# **Reactions of Aryl Isocyanates with Hydridotriosmium Carbonyl Cluster Compounds**

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The reactions of aryl isocyanates with the cluster complexes H<sub>2</sub>Os<sub>3</sub>(CO)<sub>9</sub>L [I, L = CO; II, L = P(CH<sub>3</sub>)<sub>2</sub>C<sub>6</sub>H<sub>5</sub>] are described. The major products of the reaction of I with aryl isocyanates are complexes containing N-arylformamido ligands formed by the transfer of one hydride ligand to the carbon atom of the isocyanate molecule. An X-ray crystallographic analysis of the product  $(\mu-H)(\mu_P\text{-}CH_3C_6H_4NCHO)O_{3}(CO)_{10}$  (III) was performed: space group *PI*,  $a = 7.925$  (2)  $\AA$ ,  $b = 11.705$ (3) A,  $c = 13.928$  (4) A,  $\alpha = 66.71$  (2)<sup>o</sup>,  $\beta = 80.17$  (2)<sup>o</sup>,  $\gamma = 79.18$  (2)<sup>o</sup>,  $Z = 2$ ,  $\rho_{\text{cal}} = 2.83$  g/cm<sup>3</sup>. For 3902 reflections,  $R = 0.052$  and  $R_w = 0.062$ . III contains an N-p-tolylformamido ligand which is coordinated by its nitrogen and oxygen atoms across one edge of a triangular cluster of three osmium atoms. Reaction of I11 with dimethylphenylphosphine readily gives the monosubstitution product  $(\mu-H)(\mu-\rho-H_1C_1H_4NCHO)Os_3(CO)_9[P(CH_3)_2C_6H_5]$  (V). An X-ray crystallographic analysis of **V** was performed: space group  $P\bar{I}$ ,  $a = 10.690$  (1)  $\bar{A}$ ,  $b = 13.034$  (2)  $\bar{A}$ ,  $c = 13.200$  (4)  $\bar{A}$ ,  $\alpha = 109.78$  (2)°,  $\beta = 93.05 \ (2)^{\circ}, \ \gamma = 110.25 \ (2)^{\circ}, \ Z = 2, \ \rho_{\text{calod}} = 2.28 \ g/cm^3$ . For 3985 reflections,  $R = 0.047$  and  $R_w = 0.061$ . V is similar to I11 except that it contains a dimethylphenylphosphine ligand coordinated in an equatorial position on the osmium atom coordinated to the oxygen atom of the bridging  $N-p$ -tolylformamido ligand. The major product of the reaction of II with p-tolyl isocyanate was an isomer of V which was characterized by spectroscopic and X-ray crystallographic methods as  $(\mu$ -H $)(\mu$ - $p$ -CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NHCO $)$ Os<sub>3</sub>(CO)<sub>9</sub>[P(CH<sub>3</sub>)<sub>2</sub>C<sub>6</sub>H<sub>5</sub>] (VI): space group  $P_{1}/c$ ,  $a = 12.210$  (2) Å,  $b = 14.337$  (4) Å,  $c = 17.739$  (4) Å,  $\beta = 104.49$  (2)°. For reflections,  $R = 0.034$  and  $R_w = 0.032$ . VI ligand which **is** coordinated by its carbon and oxygen atoms across an edge of a triangular cluster of osmium atoms. This product was evidently formed by transfer of a hydrogen atom from the cluster to the nitrogen atom of the isocyanate. A minor product  $(\mu-H)(\mu-N-p\text{-CH}_3C_6H_4NHCO)O_{31}(CO)_{10}$  (VII), which is similar to VI, was also isolated from the reaction of I with p-tolyl isocyanate. It was shown that **111** and VI1 cannot be interconverted under the reaction conditions. They are, thus, presumed to be formed by independent competing reactions. The changes in hydrogen transfer obtained when a carbonyl ligand is replaced with a dimethylphenylphosphine ligand are discussed.

