# Weng Kee Leong\* and Guizhu Chen

Department of Chemistry, National University of Singapore, Kent Ridge, 119260, Singapore

Received 15th August 2000, Accepted 28th September 2000 First published as an Advance Article on the web 6th November 2000

Besides the cluster  $[Os_3(\mu-H)(\mu-SbPh_2)(CO)_{10}]$ , 1, other products identified from the reaction of  $[Os_3(\mu-H)(CO)_{10}-(\mu-CO)]^-$  with  $Ph_2SbCl$  in THF were  $[Os_3(\mu-SbPh_2)_2(CO)_{10}]$ , 2a, and a dimeric form of 1, viz.  $[Os_3(\mu-H)(\mu-SbPh_2)-(CO)_{10}]_2$ , 3. Carrying out the reaction in  $CH_2Cl_2$  gave 3 as the major product, while 2a has been demonstrated to result from the reaction of 1 with  $Ph_2SbCl$ . In 2a the  $Os_3Sb_2$  unit is almost planar; one of the  $Ph_2Sb$  bridges is open while the other is closed. The two  $Ph_2Sb$ -bridged  $Os_2Os$  edges are shown by EXSY and variable-temperature  $Ph_2Sb$ -bridged  $Ph_2S$ 

## Introduction

One of the most exciting developments in organometallic cluster chemistry in recent years has been that of main grouptransition metal cluster compounds. These compounds are expected to show structural and reactivity patterns that may be quite unlike those of the homometallic main group or transition metal clusters, the result of the interplay between the differing properties of the elements. Furthermore, there is a steady movement towards the view that the main group elements in many such compounds should be better regarded as an integral part of the cluster core, rather than as ligands. 1-3 We have been interested in this view for some time now, and one of the systems that we have actively been studying is antimonyosmium.<sup>4,5</sup> We had chosen  $[Os_3(\mu-H)(\mu-SbPh_2)(CO)_{10}]$ , 1, as the starting point of our investigations in the belief that the available report on its preparation was a straightforward, high-yielding reaction.<sup>6</sup> In the course of our own attempts at the synthesis of 1, however, we have found that the products obtainable are dependent on the solvent system and the reaction time.

### Results and discussion

When the reaction of  $[Et_4N][Os_3(\mu-H)(CO)_{10}(\mu-CO)]$  and an excess of  $Ph_2SbCl$  was carried out in THF three cluster products were separated and identified, viz.  $[Os_3(\mu-H)(\mu-Sb-Ph_2)(CO)_{10}]$ , 1,  $[Os_3(\mu-SbPh_2)(CO)_{10}]$ , 2a, and  $[Os_3(\mu-H)(\mu-Sb-Ph_2)(CO)_{10}]$ , 3, in 60, 10 and 15% yields, respectively; cluster 1 was the only identified product in the earlier report. When the reaction was carried out over a longer period of time the yield of 2a increased at the expense of 1. Furthermore, when the reaction solvent was changed to  $CH_2Cl_2$  the major compound isolated was 3; no 1 was formed.

The cluster 3 may be regarded as a dimeric form of 1 in which the SbPh<sub>2</sub> moieties bridge across two triosmium units rather than across one edge of such a unit. This structural type has only been observed in one other instance, a ruthenium–phosphorus analogue  $[Ru_3(\mu-H)\{\mu-P(CF_3)_2\}(CO)_{10}]_2$ , which was obtained in 3% yield. That the formation of 3 was solvent-

dependent was an interesting point. Furthermore, we noted that the reaction in THF was almost immediate whereas that in  $CH_2Cl_2$  was much slower (as judged by the colour changes in the reaction mixtures). We have also established that 1 and 3 do not interconvert at ambient or slightly elevated temperatures, which is evidence that their formation follows different pathways. One plausible reaction scheme is shown in Scheme 1.

