## **Fluxional Ligand Migrations in Triosmium Clusters containing 2-Pyridylphosphines**

## **Antony J. Deeming\* and Martin 6. Smith**

*Department of Chemistry, University College London, 20 Gordon Street, London, UK WClH OAJ* 

The cluster  $[Os_3(CO)_{10}(\mu-L)]$  (L = Ph<sub>2</sub>PC<sub>5</sub>H<sub>4</sub>N) (X-ray structure reported), containing the diaxially coordinated 2-pyridyldiphenylphosphine bridge, undergoes rapid transfer of the pyridine between two metal atoms while the phosphorus atom remains firmly coordinated to the third.

The ligand 2-pyridyldiphenylphosphine ( $Ph_2PC_5H_4N$ , L) has been widely used to bridge metal centres and an extensive coordination and organometallic chemistry based on it has been developed.<sup>1</sup> It is generally reckoned to stabilize dinuclear compounds by being firmly attached to the metal atoms it bridges. We have now shown that the ligand L reacts with  $[Os<sub>3</sub>(CO)<sub>10</sub>(MeCN)<sub>2</sub>]$  in dichloromethane solution at room temperature to give orange crystals of the cluster  $[Os<sub>3</sub>(CO)<sub>10</sub>(\mu-L)]$  **1** (45%) and yellow crystals of  $[Os<sub>3</sub>(CO)<sub>11</sub>L]$  2 (21%) which were separated by TLC on silica from other products including  $[Os<sub>3</sub>(CO)<sub>10</sub>L<sub>2</sub>]$ . The <sup>1</sup>H NMR spectrum? shows that **2** contains the ligand L coordinated

through the phosphorus atom only [6-H pyridyl resonance at 6 **8.87** compared with 6 **8.72** for the free ligand] and its IR spectrum<sup>†</sup> is very similar to that of  $[Os<sub>3</sub>(CO)<sub>11</sub>(PPh<sub>3</sub>)]$ indicating that they are isostructural.2

The cluster **1** is very interesting because of its crystal structure and fluxional behaviour in solution. Its single-crystal X-ray structure\$ (Fig. **1)** establishes that the N and P atoms are both coordinated axially, consistent with the non-equivalence of the Ph groups as observed in the <sup>1</sup>H NMR spectra<sup>†</sup> between  $-20$  and  $-90$  °C, a result that is inconsistent with diequatorial coordination. This stereochemistry contrasts with dppm)] (dppm = Ph<sub>2</sub>PCH<sub>2</sub>PPh<sub>2</sub>), *e.g.* in which bridging dppm ligands are diequatorially coordinated.<sup>3,4</sup> Normally ligands bulkier than hydride<sup>5</sup> or acetontirile<sup>6</sup> are equatorially coordinated and we believe that the small span of L ( $N \cdots P$  = 2.650 Å)<sup>7</sup> compared with that of dppm  $(P \cdots P = 2.970$  Å)<sup>8</sup> favours diaxial coordination. The N..-P distance in **1** is 2.716 Å whereas the P···P distance in  $\left[\text{Ru}_3(\text{CO})_{10}(\mu\text{-dppm})\right]$  is **3.127 A.3** These distances might be compared with the that of  $[Ru_3(CO)_{10}(\mu\text{-dppm})]$  and  $[Os_3(CO)_{9}(\mu\text{-dppm})(\eta^{1}$ -

t *Spectroscopic data* for compound **1:** IR v(CO)/cm-l (cyclohexane): 2083s, 2032vs, 2017vs, 2004s, 1977s, 1969s, 1955m, 1938w; <sup>1</sup>H NMR (400 MHz, CD<sub>2</sub>Cl<sub>2</sub>, -60 °C): 2-pyridyl:  $\delta$  9.42 (6-H), 7.55 (4-H), 7.30 (5-H), 7.25 (3-H); phenyl: **6** 7.50, 7.45, 7.09; (1,1,2,2-C2D2C14, 80°C): 2-pyridyl: 9.41 (6-H), 7.51 (4-H), 7.22 (5-H, 3-H); phenyl: 6 7.45, 7.33 (signals at  $\delta$  7.61 and 7.09 at 20 °C have coalesced);  ${}^{31}P$ { ${}^{1}H$ } NMR (162 MHz,  $CD_2Cl_2$ , 20 °C),  $\delta$  5.6 (s).