# **Introduction**

The transfer of hydrogen atoms from metal atoms to substrates is an integral component of any catalytic cycle involving metal-hydride intermediates.' Recently transition-metal cluster compounds have attracted considerable attention as a new source of homogeneous catalysts. $2$  In an effort to delineate the nature of hydride transfers from cluster compounds to substrates, we have been investigating the reactivity of the hydridotriosmium carbonyl clusters  $H_2O_{3}(CO)_9L$  [I, L = CO; II, L =  $P(CH_3)_2C_6H_5$ ] toward small unsaturated molecules. $3$  These clusters are important because they are molecules. I hese clusters are important because they are<br>formally "electron deficient" and have been variously described<br>as a sentativities are continued and have been described cluster compounds<br>a new source of h<br>delineate the nature<br>pounds to substrate<br>of the hydridotrios<br> $L = CO$ ; II,  $L = 1$ <br>molecules.<sup>3</sup> Thes<br>formally "electron case containing an o<br> $H$ -Os-H four-cent<br>readily react with d

as containing an osmium-osmium double bond<sup>4</sup> or an Os-

 $H-Os-H$  four-center, four-electron bond.<sup>5</sup> As a result they readily react with donor molecules to form "electron saturated" 1:1 adducts. $6,10$  If the donor is an unsaturated molecule, transfer of hydrogen from the cluster to the substrate becomes a viable process. In this regard, I has been shown to be a catalyst for the hydrogenation of terminal alkenes.' We have previously investigated the nature of hydrogen transfer from these clusters to heteronuclear unsaturated molecules although

- **(1) (a) Heck, R. F. "Organotransition Metal Chemistry"; Academic Press: New York, 1974; Chapter 4. (b) James, B. R. "Homogeneous Hydrogenation"; Wiley-Interscience: New York, 1973; Chapter 11.**
- (2) (a) Muetterties, E. L.; Band, E.; Kokorin, A.; Pretzer, W. R.; Thomas, M. G. Inorg. Chem. 1980, 19, 1552. (b) Sivak, A. J.; Muetterties, E. L. J. Am. Chem. Soc. 1979, 101, 4878. (c) Thomas, M. G.; Pretzer, **W. R.; Beier, B. F.; Hirsekorn, F. J.; Muetterties, E. L.** *Ibid.* **1977, 99, 743. (d) Thomas, M. G.; Beier, B.** F.; **Muetterties, E.** L. *Ibid* **1976,98, 1296.**
- **(3) (a) Adams, R. D.; Golembeski, N. M.** *J. Am. Chem. Soc.* **1979,101, 2579. (b) Adams, R. D.; Golembeski, N. M.** *Ibid.* **1979,101, 1306. (c) Adams, R. D.; Golembeski, N. M.; Selegue, J. P.** *Ibid.* **1979,101,5862. (d) Adams, R. D.; Selegue, J. P.** *Inorg. Chem.* **1980, 19, 1791. (e)**
- **Adams, R. D.; Selegue, J. P.** *Ibid.* **1980, 19, 1795. (4) (a) Mason, R.; Mingos, D. M. P.** *J. Orgunomer. Chem.* **1973.50, 53. (b) Kaesz, H. D.** *Chem. Er.* **1973, 344. (c) Churchill, M. R.; DeBoer, B.** *G.;* **Rotella,** F. **J.** *Inorg. Chem.* **1976,** *15,* **1843.**
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- **(5) Broach, R. W.; Williams, J. M.** *Inorg. Chem.* **1979, 18, 314. (6) Shapley, J. R.; Keister, J. B.; Churchill, M. R.; DeBoer, B. G.** *J. Am. Chem.* **SOC. 1975, 97, 4145.**
- **(7) Keister, J. B.; Shapley, J. R.** *J. Am. Chem. Soc.* **1976, 98, 1056.**

the processes were not catalytic.<sup>3</sup> Here, we report the results of our studies of the reactions of the clusters I and I1 with aryl isocyanates. A preliminary report of this work has been published.8

# **Experimental Section**

General Information. Although the cluster complexes were generally air stable, reactions were routinely performed under a prepurified nitrogen atmosphere. Hexane was purified by distillation from **so**dium-benzophenone; other solvents were stored over **4-A** molecular sieves and degassed with a dispersed stream of nitrogen. Tolyl isocyanate was vacuum distilled and stored under nitrogen. Dimethylphenylphosphine and triethylamine were distilled from sodium metal and stored under nitrogen. Osmium carbonyl was obtained commercially (Strem) or prepared from **OsO,.** Alumina for chromatography was Baker acid-washed aluminum oxide deactivated with 6% water, unless otherwise specified.  $H_2Os_3(CO)_{10}^9$  and  $H_2Os_3(C O$ <sub>9</sub>[P(CH<sub>3</sub>)<sub>2</sub>C<sub>6</sub>H<sub>5</sub>]<sup>10</sup> were prepared by published methods. Other reagents were used as received from commercial sources.