Scheme 1

The presence of moisture or possibly acid (from hydrolysis of Ph<sub>2</sub>SbCl) when the more polar solvent THF is employed may have resulted in protonation of  $[Os_3(\mu-H)(CO)_{10}(\mu-CO)]^-$  to form the cluster  $[Os_3H(\mu-H)(CO)_{11}]$ . This species is known to decarbonylate easily to form the unsaturated cluster  $[Os_3-(\mu-H)_2(CO)_{10}]$  which can undergo nucleophilic addition with Ph<sub>2</sub>SbCl to generate the species  $[Os_3H(CO)_{10}(SbPh_2Cl)]$ ; subsequent elimination of HCl would result in the formation of 1. The protonation step would account for the accelerated rate of

<sup>†</sup> Electronic supplementary information (ESI) available: EXSY spectrum of compound **2b**. See http://www.rsc.org/suppdata/dt/b0/b006669i/

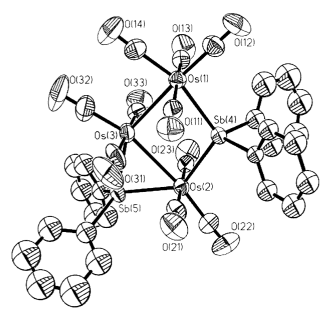


Fig. 1 An ORTEP diagram (50% thermal ellipsoids) for compound 2a

colour change in THF. In accord with this proposed route, we have observed that the reaction of the unsaturated cluster  $[Os_3(\mu-H)_2(CO)_{10}]$  with  $Ph_2SbCl$  indeed gave 1 in good yield. We have previously observed that 1 does have a tendency to undergo nucleophilic addition by cleavage of the  $SbPh_2$ -bridged OsOs bond, *i.e.* antimony prefers formation of an open  $SbPh_2$  bridge. It is therefore possible that the proposed intermediate  $[Os_3H(SbPh_2)(CO)_{11}]$  in route **b** may prefer dimerisation to form 3 (with open  $SbPh_2$  bridges) over the formation of 1 (with a closed  $SbPh_2$  bridge). In accord with this argument we have found that, unlike 1, cluster 3 did not undergo nucleophilic addition with  $PPh_3$  at ambient temperatures.

Cluster 2a comprised a planar Os<sub>3</sub>Sb<sub>2</sub> metal core (Fig. 1); this planarity persisted in solution, as was evident by the presence of resonances corresponding to only two different phenyl environments in the aromatic region. We established that 1 reacted with Ph<sub>2</sub>SbCl to afford 2a in essentially quantitative yield. A plausible mechanism for the formation of 2a is therefore that depicted in Scheme 2. The structure of the intermedi-

ate was suggested by our earlier observation that nucelophiles (including SbPh<sub>3</sub>) reacted with 1 to form adducts with the structure shown.<sup>4</sup> Indeed, monitoring of the reaction by  $^{1}H$  NMR spectroscopy showed the formation of an intermediate with a resonance at  $\delta$  -7.72. Although we did not obtain an IR

Scheme 2

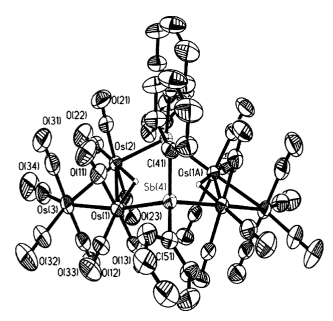
spectrum of the intermediate, we did manage to do so for the corresponding intermediate formed from the reaction between 1 and  $(p\text{-MeC}_6H_4)_2\text{SbCl}$  (see also below); the difference IR spectrum showed CO absorptions at 2114w, 2081m, 2069m, 2031s, 2007m and 1970w,br cm<sup>-1</sup>, which has the same pattern as that of  $[\text{Os}_3\text{H}(\text{SbPh}_2)(\text{CO})_{10}(\text{PPh}_3)]$ ; the <sup>1</sup>H NMR spectrum for this showed a resonance at  $\delta$  –7.76. Both the <sup>1</sup>H NMR and IR data are thus consistent with the proposed structure.