Compound **2,** IR v(CO)/cm-l (cyclohexane): 2108m, 2053s, 2042m, 2020vs, 1999m, 1987m, 1971m; <sup>1</sup>H NMR (400 MHz, CDCl<sub>3</sub>, 20°C): 2-pyridyl: 6 8.87 (6-H), 7.57 (5-H), 7.27 (4-H), 7.09 (3-H); phenyl: δ 7.40-7.50; <sup>31</sup>P{<sup>1</sup>H} NMR (162 MHz, CDCl<sub>3</sub>, 20 °C): 4.2(s).

Compound  $3: \text{IR } v(\text{CO})/\text{cm}^{-1}$  (cyclohexane): 2083s, 2034vs, 2018vs, 2003m, 1977s, 1968s, 1952w, 1939w; <sup>1</sup>H NMR (400 MHz, CD<sub>2</sub>Cl<sub>2</sub>, -20 °C): N-bonded 2-pyridyl:  $\delta$  9.40 (6-H), 7.91 (3-H), 7.59 (4-H), 7.30 (5-H), N-non-bonded 2-pyridyl: 6 8.70, 8.59 (6-H), 8.03, 7.88 (5-H), 7.74, 7.39, 7.29, 7.19 (3-H, 4-H); 31P{1H} NMR  $(162 \text{ MHz}, \text{CD}_2\text{Cl}_2, 20 \text{ °C})$ :  $\delta$  12.7(s).

Compound 4: IR v(CO)/cm<sup>-1</sup> (cyclohexane): 2083s, 2033vs, 2018vs, 2003m, 1978s, 1969s, 1952w, 1938w; <sup>1</sup>H NMR (400 MHz,  $CD_2Cl_2$ ,  $20 °C$ ): two isomers: N-bonded 2-pyridyl:  $\delta$  9.49 and 9.37 (6-H); N-non-bonded 2-pyridyl: 6 8.72 and 8.67 **(6-H);** other resonances at 6 8.12, 8.03, 7.86, 7.81, 7.62, 7.51, 7.41, 7.35, 7.27, 7.19, 7.01; 31P{1H} NMR (162 MHz, CD2C12, 20°C): 6 11.0 **(s)** and 8.4 **(s)** *(ca.* 1 : 1 ratio).

<sup>\$</sup> *Structure determination:* Nicolet R3v/m diffractometer, Mo-radiation  $(\lambda = 0.71073 \text{ Å})$ , room temp. data corrected empirically for absorption. Orange crystal,  $C_{27}H_{14}NO_{10}POs_3$ ,  $M = 1113.99$  g mol<sup>-1</sup>, size =  $0.40 \times 0.35 \times 0.30$  mm<sup>3</sup>, monoclinic,  $P2_1/n$ ,  $a = 9.278(4)$ ,  $b =$  $25.508(7)$ ,  $c = 13.096(6)$  Å,  $\beta = 107.31(4)$ °,  $U = 2959(2)$  Å<sup>3</sup>,  $Z = 4$ ,  $D_c = 2.50$  g cm<sup>-3</sup>,  $\mu$ (Mo-K $\alpha$ ) = 129.7 cm<sup>-1</sup>,  $F(000) = 2024$ . Direct methods,  $R = 0.0556$ ,  $R' = 0.0545$ , where  $R' = [\Sigma w(|F_o] |F_c|$ <sup>2</sup>/ $\sum$  $|F_o|^2$ <sup> $\}$ </sup>, after refinement 379 parameters using 4107 data with  $I_o$  $> 1.5\sigma(I_0)$  in the range  $5 \le 2\theta \le 50^\circ$ . All non-H atoms were refined anisotropically with H-atoms included in idealised positions (C-H 0.96 Å,  $U = 0.08$  Å<sup>3</sup>). Atomic coordinates, bond lengths and angles, and thermal parameters for both structures have been deposited at the Cambridge Crystallographic Data Centre. See Notice to Authors, Issue No. 1.



Fig. 1 Molecular structure of  $[Os<sub>3</sub>(CO)<sub>10</sub>(\mu-Ph<sub>2</sub>PC<sub>5</sub>H<sub>4</sub>N)]$ , cluster 1.  $(CO)<sub>3</sub>CO$ Selected bond lengths (Å) and angles (°): Os(1)-Os(2), 2.827(2); Os(2)-Os(3), 2.872(2); Os(1)-Os(3), 2.929(2); Os(1)-P, 2.380(4); Os(2)-N, 2.22(1); P-C(41), 1.85(2); N-C(41), 1.32(2); Os(1)-C(11), 1.92(2);  $Os(2)-C(21)$ , 1.87(2);  $Os(3)-C(31)$ , 2.01(2);  $Os(3)-C(34)$ ,  $Os(2)-Os(1)-P, 81.9(1); Os(1)-P-C(41), 117.3(5); Os(2)-N-C(41),$  $125(1)$ ; N-C(41)-P, 117(1). 1.94(2); av. Os-C (equatorial), 1.894; Os(1)-Os(2)-N, 93.2(4);



