	6	7	
Chemical formula	$C_{43}H_{49}O_5P_2Sb_2$	$C_{43}H_{49}As_2O_5Sb_2$	C ₆₁ H ₅₁ As ₂ Cl ₃ O ₅ Sb ₂	$C_{49}H_{61}O_5P_2Sb_2$	
M	951.30	1039.16	1363.77	1035.47	
Crystal size/mm	0.20 imes 0.18 imes 0.14	0.32 imes 0.20 imes 0.16	0.15 imes 0.10 imes 0.10	0.24 imes 0.22 imes 0.18	
Space group	C2/c	C_2/c	$P2_1/a$	$P2_1/n$	
aĺÅ	26.704(3)	27.176(3)	11.804(12)	12.038(12)	
b/Å	16.879(3)	16.826(2)	40.910(4)	17.127(13)	
c/Å	22.1874(12)	22.637(3)	12.738(4)	23.35(2)	
β/°	125.35(1)	126.887(8)	115.71(4)	90.27(3)	
$U/Å^3$	8157.0(6)	8278.7(2)	5542.2(6)	4814.6(3)	
Ζ	8	8	4	4	
$D_{\rm c}/{\rm g~cm^{-3}}$	1.549	1.667	1.635	1.429	
μ/cm^{-1}	14.52	29.34	23.57	12.36	
F(000)	3832	4120	2704	2108	
θ Limits/°	3–25	2-25	3-25	3-25	
Index ranges (for unique data)	-29 < h < 23,	-29 < h < 30,	-13 < h < 11,	-13 < h < 13,	
	0 < k < 18, 0 < l < 25	0 < <i>k</i> < 18, 0 < <i>l</i> < 24	0 < k < 45, 0 < l < 14	0 < <i>k</i> < 19, 0 < <i>l</i> < 26	
<i>T</i> /K	120	150	120	150	
Total data collected	17 558	17 594	21 169	23 252	
Unique data	6171	6406	7933	7414	
R _{int}	0.0481	0.0913	0.1616	0.0937	
Observed data $[I > 3\sigma(I)]$	4315		6128	4900	
Absorption correction	DIFABS	None	DIFABS	DIFABS	
minimum	0.855		0.773	0.849	
maximum	1.160		1.115	1.118	
Structure solution	Direct (SIR 92)	Direct (SHELXS 86)	Direct (SIR 92)	Direct (SIR 92)	
Data, variables	4315, 455	6406, 467	6128, 658	4900, 517	
Goodness of fit (S)	1.080	0.935	1.073	1.134	
Final difference map features/e Å ⁻³	+1.10, -0.73	+1.736, -0.643	+2.09, -2.05	+0.74, -0.49	
R	0.0344		0.0702	0.0367	
R'	0.0698		0.0942	0.0912	
<i>R</i> observed data $[I > 2\sigma(I)]$ (all data)		0.0422 (0.0633)			
R' observed data (all data)		0.0940 (0.1247)			
* Details in common: monoclinic; Mo-K α radiation (λ 0.710 69 Å); full-matrix least-squares refinement on F^{2} .					

 $[(SbPh_3)_2(\mu-O)(\mu-O_2AsMe_2)_2]$ 5. A mixture of μ -oxobis[bromotriphenylantimony(v)] (0.74 g, 0.84 mmol), sodium dimethylarsinate (0.36 g, 2.3 mmol) and dichloromethane (35 cm³) was stirred at reflux for 20 h. After filtration and removal

of the solvent in a vacuum, the product was obtained by diffusion of hexane vapour into a concentrated chloroform solution. Yield 0.62 g (74%), m.p. 241–245 °C (Found: C, 47.85; H, 4.3. Calc. for $C_{40}H_{42}As_2O_5Sb_2$: C, 48.2; H, 4.25%); $\delta_H(250~MHz,$