In an attempt to verify the last step, which involved rearrangement of an Os-Os bond, cluster 1 was treated with Ph<sub>2</sub>PCl with the view that if the proposal was correct we would form cluster 4 as the isomer A (Scheme 2); the phosphorus atom would also provide a useful <sup>31</sup>P NMR probe. As it turned out, cluster 4 was found to have the structure of isomer B. It may be that a cluster with structure A was indeed formed initially but it isomerised quickly to isomer B, the driving force for this rearrangement being the lesser tendency for the smaller phosphorus atom in PR<sub>2</sub> bridges to span open Os-Os edges.

To circumvent this isomerisation process, [Os<sub>3</sub>(μ-SbPh<sub>2</sub>)- $\{\mu\text{-Sb}(C_6H_4Me-p)\}_2(CO)_{10}\}$ , was prepared from the reaction of 1 with (p-MeC<sub>6</sub>H<sub>4</sub>)<sub>2</sub>SbCl. The <sup>1</sup>H NMR spectrum of 2b showed two sets of signals due to the CH<sub>3</sub> groups, indicating the presence of two isomers in solution which differed in whether the SbPh<sub>2</sub> or the Sb( $C_6H_4Me-p$ )<sub>2</sub> group formed the open bridge. A variable temperature <sup>1</sup>H NMR spectroscopic study (in d<sub>8</sub>toluene) showed that these two sets of signals coalesced at 85 °C, corresponding to an exchange rate of 14.6 s<sup>-1</sup> or a  $\Delta G^{\ddagger}$ of 80.2 kJ mol<sup>-1</sup> at that temperature; the coalescence was reversible and indicated no decomposition. An EXSY experiment was also carried out to confirm chemical exchange between these two sets of signals. Although the two CH<sub>3</sub> signals were too close to each other ( $\delta$  2.35 and 2.38, respectively) so that any crosspeaks between them would lie very near to the main diagonal and difficult to observe, it was fortuitous that we were able to identify the *ortho-H* signals of the *p*-tolyl groups in the two isomers, which were traced via NOE crosspeaks. Hence, the two CH<sub>3</sub> signals at  $\delta$  2.35 and 2.38 (labelled I and II, respectively) showed NOE crosspeaks with the aromatic signals at  $\delta$  7.17 and 7.21, respectively, which were therefore assignable to the respective *meta-H* of the tolyl groups. These in turn showed NOE crosspeaks to signals at  $\delta$  7.43 and 7.58, which may thus be assigned to the corresponding *ortho*-H of the *p*-tolyl groups in the two isomers I and II, respectively. It was not possible to decide which of I and II has the structure A (and hence which B), but these ortho-H resonances were sufficiently apart in their chemical shifts so that exchange crosspeaks between them were clearly observed, confirming the presence of chemical exchange between I and II. We have attributed this chemical exchange as being mediated through a metal-metal bond fluxion, one of as-yet relatively few examples. 10

## Crystallographic study

The molecule of compound 3 is located on a crystallographic twofold axis; the ORTEP diagram and selected bond parameters are given in Fig. 2. The most obvious feature is the two SbPh<sub>2</sub> bridges that link together two triosmium units. The central six-membered Os<sub>4</sub>Sb<sub>2</sub> ring is puckered about the Sb atoms into a "butterfly-like" conformation; the dihedral angle between the two SbOs<sub>2</sub>Sb planes is 39.8°. This brings one phenyl ring from each SbPh, unit into a mutually cis orientation. The OsSbOs angle is rather large; at 121.61(2)°, it is very much larger than the value of just below 106° found in clusters of the type Os<sub>3</sub>H(SbPh<sub>2</sub>)(CO)<sub>10</sub>(L) which also contain an SbPh<sub>2</sub> unit bridging two non-bonded osmium atoms. The structural features about each of the triosmium units are fairly typical: the Os-Os edge believed to be bridged by the metal hydride is appreciably longer (3.0693(4) Å) than the other two edges (2.8881(4) and 2.8950(4) Å), and the Os···O distances (sum of Os-C and C-O bond lengths) are longer for the axial



**Fig. 2** An ORTEP diagram (50% thermal ellipsoids) and selected bond parameters for compound **3**. Os(1)–Os(2) 3.0693(4); Os(2)–Os(3) 2.8881(4); Os(1)–Os(3) 2.8950(4); Os(1)–Sb(4) 2.6881(6); and Os(2A)–Sb(4) 2.6823(5) Å; Os(1)–Sb(4)–Os(2A) 121.61(2); Os(1)–Os(2)–Os(3) 58.055(10); Os(2)–Os(3)–Os(1) 64.110(10); and Os(3)–Os(1)–Os(2) 57.835(10)°.