Melting points were determined in evacuated capillary tubes with a Thomas-Hoover and are **uncorrected.** Infrared **spectra** were recorded on a Perkin-Elmer **237B** spectrophotometer. Fourier transform 'H NMR spectra were obtained at **270** MHz on a Bruker **HX270.** Mass spectra were obtained at 20 eV on a Hewlett-Packard **5985** GC/MS with use of a direct-inlet, electron-impact mode.

**Reaction of**  $H_2Os_3(CO)_{10}$  **with**  $p\text{-CH}_3C_6H_4NCO$ **.** A solution of H2Ck3(CO)10 **(1 13** mg, **0.133** mmol) andp-CH3C6H4NC0 **(0.75** mL, ca. **5.3** mmol) in hexane **(50** mL) was heated to reflux for **3** h and **45 min.** All volatiles were removed in vacuo, and the oily yellow residue was dissolved in CDCl<sub>3</sub> and examined by <sup>1</sup>H NMR. This showed four metal hydride containing products (8 **-1 1.26 (>90%); 8 -1 1.46, -14.27, -20.18 (<5%** each)). The mixture was chromatographed on alumina. The first yellow band was eluted with hexane/benzene **(6/1**   $v/v$ ), reduced to dryness, and crystallized from pentane at  $-20$  <sup>o</sup>C to give  $(\mu$ -H) $(\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NCHO)Os<sub>3</sub>(CO)<sub>10</sub> (III) as yellow crystals **(89** mg, **68%).** The second yellow band was eluted with benzene and dichloromethane and then reduced in vacuo and crystallized from

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- *Chem. Soc.* **1975, 97, 3942. (10) Deeming, A. J.; Hasso, S.** *J. Orgammet. Chem.* **1976, 114, 313.**

<sup>(8)</sup> **Adams, R. D.; Golembeski, N. M.** *J. Orgummet. Chem.* **1979, 171, C21.** 

Hydridotriosmium Carbonyl Cluster Compounds



**7.36** br **s (1 H), 7.09** m **(5 H), 3430** w;~ **2105** m, **2065 s, 2055 s,**  170-141 **2.30 s (3 H),-14.07 s (1 H)b 2020** sh, **2010** vs, **1990 s, 1975** mc

 $a$  (CD<sub>3</sub>)<sub>2</sub>CO.  $b$  CDCl<sub>3</sub>. <sup>*c*</sup> Hexane.

dichloromethane/hexane to give  $(\mu$ -H)( $\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NHCO)Os<sub>3</sub>-(CO),, **(VII) (1.5** mg, **1%)** as yellow crystals. **A** mass spectrum of this compound showed a molecular ion *(m/e* **991** corresponding to  ${}^{12}C_{18}{}^{1}H_9{}^{14}N{}^{16}O_{11}{}^{192}Os_3$ ) and fragment ions corresponding to the loss of each of nine carbonyl ligands, all with the expected complex isotope distribution pattern.

This reaction was performed similarily in a solvent consisting of **99/ 1** v/v hexane/triethylamine. There was no significant difference in the products or their ratios.

 $\text{Os}_3(\mu\text{-H})(\mu\text{-C}_6\text{H}_3\text{NCHO})(CO)_{10}$  (IV) was similarly prepared from **I** and C6H5NC0. t-BUNCO does not react with **I** even after **19** h in refluxing hexane.