CDCl₃, r.t.) 0.71 (br s, AsMe), 1.89 (br s, AsMe) (integrals unreliable due to peak broadness); phenyl region, species A, 7.17 [12 H, t, ³J(HH) 7, m-H of Ph], 7.27 [6 H (masked), t, p-H of Ph] and 7.61 [12 H, d, ³J(HH) 7 Hz, o-H of Ph]; species B, 7.27 (18 H, m, m- and p-H of Ph) and 7.71 (12 H, m, o-H of Ph); ṽ_{max}/cm⁻¹ 1429s, 1265m, 1185m, 1069s, 1023w, 852vs (br), 738vs, 698vs, 657m, 641m, 465s and 421m; FAB mass spectrum m/z 996 (0.4, M^+), 859 (6, $[M - O_2AsMe_2]^+$), 489 {61, $[SbPh_3(O_2AsMe_2)]^+$, 335 {28, $[SbPh(O_2AsMe_2)]^+$ } and 275 $(39\%, \text{SbPh}_2^+).$

 $[(SbPh_3)_2(\mu-O)(\mu-O_2AsPh_2)_2]$ 6. A mixture of μ -oxobis[bromotriphenylantimony(v)] (0.60 g, 0.68 mmol), sodium diphenylarsinate (0.49 g, 1.7 mmol) and dichloromethane (35 cm³) was stirred at reflux for 24 h. After filtration and removal of the solvent, the residue was purified by diffusion of hexane into a concentrated chloroform solution. Yield 0.63 g (74%), m.p. 244-247 °C (Found: C, 53.1; H, 3.7. Calc. for C₆₀H₅₀-As₂O₅Sb₂: C, 57.9; H, 4.05. Calc. for C₆₀H₅₀As₂O₅Sb₂·CHCl₃: C, 53.7; H, 3.8%); δ_H(250 MHz, CDCl₃, r.t.) 6.97 (m), 7.08 (m), 7.33 (m), 7.43 (m), 7.58 [d, ³J(HH) 7 Hz] and 8.04 (m); $\tilde{\nu}_{max}/cm^{-1}$ (Nujol) 1440s, 1430s, 1306m, 1183m, 1090s, 1069s, 1023m, 999m, 831vs (br), 735vs, 692vs, 658m, 609w, 536m, 464s and 410m.

 $[{Sb(p-MeC_6H_4)_3}_2(\mu-O)(\mu-O_2PMe_2)_2]$ 7. A solution of μ -oxobis[bromotris(p-tolyl)antimony(v)] (0.59 g, 0.61 mmol) in dichloromethane (20 cm³) was added to a suspension of sodium dimethylphosphinate (0.15 g, 1.3 mmol) in dichloromethane (10 cm³) which was stirred at room temperature for 18 h. After filtration and removal of the solvent in a vacuum, the remaining white solid was purified by overlayering a chloroform solution with three times the volume of hexane. Yield 0.35 g (58%) (Found: C, 55.2; H, 5.4. Calc. for C46H54O5P2Sb2: C, 55.7; H, 5.5%); 8H(250 MHz, CDCl3, r.t.) moist solvent, 0.97 [6 H, d, ² J(HP) 14, PMe], 2.40 (9 H, s, MeC₆H₄), 7.51 [6 H, d, ³ J(HH) 8, m-H of aryl] and 8.12 [6 H, d, ³J(HH) 8, o-H of aryl]; anhydrous solvent, species A, 0.70 [12 H, d, 2J(HP) 14, PMe], 2.38 (18 H, s, MeC₆H₄), 7.10 [12 H, d, ³J(HH) 8, *m*-H of aryl] and 7.55 [12 H, d, ³J(HH) 8, o-H of aryl]; species B, 1.05 (12 H, br, PMe), 2.39 (18 H, s, MeC₆H₄), 7.25 [12 H, d, ³J(HH) 8, m-H of aryl] and 7.61 [12 H, d, ${}^{3}J$ (HH) 8 Hz, o-H of aryl]; \tilde{v}_{max} /cm⁻¹ (Nujol) 1298m, 1187m, 1120s, 1111vs, 1048vs, 1033vs, 863s, 801s, 732s, 705w, 576w and 487s; FAB mass spectrum m/z 962 (4, $[M - 2Me]^+$), 899 (0.4, $[M - O_2PMe_2]^+$) and 487 {100%, $[Sb(p-MeC_6H_4)_3(O_2PMe_2)]^+$.