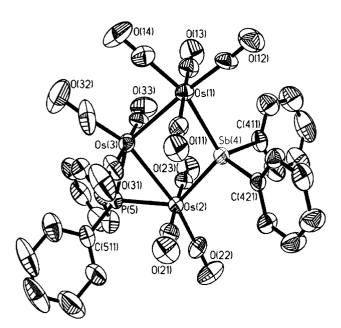


Fig. 3 An ORTEP diagram (50% thermal ellipsoids) for compound 4.

(3.080 to 3.101 Å) than for the equatorial carbonyls (3.034 to 3.055 Å).

Clusters **2a** and **4** are isostructural; the ORTEP diagram for **4** showing the atomic labelling scheme is given in Fig. 3, and selected bond parameters for **2a** and **4** are given in Table 1. The  $Os_3SbE$  unit is almost coplanar, the dihedral angle between Os(1)Os(3)Os(2)Sb(4) and Os(2)Os(3)E(5) being 1.2 and 1.6° for **2a** and **4**, respectively. This is in sharp contrast to that of the corresponding selenium analogues,  $[Os_3(\mu\text{-SeR})_2(CO)_{10}]$  (R = Me or Ph), which have the SeR group bridging the closed Os–Os edge far out of the  $Os_3Se$  plane, in an axial-type position. This difference in structure is presumably due to the steric constraint imposed by the  $SbR_2$  group; the presence of another R group in  $SbR_2$  as compared to SeR implies that any bending of one of the  $SbR_2$  groups out of the  $Os_3Se$  plane would meet with steric hindrance from the axial carbonyls. The

Table 1 Selected bond lengths (Å) and bond angles (°) for compounds 2a and 4

	2a (E = Sb)	4(E = P)
Os(1)–Os(3)	2.9967(6)	2.9968(5)
Os(2)-Os(3)	3.0408(6)	2.9436(5)
Os(1)-Sb(4)	2.7132(8)	2.6993(6)
Os(2)-Sb(4)	2.6384(8)	2.6334(6)
Os(2)-E(5)	2.6164(9)	2.350(2)
Os(3)-E(5)	2.6023(9)	2.311(2)
Os(1)–Sb(4)–Os(2)	110.61(3)	109.07(2)
Os(1)-Os(3)-Os(2)	93.574(17)	93.965(14)
Os(3)-E(5)-Os(2)	71.28(2)	78.33(7)

most interesting feature is the large difference in the Os(2)—Os(3) bond lengths (3.0408(6) Å and 2.9436(5) Å for **2a** and **4**, respectively), which is in sharp contrast with the essentially identical Os(1)—Os(3) bond lengths. This is a clear example of the effect of the size of the bridging atom on the length of the bridged edge.

## **Experimental**

### General procedures

All reactions and manipulations were performed under a nitrogen atmosphere by using standard Schlenk techniques. Solvents were purified, dried, distilled, and kept under nitrogen prior to use. Routine <sup>1</sup>H and <sup>31</sup>P NMR spectra were recorded on a Bruker ACF300 NMR spectrometer in CDCl<sub>3</sub> unless otherwise stated. The variable-temperature study was carried out on a Bruker DPX300 NMR spectrometer while the EXSY was carried out on a Bruker AVANCE DRX500 NMR spectrometer using a mixing time of 450 μs. Microanalyses were carried out by the microanalytical laboratory at the National University of Singapore. The cluster [Et<sub>4</sub>N]-[Os<sub>3</sub>(μ-H)(CO)<sub>10</sub>(μ-CO)], Ph<sub>2</sub>SbCl and (*p*-MeC<sub>6</sub>H<sub>4</sub>)<sub>2</sub>SbCl were prepared according to literature methods; <sup>12,13</sup> all other reagents were from commercial sources and used as supplied.