Fig. 2 400 MHz <sup>1</sup>H NMR spectra of the cluster  $[Os<sub>3</sub>(CO)<sub>10</sub>{\mu}$ - $P(C_5H_4N)_3$ } recorded in CD<sub>2</sub>Cl<sub>2</sub> (-20 and 20 °C) or in 1,1,2,2- $C_2D_2Cl_4$  (80 °C). Signals for the 6-H protons of the pyridyl rings are indicated for the free and coordinated groups; \* indicates peaks for decomposition products at 80 "C.

shortest axial-axial C...C distances (av = 2.875 Å) and the shortest equatorial-equatorial distances (av  $= 3.424 \text{ Å}$ ) in  $[Os<sub>3</sub>(CO)<sub>12</sub>]$  (calculated from data given in ref. 5). Dppm is a more flexible ligand and has been found to span metal-metal distances from 2.1 to 4.4 Å,<sup>9</sup> while  $Ph_2PC_5H_4N$  has never been found associated with a metal-metal distance greater than about 2.9 A.

The <sup>1</sup>H and <sup>13</sup>C $\{$ <sup>1</sup>H $\}$  NMR spectra† of **1** are temperature dependent. At 20 °C sharp 2-pyridyl<sup>1</sup>H NMR resonances are observed while the phenyl resonances are very broad (linewidths up to 30Hz). Lower temperatures lead to better resolution whereas higher temperatures lead to coalescence of peaks at  $\delta$  7.61 and 7.09 to a new broad resonance at  $\delta$  7.33. We have not been able to assign these phenyl signals with confidence but the observed changes are consistent with dynamic exchange of the two non-equivalent phenyl groups leading to a time-averaged plane of symmetry.



**Scheme 1** Possible mechanisms for  $R^a-R^b$  exchange in compounds 1, 3 and **4** 

The corresponding tri(2-pyridy1)phosphine complex  $[Os<sub>3</sub>(CO)<sub>10</sub>{\mu \cdot \hat{P}(C_5H_4N)_3}]$  3, prepared similarly, gives more easily interpreted <sup>1</sup>H NMR spectra. At  $-20$  °C there are three sharp sets of 2-pyridyl resonances [6-H resonances: 6 8.70 and 8.59 for the non-coordinated groups and 6 **9.40** for the coordinated group] (Fig. 2). The pairwise coalescence of the non-coordinated 2-pyridyl resonances without the coordinated 2-pyridyl resonances changing very much shows that the exchange process does not involve the formation of monodentate P-bonded tri(2-pyridy1)phosphine which would lead to the mutual exchange of all three groups. Therefore we propose that Ra exchanges with Rb while the N and **P** atoms remain coordinated (Scheme 1). Processes **A,** B and C could account for the coalescences but we feel that process C is less likely than the others. Process A corresponds to a classical CO rearrangement process but with the pyridine and a CO ligand migrating *via* bridging positions instead of two CO ligands. However, a phosphine migration is an alternative and has been observed in triplatinum systems.10

We examined 13C NMR spectra of cluster **1,** hoping to discriminate between these processes. **A** set of ten 13C carbonyl resonances consistent with the static structure is observed for a  $CD_2Cl_2$  solution at  $-85$  °C but there are fast processes leading to CO exchange which prevent our specifically identifying the CO exchanges corresponding to  $R^a-R^b$ exchange. At room temperature nine of the ten resonances have coalesced to a broad resonance at  $\delta$  183.5 while that for the CO ligand *trans* to the PPh<sub>2</sub> group remains as a doublet  $[\delta 184.2 \, (\bar{d}, J = 4.5 \, \text{Hz})]$ . We suggest that process A together with other processes including a merry-go-round process in the  $M_3$  plane over all three metal atoms as identified for [ $M_3(CO)_{10}(\mu$ -1,2-diazine)] (M = Ru or Os),<sup>11,12</sup> also containing diaxially coordinated ligands, would lead to just the one CO ligand not exchanging. Process B or C could not account for this observation. Final confirmation that the P-atom remains bonded to the same 0s atom throughout came from the observation of sharp <sup>187</sup>Os satellites<sup>13,14</sup> to the <sup>31</sup>P $\{^1H\}$ NMR singlet  $[J(187Os<sup>31</sup>P)$  165 Hz] at temperatures from -40 °C up to 50 °C. Any process involving the migration of the P-atom between the 0s sites would have led to the collapse of the 1870s satellites. Only process A, or the very unlikely process C, would be consistent with the observed  $31P\{1H\}$ NMR spectra.