 $[(SbMe_3)_2(\mu-O)(\mu-O_2PMe_2)_2]$ 8. A solution of μ -oxobis[chlorotrimethylantimony(v)] (0.84 g, 2 mmol) in dichloromethane (20 cm³) was added to a suspension of sodium dimethylphosphinate (0.46 g, 4.4 mmol) also in dichloromethane (20 cm³) at 0 °C under an atmosphere of argon and the mixture stirred at room temperature for 6 h. The mixture was filtered and the clear filtrate concentrated to ca. 5 cm³. Overlayering with hexane (15 cm³) gave extremely moisture-sensitive crystals of the required compound, which were isolated by decantation and dried in a vacuum. Yield 0.90 g (84%). Owing to the extreme air sensitivity of this compound, satisfactory microanalytical results could not be obtained. $\delta_{\rm H}(250 \text{ MHz},$ CDCl₃, r.t., Teflon-sealed tube under N₂) 1.13 [12 H, d, ²J(HP) 14, PMe] and 1.66 (18 H, s, SbMe); 8c (69 MHz, CDCl₃, r.t.) 16.3 (s, SbMe) and 18.8 [d, ${}^{1}J(CP)$ 104 Hz, PMe]; \tilde{v}_{max}/cm^{-1} 1421s, 1286s, 1262w, 1221s, 1172vs (br), 1020vs (br), 851vs (br), 753vs (br), 692s, 578s, 531m, 512w, 482vs and 420vs (possible decomposition in Nujol).

[(SbMe₃)₂(µ-O)(µ-O₂AsPh₂)₂] 9. A mixture of µ-oxobis[chlorotrimethylantimony(v)] (0.39 g, 0.93 mmol), sodium diphenylarsinate (0.28 g, 1.0 mmol) and dichloromethane (35 cm³) at 0 °C was treated as described above to yield moisturesensitive crystals of the required compound, shown by ¹H NMR spectroscopy to be a dichloromethane solvate. Again satisfactory microanalytical results could be obtained. Yield 0.60 g (75%). $\delta_{\rm H}(250~{\rm MHz},~{\rm CDCl}_3,~{\rm r.t.},~{\rm Teflon-sealed}~{\rm tube}~{\rm under}~{\rm N_2})$ 1.66 (18 H, s, MeSb), 5.27 (2 H, s, CH₂Cl₂), 7.41 (12 H, m, mand p-H of Ph) and 7.70 (8 H, m, o-H of Ph); δ_c (69 MHz, CDCl₃, r.t.) 16.8 (SbMe), 128.7 (m-C of Ph), 130.4 (p-C of Ph), 131.2 (o-C of Ph) and 138.2 (ipso-C of Ph); \tilde{v}_{max}/cm^{-1} 1307m, 1216w, 1180w, 1157w, 1088s, 1066m, 1025m, 998m, 896s, 795vs (br), 740vs (br), 720vs (br), 692vs (br), 574s, 526w, 481s and 462s (possible decomposition in Nujol).

Crystallography

Crystallographic quality single crystals for all compounds were obtained by slow diffusion of hexane vapour into concentrated chloroform solutions of the compounds.

Crystallographic data are summarised in Table 7 for the quadruply bridged compounds and in Table 8 for the triply bridged species. The tables also contain details of the method of solution and the refinement conditions. Approximately one hemisphere of data was collected at low temperature using either a Delft Instruments FAST TV area-detector diffractometer, equipped with rotating-anode FR591 generator (for 1, 3 and 5-7) or a Siemens SMART area-detector diffractometer (for 4). The data were corrected for Lorentz-polarisation effects, merged and systematically absent reflections removed. An absorption correction was usually applied (see Tables 7 and 8). Structures were solved by direct methods (SIR 92²⁸ or SHELXS 86²⁹) and refined by full-matrix least squares (SHELXL 93³⁰ or CRYSTALS³¹); hydrogen atoms were placed at their calculated positions and refined riding on their respective carbon atoms. A standard weighting scheme was applied and corrections were made for extinction where appropriate.

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