## **Cluster Chemistry. Reactions between Metal Carbonyl Clusters and Lewis Bases Initiated by Radical Ions: Improved Syntheses of Substituted Derivatives of M, and M, Clusters (M** = **Fe, Ru, Os, or Co)**

**Michael 1. Bruce,\*a David C. Kehoe,a Janis G. Matisons,a Brian K. Nicholson,a Philip H. Rieger,b and Michael L. Williamsa** 

**<sup>a</sup>***Department of Physical and Inorganic Chemistry, University of Adelaide, Adelaide, South Australia 5007*  **<sup>b</sup>***Department of Chemistry, Brown University, Providence, Rhode Island 029 72, USA.* 

Addition of catalytic amounts of  $Ph<sub>2</sub>CO $\cdot$  to a mixture of the appropriate cluster with stoichiometric amounts$ of Lewis base in tetrahydrofuran gave the substituted derivatives  $[Fe_3(CO)_{11}(L)]$ ,  $[Ru_3(CO)_{12-n}(L)_n]$ ,  $[Os_3(CO)_{11}(L)]$ , and  $[H_4Ru_4(CO)_{12-n}(L)_n]$   $[n = 1-3; L = Bu^tNC, PR_3, P(OR)_3]$  in greatly improved yields.

**A** continuing hindrance to the study of metal carbonyl clusters, including their reactions with simple donor molecules, is the severe reaction conditions required to bring about substitution. These often result in further transformations of first-formed reaction products, often with concomitant bond cleavage in the reactant molecules, in further substitution of an activated monosubstituted product, as found in reactions between  $\text{[Ru}_{3}(\text{CO})_{12}\text{]}$  and many tertiary phosphines or arsines, or in the formation of all possible substitution products, as with  $[H_4Ru_4(CO)_{12}]$  and tertiary phosphines or phosphites, which require extensive chromatography to isolate low yields of products. We describe below a novel method of introducing a variety of 2e donor ligands into metal cluster complexes in specific, stepwise reactions which generally proceed rapidly and in high yields.

Recent studies have shown that electrochemically or chemically generated  $[Co_2(CO)_6(\mu-C_2R_2)]$  or  $[Co_3(\mu_3-CR) (CO)_9$ <sup>--</sup> radical anions react extremely rapidly with tertiary phosphines or phosphites to give substituted complexes.<sup>1</sup> We have now found that this reaction can be extended to a wide range of polynuclear metal carbonyl clusters that are not normally labile towards substitution.

Typically, a mixture of the cluster carbonyl and the stoich-

eiometrically required amount of ligand are dissolved in tetrahydrofuran (THF), and the mixture is treated dropwise with a solution of sodium diphenylketyl in **THF** *(ca.* 0.025 **M)** until reaction is complete (i.r. confirmation). For the more reactive systems (see below),  $2-4$  drops of catalyst solution ( $\leq 0.1$  mol  $\%$ ) were sufficient, the reactions being completed within five minutes at ambient temperature. Since the reactions are essentially specific and quantitative, work-up involves only a simple extraction-crystallisation sequence, thus avoiding tedious chromatographic separations. Table 1 summarises some typical reactions and products, and can be amplified by the following comments concerning particular cluster systems :

(i)  $[Fe<sub>3</sub>(CO)<sub>12</sub>]$  required somewhat more catalyst than indicated above, and some fragmentation of the Fe, cluster was observed, with concomitant lowering of yields of the substituted cluster. Nevertheless, yields were significantly higher than those obtained by a much more laborious thermal route.<sup>2</sup>

(ii)  $[Ru_3(CO)_{12}]$  has been the most studied cluster, and reacts quickly to give excellent yields of mono-, di-, tri-, or tetra- (in one case) substituted products. The thermal route often gives only the trisubstituted complex,  $\text{[Ru}_{3}(\text{CO})_{9}(\text{L})_{3}$ ].<sup>3</sup>

(iii)  $[H_4Ru_4(CO)_{12}]$  also reacts readily to give the appropri-



**Table 1.** Products obtained from radical ion-initiated reactions between metal carbonyl clusters and Lewis bases.

**a** All new compounds have satisfactory elemental microanalyses, i.r., and <sup>1</sup>H n.m.r. spectra; dppe  $= Ph_2PCH_2CH_2PPh_2$ , dpam  $= CH_2$ -(AsPh<sub>2</sub>)<sub>2</sub>. <sup>b</sup> Reference 2; Otol = OC<sub>6</sub>H<sub>4</sub>Me-p. <sup>c</sup> M. I. Bruce, D. Schultz, R. C. Wallis, and A. D. Redhouse, J. Organomet. Chem., 1979, 169, C15. <sup>d</sup> Reference 3. <sup>e</sup> F. Piacenti, M. Bianchi, P. Frediani, and E. Ben *Soc., Dalton Trans.,* 1974, 2549. Reference 2; Otol  $=$  OC<sub>6</sub>H<sub>4</sub>Me-*p*.

ate stoicheiometric product, in contrast with the thermal reactions which often give all possible derivatives  $[H_4Ru_4 (CO)_{12-n}(L)_n$   $(n = 0-4)$ .