Reaction of  $(\mu-H)(\mu-p-CH_3C_6H_4NCHO)Os_3(CO)_{10}$  (III) with  $P(CH<sub>3</sub>)<sub>2</sub>C<sub>6</sub>H<sub>5</sub>$ . A solution of III (89 mg, 0.090 mmol) and dimethylphenylphosphine **(0.3** mL, mmol) in heptane (40 mL) was heated to reflux for **3.5** h and then reduced to dryness in vacuo. A <sup>1</sup>H NMR spectrum of the crude product showed three osmium hydride Chromatography on alumina gave two major yellow bands, eluted with hexane/benzene **(7/3** to 4/6, v/v) and benzene, respectively. Both were reduced to dryness in vacuo and crystallized from pentane at  $-20$  °C. Band 1 produced yellow crystals of  $(\mu$ -H) $(\mu$ - $CH_3C_6H_4NCHO$ ) $Os_3(CO)_9[P(CH_3)_2C_6H_5]$  (V) (50.3 mg, 51%). The second band produced **9.3** mg of a yellow solid which was evidently a phosphine disubstitution product ( $\delta$  -10.79 t,  $J_{PH}$  = 11.6 Hz) and was not fully characterized. **species:** *6* **-10.10** t (a. **lo%), -10.82** t (a. **20%), -1 1.22** <sup>d</sup>**(a. 70%).** 

**Reaction of**  $(\mu \cdot H)_2 Os_3(CO)_9 [P(CH_3)_2 C_6 H_5]$  (II) with *p*-CH3C6H,NCO. A solution of 11 (44.5 mg, **0.046** mmol) and p-tolyl isocyanate (0.1 mL, ca. **0.75** mmol) in hexane (20 mL) was heated to reflux for **22** h. The volatiles were removed in vacuo, and the oily residue was chromatographed on alumina. The major yellow band eluted with hexane/benzene **(3/2,** v/v) was shown by 'H NMR to consist of major (6 **-13.92** d, ca. **75%)** and minor (6 **-11.22,** ca. **25%)**  products. Rechromatography on alumina (acid washed,  $2.5\%$   $H_2O$ ) gave a broad band which was collected in two portions. The front yellow portion was eluted with benzene/ethyl ether **(4/1,** v/v), and the orange tail was eluted with methanol. The first portion was reduced to dryness and crystallized from pentane at **-20** 'C to give yellow crystals of the major product  $(\mu-H)(\mu-p-CH_3C_6H_4NHCO)Os_3$ -(C0)9[P(CH3)2C6HS] **(VI) (16.1** mg, **32%).** Recrystallization from CH2Cl&H14 gave yellow crystals of **VI** suitable for X-ray diffraction. The minor product  $(\delta -11.22 d)$  was spectroscopically characterized as **V.** 

**Thermolyses of** Ill **and W.** Compound **111 (89.1** *mg)* was dissolved in **40** mL of heptane and heated to reflux for **4** h. The solvent was removed in vacuo, and the residue was dissolved in ca. **0.3** mL of  $CDCl<sub>3</sub>$ . A <sup>1</sup>H NMR spectrum showed that the sample was unchanged. Compound **VI1** was treated similarly and was also recovered unchanged.

Crystallographic **Analyses.** Crystals of **111, V** and **VI** suitable for diffraction measurements were obtained from hexane solutions by cooling to  $-20$  °C. All crystals were mounted in thin-walled glass capillaries. All diffraction measurements were made on an Enraf-Nonius CAD-4 fully automated four-circle diffractometer using graphite-monochromatized Mo Ka radiation. Unit cells were determined and refined from **25** randomly selected reflections obtained by using the CAD-4 automatic search, center, index and least-squares routines. The space groups were determined from the systematic absences observed during data collection. In the triclinic cases the space groups *Pi* were assumed and were confirmed by the successful solution and refinement of the structures. Crystal data and data collections parameters are listed in Table **VIII.** All data processing was performed on a Digital PDP 11/45 computer with use of the Enraf-Nonius SDP program library. Absorption corrections of a Gaussian integration type were done for all structures. Neutral-atom scattering factors were calculated by the standard procedures.<sup>11a</sup> Anomalous dispersion corrections were applied for all nonhydrogen atoms.<sup>11b</sup> All structures were solved by a combination of Patterson and difference Fourier techniques. Hydrogen atom positions were either obtained from difference Fourier syntheses or calculated on the basis of geometric considerations. Hydrogen atom contributions were included in structure factor calculations, but their positions were not refined. Full-matrix least-squares refinements minimized the function  $\sum_{hkl} w(|F_{\text{obad}}| - |F_{\text{calcd}}|)^2$  where  $w = 1/(\sigma(F))^2$ ,  $\sigma(F) =$ Structure **I11** was refined by using unit weights for all reflections. All atoms heavier than oxygen were refined with anisotropic temperature factors. All other atoms were refined with isotropic temperature factors only. Final fractional atomic coordinates, thermal parameters, and structure factor amplitudes are available for all structures (see sup plementary material). Ineratomic distances and angles with errors obtained from the inverse matrix calculated on the final cycle of least-squares refinement are listed in Tables **11-VII.** The **Os( 1)-C(2)**  bond length of  $1.64$  (1)  $\hat{A}$  in V is anomalously short. The C(2)–O(2) at  $1.23$  (1)  $\AA$  is a little long. It seems atom  $C(2)$  may be slightly incorrectly positioned in this structure. In spite of the final refinement, we feel that these distances should not be taken too seriously.  $(F_{\text{obsd}}^2)/2F_{\text{obsd}}$  and  $\sigma(F_{\text{obsd}}^2) = [(\sigma(I_{\text{raw}}))^2 + (PF_{\text{obsd}}^2)^2]^{1/2}/Lp$ .

