## JOM 24134PC

# **Preliminary Communication**

Synthesis and structure of Ir<sub>4</sub>Sn<sub>3</sub>(CO)<sub>6</sub>(CH(SiMe<sub>3</sub>)<sub>2</sub>)<sub>5</sub>O<sub>3</sub>; a heterometallic 'raft' with sail hoisted! \*

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#### **Abstract**

Heterometallic raft clusters have been obtained previously for a variety of metals but none for tin and iridium, and more significantly none to date have had metal groups bonded above the raft plane. We report a hexametallic Ir<sub>4</sub>Sn<sub>2</sub> raft to which a third tin group is attached by a single short Ir-Sn bond and three Ir-O-Sn bridges.

There is much current interest in the interface between Main Group and Transition Metal cluster chemistry [1]. By employing the low valent tin-reagent  $[SnR_2^1]$   $[R^1 = CH(SiMe_3)_2]$  we sought to prepare the first tin-iridium clusters. This tin reagent, first prepared and characterised by Lappert and his co-workers [2], proved valuable in the preparation of mixed metal cluster of iron, ruthenium, and osmium having a variety of frameworks [3].

We have now studied the reactions of several tin(II) reagents  $SnR_2$  (including  $R = R^1$ , or 2,4,6-triisopropylphenyl) with  $[Ir_4(CO)_{11}Br]^-$ , a labile derivative of the insoluble and rather unreactive parent  $[Ir_4(CO)_{12}]$ . We have isolated and characterised a variety of Ir-Sn clusters with nuclearities varying from 5 to 9 [4].

With strict exclusion of air and moisture [NEt<sub>4</sub>][Ir<sub>4</sub>-(CO)<sub>11</sub>Br] was suspended in degassed Et<sub>2</sub>O and a solution of [SnR<sup>1</sup><sub>2</sub>] in Et<sub>2</sub>O was added. The mixture

was allowed to warm to ambient temperature, stirred for 8 h, filtered to remove small amounts of  $[Ir_4(CO)_{12}]$ , and concentrated to ca. 30 ml; TLC of the product (silica; eluant hexane) showed the presence of four compounds with  $R_{\rm f}$  values of 0.75, 0.65, 0.6 and 0.4. Solvent was removed under vacuum, the residue redissolved in the minimum of hexane and placed on a flash silica column under pure hexane. The first three products were collected with hexane as eluant, and gradual increase of the polarity (to ca. 80:20 hexane: dichloromethane) afforded the slowest moving species as a green solution. Evaporation of the solvent, followed by crystallisation of the residue from hexane/toluene, afforded the dark green-black crystals of the fourth product (variable yield 15-35%), suitable for an X-ray diffraction study. The samples gave a satisfactory elemental analysis; IR (KBr)  $\nu$ (CO) 2064m, 2031s, 2020sh, 2008m, 1998w, 1982m, and 1976sh cm<sup>-1</sup>. Suitable crystals of approximate dimensions  $0.25 \times 0.15 \times 0.15$  mm were mounted on an Enraf-Nonius CAD4 diffractometer, and data were taken from 2 separate crystals owing to substantial decomposition occurring during data collection. Crystals became black and opaque following exposure, probably associated with metal-metal bond fission during X-ray exposure.

The three products which were eluted first were identified as  $Ir_3$  and  $Ir_4$  clusters having bridging  $SnR^1_2$  groups [4]. Of particular interest is the fourth product, obtained as a dark green cluster of stoichiometry  $[Ir_4Sn_3(CO)_6R^1_5O_3]$ . The yield of this species is variable, but the highest yields were when a substantial excess of  $[SnR^1_2]$  was used. Figure 1 shows the molecular structure of the molecule, which can be described as a hexametallic  $Ir_4Sn_2$  planar 'raft' [5], surmounted by a  $\mu_4$ -R¹SnO<sub>3</sub> 'sail'. Although molecular rafts of 6 and more metal atoms are known [5] none has been reported to date with a capping group.

The geometry about the capping tin of the sail is that of a highly distorted trigonal bipyramid, with normal Sn-O distances, though it is noteworthy that the Ir-Sn distance for the sail is the shortest such separation in the cluster. It may be mentioned that structures involving Group 14 element-O-Os bonds have been postulated in deposits of osmium clusters on silica [6]. The incorporation of oxygen atoms in the reactions we describe is unique, though it is probable that they come from the silica used in chromatography. Thus, the

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<sup>\*</sup> Dedicated to Professor M.F. Lappert on the occasion of his 65th birthday, with our warmest good wishes for the future.

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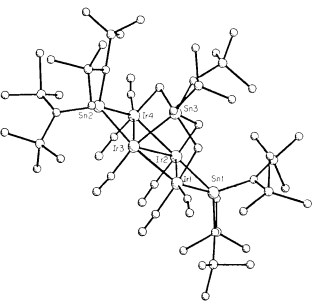


Fig. 1. A single molecule of the cluster. Selected bond lengths (Å) and angles (°): Sn(3)-Ir(3) 2.637(6), Sn(2)-Ir(3) 2.676(6), Sn(2)-Ir(4) 2.680(6), Sn(3)-O 2.008(26), 2.192(24), and 2.194(25), (to the three O-atoms); Ir-O distances 2.120(26), 2.134(25), and 2.130(26); Ir(3)-Sn(3)-C(11) 150.1(11).

product from chromatography is not the dark green species initially obtained from the reaction (IR data) [7], and reaction with  $[SnAr_2]$  (Ar = 2,4,6-triisopropylphenyl) gives a similar product distribution, in which the analogous green compound observed decomposes on a silica column, precluding isolation. There is precedent for a cluster having a  $SnR^1$  ligand [8], formed by an unusual rearrangement reaction, and having a unique framework geometry, but in that case, the bonding to the transition metal framework is by two Sn-Os bonds.

A single cluster has previously been reported bearing an SnR<sup>1</sup> substituent, also formed by rearrangement and having an unusual framework geometry [7\*].

Crystal data:  $C_{41}H_{95}Ir_4O_9Si_{10}Sn_3 \cdot C_7H_8$ , M=2230.23, space group  $P\overline{1}$ . Triclinic, a=12.095(9), b=15.511(11), c=22.51(2) Å,  $\alpha=82.48(5)$ ,  $\beta=87.86(5)$ ,  $\gamma=75.56(6)^\circ$ ; U=4054(5) Å<sup>3</sup>. Z=2;  $D_{calc}=1.83$  g

cm<sup>-3</sup>. A total of 8307 reflections were measured to a maximum  $\theta$  of 20°, using Mo-K $\alpha$  radiation, with a graphite monochromator on a CAD-4 diffractometer  $(\lambda = 0.71069 \text{ Å})$ . Data were corrected for both decay and absorption, using a linear decay correction, and empirical absorption correction (merging R = 6.05%). A second crystal was mounted when the first had decayed to 60% of the initial intensity. Merging of the two datasets gave 6695 independent reflections of which 5211 were observed using the criterion  $|F_{\alpha}| > 4\sigma(F_{\alpha})$ . The structure was solved using SHELXS (courtesy of Professor G.M. Sheldrick, University of Göttingen), and refined using full-matrix least squares (SHELX-76) to a conventional R-factor of 8.8%. The moderate quality of the data is due to serious absorption and decay problems. An absorption correction using  $\psi$ -scan data was attempted, but not used, as it made no detectable improvement in the fit to the data. The data have been deposited with the Cambridge Crystallographic Data Centre.

#### Acknowledgement

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## References and notes

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<sup>\*</sup> Reference number with asterisk indicates a note in the list of references.