(iv)  $[Os_3(CO)_{12}]$  and  $[H_4Os_4(CO)_{12}]$  are significantly less reactive than the analogous ruthenium complexes, and tend to give mixtures of products; however, the lower temperature required gives improved yields of primary substitution products, and little or no decomposition is found.

(v)  $[Co_3(\mu_3-CCl)(CO)_9]$  reacts quickly and cleanly<sup>1</sup> to give the stoicheiometric product, with up to three ligands being incorporated per cluster.

(vi)  $[M_4(CO)_{12}]$  (M = Co, Rh, or Ir) readily undergoes substitution by tertiary phosphines or phosphites, and the present method offers no advantage.

(vii)  $[H_3M_3(CO)_{12}]$  (M = Mn or Re) does not give substituted clusters; the manganese complex readily forms  $[Mn_2(CO)_{10}]$  and  $[Mn_2(CO)_8(L)_2]$  under the reaction conditions, while the rhenium complex undergoes a slow, nonspecific reaction.

This radical-catalysed reaction described above appears to be general for systems which satisfy the following requirements: (i) the cluster carbonyl can be reduced to an anion which has a significant lifetime without fragmentation occurring; (ii) the incoming ligand is  $\text{less } \pi$ -acidic than CO (to facilitate electron transfer from the substituted to the unsubstituted cluster radical anion); (iii) the ligand is not reduced by  $Ph_2CO^{-1}$ .

The mechanism proposed for these reactions is essentially that discussed earlier, $<sup>1</sup>$  and is related to the electron-transfer</sup> chain-catalysed reactions found in organic systems **;4** as an example, equations (l), **(2),** and **(3)** give the chain reaction for  $[Ru_3(CO)_{12}]$ . The reactive species is the radical anion

$$
[Ru_3(CO)_{12}] + Ph_2CO^{\bullet-} \longrightarrow [Ru_3(CO)_{12}]^{\bullet-} + Ph_2CO \qquad (1)
$$

$$
[Ru_3(CO)_{12}]^-+L \longrightarrow [Ru_3(CO)_{11}L]^{--}+CO \qquad (2)
$$

$$
[Ru_3(CO)_{11}L]^{\centerdot -} + [Ru_3(CO)_{12}] \longrightarrow [Ru_3(CO)_{11}L] + [Ru_3(CO)_{12}]^{\centerdot -} (3)
$$

 $[Ru_3(CO)_{12}]$ <sup>-</sup> where the excess electron is in an Ru-Ru  $(\sigma^*)$ antibonding orbital, which leads to a weakened bond. Cleavage of an Ru-Ru bond gives a 17e Ru centre which readily undergoes attack by L and elimination of CO (equation **2).**  Reformation of the Ru-Ru bond gives  $[Ru_3(CO)_{11}L]^{\text{-}}$  which rapidly transfers an electron to  $[Ru_3(CO)_{12}]$  to continue the catalytic cycle (equation **3).** 

**As** can be seen from Table **1,** this method affords many complexes which cannot be obtained from the analogous thermal reactions. For systems where the reaction is favourable, the short reaction times, mild conditions, and easy product isolation will all facilitate further study of the initial reactions occurring between ligands and metal carbonyl clusters.

We thank the Australian Research Grants Committee for support of this work, and Waikato University, Hamilton, New Zealand for study leave (B.K.N.).

*Received, 29th December 1981* ; *Corn. <sup>1470</sup>*

## **References**

- **1 G. J. Bezems, P. H. Rieger, and S. J. Visco,** *J. Chem. Suc., Chem. Commun.,* **1981, 265;** *J. Am. Chem. SOC.,* **submitted for publication; M. Arewgoda, P. H. Rieger, B. H. Robinson, J.**  Simpson, and S. J. Visco, *J. Am. Chem. Soc.*, submitted for **publication.**
- **2 S. M. Grant and A. R. Manning,** *Inorg. Chim. Acta,* **1978, 31, <sup>41</sup>**; **R. J. Angelici and E. E. Siefert,** *Inorg. Chem.,* **1966,5, 1457.**
- *3* **M. I. Bruce, G. Shaw, and F. G. A. Stone,** *J. Chem. Suc., Dalton Trans.,* **1972, 2094.**
- **4 R. W. Alder,** *J. Chem. SOC., Chem. Commun.,* **1980,** *1* **184.**