# Reactions of [Et<sub>4</sub>N][Os<sub>3</sub>( $\mu$ -H)(CO)<sub>10</sub>( $\mu$ -CO)] and Ph<sub>2</sub>SbCl

In THF. To a solution of  $[Et_4N][Os_3(\mu-H)(CO)_{10}(\mu-CO)]$  (100 mg, 0.099 mmol) in THF (20 mL) was added Ph<sub>2</sub>SbCl (60 mg, 0.193 mmol). The solution changed from dark red to yellow immediately. After stirring for 30 min the solvent was removed in vacuo and the residue separated by TLC with CH<sub>2</sub>Cl<sub>2</sub>-hexane (20:80, v/v) as eluent to afford two major bands. Band 1 gave orange crystals of  $[Os_3(\mu-H)(\mu-SbPh_2)(CO)_{10}]$ , 1 (67.6 mg, 60%). IR (v(CO), hexane): 2102mw, 2068w, 2054vs, 2022s, 2012ms, 1988mw and 1990m cm<sup>-1</sup>. <sup>1</sup>H NMR (CDCl<sub>3</sub>):  $\delta$  –19.84 (s, OsHOs). Calc. for  $C_{22}H_{11}O_{10}Sb$ : C, 23.43; H, 0.98. Found: C, 23.67; H, 0.98%. Band 2 gave a mixture of [Os<sub>3</sub>(Sb- $Ph_2_2(CO)_{10}$ , **2a** (14 mg, 10%), and  $[Os_3(\mu-H)(\mu-SbPh_2)(CO)_{10}]_2$ , 3 (17 mg, 15%). These two compounds were separated by fractional recrystallisation. 2a: IR (v(CO), hexane) 2102mw, 2054mw, 2033 (sh), 2026s, 2001w, 1983mw, 1972w and 1960mw cm<sup>-1</sup> (Calc. for  $C_{34}H_{20}O_{10}Os_3Sb_2\cdot\frac{1}{2}C_6H_{14}$ : C, 30.74; H, 1.80. Found: C, 30.86; H, 2.09%. 3:  $\bar{IR}$  ( $\nu$ (CO), hexane) 2097s, 2066m, 2051m, 2032vs, 2017m and 2006s cm<sup>-1</sup>; <sup>1</sup>H NMR (toluene)  $\delta - 18.68$  (s, OsHOs) (Calc. for  $C_{22}H_{11}O_{10}Os_3Sb$ : C, 23.43; H, 0.98. Found: C, 23.62; H, 1.24%).

In CH<sub>2</sub>Cl<sub>2</sub>. To a solution of [Et<sub>4</sub>N][Os<sub>3</sub>( $\mu$ -H)(CO)<sub>10</sub>( $\mu$ -CO)] (100 mg, 0.099 mmol) in dichloromethane (20 mL) was added Ph<sub>2</sub>SbCl (60 mg, 0.193 mmol). The reaction mixture was allowed to stir overnight. TLC separation using CH<sub>2</sub>Cl<sub>2</sub>—hexane (20:80, v/v) as eluent gave cluster **3** as the only major compound (34 mg, 30%).