The corresponding cluster  $[Os<sub>3</sub>(CO)<sub>10</sub>(\mu-PhP(C<sub>5</sub>H<sub>4</sub>N)<sub>2</sub>)]$  4 from **di(2-pyridy1)phenylphosphine** shows two isomers both in

the 1H and **31P** NMR spectra as expected but a rapid equilibrium between them at room temperature leads to broad resonances. This is completely consistent with what we have concluded for the clusters **1** and **3.** 

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## **References**

1 C. G. Arena, E. Rotondo, **F.** Faraone, M. Lanfranchi and A. Tiripicchio, *Organometallics,* 1991, 10, 3877; **S.** Lo Schiavo, E. Rotondo, G. Bruno and F. Faraone, *Organometallics,* 1991, 10, 1613; E. Rotondo, **S.** Lo Schiavo, G. Bruno, C. G. Arena, R. Gobetto and F. Faraone, *Inorg. Chem.,* 1989, 28, 2944; J. L. Roustan, N. Ansari and F. R. Ahmed, *Inorg. Chem. Acta,* 1987, 129, L11; Z.-Z. Zhang, H.-K. Wang, H.-G. Wang and R.-J. Wang, J. *Organomet. Chem.* , 1986, 314, 357; T. **J.** Barder, F. A. Cotton, G. L. Powell, **S.** M. Tetrick and R. A. Walton, J. *Am. Chem. SOC.,* 1984, **106,** 1323; J. P. Farr, M. M. Olmstead, N. M. Rutherford, **F.** E. Wood and A. L. Balch, *Organometallics,* 1983, 2,1758; **J.** P. Farr, M. M. Olmstead, **F.** E. Woodand A. L. Balch, *J. Am. Chem. SOC.,* 1983, 105,792.

- 2 M. **1.** Bruce, D. C. Kehoe, J. G. Matisons, B. K. Nicholson, P. H. Rieger and M. L. Williams, *J. Chem. SOC., Chem. Commun.,*  1982, 442; B. **F.** G. Johnson, J. Lewis and D. **A.** Pippard, *J. Chem. SOC., Dalton Trans.,* 1981, 407.
- 3 A. W. Coleman, D. **F.** Jones, P. H. Dixneuf, C. Brisson, J.-J. Bonnet and G. Lavigne, *Inorg. Chem.,* 1984,23,952.
- **4 S.** Cartwright, J. **A.** Clucas, R. H. Dawson, D. **F.** Foster, M. M. Harding and **A.** K. Smith, *J. Organomet. Chem.,* 1986, 302,403.
- *5* M. R. Churchill and B. G. DeBoer, *Inorg. Chem.,* 1977, 16, 878. 6 P. **A.** Dawson, B. F. G. Johnson, J. Lewis, J. Puga, P. R. Raithby
- and M. J. Rosales, J. *Chem. SOC., Dalton Trans.,* 1982, 233. 7 Calculated from data given by J.-P. Charland, J.-L. Roustan and N. Ansari, *Acta Crystallogr., Sect. C,* 1989, 45, 680.
- 8 Calculated from data given by H. Schmidbaur, G. Reber, A. Schier, F. E. Wagner and G. Miiller, *Inorg. Chim. Acta,* 1988, 147, 143.
- 9 A. L. Balch, R. R. Guimerans, J. Linehan and F. E. Wood, *Inorg. Chem.,* 1985, 24, 2021.
- 10 A. M. Bradford, G. Douglas, L. Manojlović-Muir, K. W. Muir and R. J. Puddephatt, *Organometallics,* 1990, 9, 409.
- 11 F. A. Cotton, B. E. Hanson and J. D. Jamerson, *J. Am. Chem.*  **SOC.** , 1977, 99, 6588.
- 12 F. A. Cotton and B. E. Hanson, *Inorg. Chem.,* 1977, 16,2820.
- 13 M. A. Gallop, B. F. G. Johnson and **J.** Lewis, *J. Chem.* Soc., *Chem. Commun.,* 1987, 1831.
- 14 R. Benn, H. Brenneke, E. Joussen, H. Lehmkuhl and F. López Ortiz, *Organometallics,* 1990, 9, 756.