Table 2 Crystal and refinement data for  $[Os_3(\mu-SbPh_2)_2(CO)_{10}]$  2a,  $[Os_3(\mu-H)(\mu-SbPh_2)(CO)_{10}]$ , 3 and  $[Os_3(\mu-SbPh_2)(\mu-PPh_2)(CO)_{10}]$  4

	2a	3	4
Formula	$C_{34}H_{20}O_{10}Os_3Sb_2\cdot\frac{1}{2}C_6H_{14}$	$C_{44}H_{22}O_{20}Os_6Sb_2 \cdot 1.5C_6H_{14}$	$C_{34}H_{20}O_{10}Os_3PSb\cdot\frac{1}{2}CH_2Cl_2$
M	1445.69	2384.57	1354.28
Crystal system	Monoclinic	Monoclinic	Monoclinic
Space group	$P2_1/c$	P2/c	$P2_1/c$
a/Å	13.3803(2)	14.8356(4)	12.6310(1)
b/Å	17.9495(2)	11.8116(3)	18.0529(2)
c/Å	17.6996(1)	19.4734(4)	17.1885(1)
β/°	98.097(1)	101.587(1)	94.469(1)
$V/Å^3$	4208.53(8)	3342.82(14)	3907.45(5)
Z	4	2	4
$\mu(\text{Mo-K}\alpha)/\text{mm}^{-1}$	10.342	12.213	10,569
Reflections collected	26494	19021	25323
Independent reflections	$10238 (R_{int} = 0.0533)$	$7128 (R_{int} = 0.0304)$	$9701 (R_{int} = 0.0420)$
Data/restraints/parameters	10238/0/341	7128/8/358	9701/1/453
Final R1, wR2 indices $[I > 2\sigma(I)]$	0.0576, 0.1212	0.0323, 0.0704	0.0462, 0.0857
(all data)	0.1125, 0.149	0.0506, 0.0808	0.0809, 0.0983

## Reaction of compound 1 and Ph<sub>2</sub>SbCl

An excess of Ph<sub>2</sub>SbCl (31 mg, 0.10 mmol) and compound 1 (60 mg, 0.0522 mmol) were stirred in THF at room temperature until 1 had been consumed (by IR spectroscopy, ≈1 d). Removal of the solvent in vacuo, followed by thin-layer chromatographic separation with CH<sub>2</sub>Cl<sub>2</sub>-hexane (20:80, v/v) as the eluent, gave **2a** as a yellow band (47.6 mg, 65%).

A similar procedure was used to prepare [Os<sub>3</sub>(μ-SbPh<sub>2</sub>)- $\{\mu\text{-Sb}(C_6H_4Me\text{-}p)_2\}(CO)_{10}],\ \ \textbf{2b},\ \ \text{in}\ \ 70\%\ \ \text{yield}\ \ (56\ \ \text{mg})\ \ \text{from}$ compound 1 (63 mg, 0.055 mmol) and an excess of (p-MeC<sub>6</sub>- $H_4)_2SbCl$  (33 mg, 0.097 mmol), and  $[Os_3(\mu-SbPh_2)(\mu-PPh_2) (CO)_{10}$ , 4, in 85% yield (49 mg) from 1 (50 mg, 0.044 mmol) and an excess of Ph<sub>2</sub>PCl (3–4 drops). **2b**: IR ( $\nu$ (CO), hexane) 2101w, 2054 mw, 2025 s, 2000 w, 1982 mw, 1971 vw and 1958 mw  $cm^{-1}$ (Calc. for C<sub>36</sub>H<sub>24</sub>O<sub>10</sub>Os<sub>3</sub>Sb<sub>2</sub>: C, 30.22; H, 1.69. Found: C, 30.53; H, 1.66%). **4**: IR ( $\nu$ (CO), hexane) 2104w, 2062mw, 2027s, 2000mw, 1992mw, 1967w and 1955m cm $^{-1}$ ;  $^{31}P-\{^{1}H\}$  NMR  $\delta$  46.44s (Calc. for C<sub>34</sub>H<sub>20</sub>O<sub>10</sub>Os<sub>3</sub>PSb: C, 31.12; H, 1.52. Found: C, 30.97; H, 1.98%).

### Crystal structure determinations

Crystal data and structure refinement details are given in Table 2. All organic hydrogen atoms were placed in calculated positions and allowed to ride on the attached carbon atoms; their isotropic thermal parameters were given a value 1.5 times that of the attached carbon. For compound 3 the metal hydride was placed in a calculated position at 1.76 Å from Os(1) and Os(2) and trans to Os(3); the isotropic thermal parameter was fixed at 0.08 Å<sup>2</sup> and the Os-H distances were also fixed at 1.76 Å. All non-hydrogen atoms (except those of the solvent molecules) were given anisotropic displacement parameters in the final refinement.

All three compounds showed the presence of solvent molecules in the crystals, confirmed by <sup>1</sup>H NMR. Compound **2a** has half a molecule of hexane which was modelled as disordered about an inversion centre; the carbon atoms were given isotropic thermal parameters. Compound 3 has one and a half molecule of hexane modelled as one half molecule and a full molecule with two disordered sites; the carbon atoms were given a fixed isotropic thermal parameter of 0.15 Å<sup>2</sup>. Appropriate restraints were placed on the geometries of the disordered solvent molecules. There was a half molecule of CH<sub>2</sub>Cl<sub>2</sub> in compound 4 which was refined isotropically; the thermal parameters for the chlorine atoms were constrained to be equal.

CCDC reference number 186/2208.

See http://www.rsc.org/suppdata/dt/b0/b006669i/ for crystallographic files in .cif format.

### Acknowledgements

This work was supported by the National University of Singapore (Research Grant No. RP 982751) and one of us (G. C.) thanks the University for a Research Scholarship. Technical assistance with the EXSY experiment, rendered by Ms Wong S. Y. and Ms Shannon Sng of the Crystal and Molecular Analysis Centre (NUS), is also gratefully acknowledged.

### References

- 1 For example, T. P. Fehlner, Inorganometallic Chemistry, Plenum, New York, 1992; W. A. Herrmann, Angew. Chem., Int. Ed. Engl., 1986, 25, 56; J. N. Nicholls, Polyhedron, 1984, 3, 1307.
- 2 K. H. Whitmire, Adv. Organomet. Chem., 1998, 42, 1.
- 3 W. K. Leong, Bull. Sing. N. I. C., 1996, 24, 51.
- 4 G. Chen and W. K. Leong, J. Chem. Soc., Dalton Trans., 1998,
- 5 G. Chen and W. K. Leong, J. Organomet. Chem., 1999, 574, 276.
- 6 B. F. G. Johnson, J. Lewis, A. J. Whitton and S. G. Bott, J. Organomet. Chem., 1990, 389, 129.
- 7 H. G. Ang, S. G. Ang and S. Du, J. Organomet. Chem., 1999, 590, 1.
- 8 A. J. Deeming and S. Hasso, J. Organomet. Chem., 1975, 88, C21; J. R. Shapley, J. B. Keister, M. R. Churchill and B. G. DeBoer, J. Am. Chem. Soc., 1975, 97, 4145.
- 9 C. K. Johnson, ORTEP II, Report ORNL-5138, Oak Ridge
- National Laboratory, Oak Ridge, TN, 1976.

  10 D. A. Bailey, A. L. Balch, L. A. Fossett, M. M. Olmstead and P. E. Reedy, Jr., Inorg. Chem., 1987, 26, 2413; M. H. Chisholm, K. Folting, J. C. Huffman and J. A. Klang, Organometallics, 1988, 7, 1033; G. S. Rodman and K. R. Mann, J. Organomet. Chem., 1989, 378, 255; M. Akita, M. Terada and Y. Morooka, Organometallics, 1992, 11, 1825; D. Cauzzi, C. Graiff, M. Lanfranchi, G. Predieri and A. Tiripicchio, J. Organomet. Chem., 1997, 536-7, 497; D. Cauzzi, C. Graiff, G. Predieri, A. Tiripicchio and C. Vignali, J. Chem. Soc., Dalton Trans., 1999, 237
- 11 A. J. Arce, P. Arrojo and Y. De Sanctis, Polyhedron, 1992, 9, 1013.
- 12 J. B. Keister, J. R. Shapley and D. A. Strickland, *Inorg. Synth.*, 1988,
- 13 M. Nunn, D. B. Sowerby and D. M. Wesolek, J. Organomet. Chem., 1983, 251, C45.