#### **Results**

**The Formamido Complexes.** The principal products isolated from the reaction of I with aryl isocyanates are the complexes  $(\mu-H)(\mu-ArNCHO)Os<sub>3</sub>(CO)<sub>10</sub>$ , which contain formamido ligands. These complexes were formed by an addition of one isocyanate molecule to the cluster, which was accompanied by a transfer of one hydrogen atom from the cluster to the carbon atom of the isocyanate. The presence of the hydrogen atom on the carbon atom was indicated by the 'H **NMR** 

**<sup>(1 1) &</sup>quot;International Tables for X-ray Crystallography"; Kynoch Press: Birmingham, England, 1975; Vol. IV: (a) Table 2.2B, pp 99-101; (b) Table 2.3.1, pp 149-150.** 

**Table 11.** Interatomic Distances **(A)** with Esd's for  $(\mu$ -H $)(\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NCHO $)$ Os<sub>3</sub>(CO)<sub>10</sub>(III)

$Os(1)-Os(2)$	2.903(1)	$C(11) - C(12)$	1.40(1)
$Os(1) - Os(3)$	2.909(1)	$C(12) - C(13)$	1.39 (1)
$Os(2)-Os(3)$	2.888(1)	$C(13) - C(14)$	1.36(2)
$Os(1)$ –C $(5)$	1.912(9)	$C(14) - C(15)$	1.39(2)
Os(1) – C(6)	1.898 (9)	$C(15) - C(16)$	1.41(1)
$Os(1)-C(7)$	1.910(8)	C(16)–C(11)	1.39(1)
Os(1)–N	2.156(6)	$C(14) - C(18)$	1.56 (2)
Os(2) – C(1)	1.963(8)	$C(1) - O(1)$	1.13(1)
Os(2) – C(2)	1.948(9)	$C(2)-O(2)$	1.15(1)
$Os(2) - C(3)$	1.901(8)	$C(3) - O(3)$	1.15(1)
Os(2) – C(4)	1.924 (10)	$C(4)-O(4)$	1.17(1)
$Os(3)-C(8)$	1.902 (10)	$C(5)-O(5)$	1.13(1)
Os(3)–C(9)	1.889 (9)	$C(6)-O(6)$	1.17(1)
Os(3) – C(10)	1.903 (11)	$C(7)-O(7)$	1.14(1)
Os(3)–O(11)	2.145(6)	$C(8)-O(8)$	1.18(1)
C(17)–O(11)	1.27(1)	$C(9)-O(9)$	1.16(1)
C(17)–N	1.33(1)	$C(10)-O(10)$	1.16 (1)
N–C(11)	1.45(1)		

**Table 111.** Interatomic Angles (Deg) with Esd's for  $(\mu$ -H)( $\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NCHO)Os<sub>3</sub>(CO)<sub>10</sub> (1II)



spectra, which showed very low-field resonances ( $\delta$  8-9), characteristic of formamido and related ligands,<sup>12</sup> and indirectly supported by an X-ray crystallographic analysis of the p-tolyl derivative.

An ORTEP drawing of the molecular structure of  $(\mu$ -H) $(\mu$  $p$ -CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NCHO)Os<sub>3</sub>(CO)<sub>10</sub> (III) is shown in Figure 1. The formamido hydrogen atom was not observed crystallographically, but it is shown in an idealized position in the figure. Interatomic distances and angles are listed in Tables I1 and 111. The molecule consists of a triangular cluster of three osmium atoms. The three osmium-osmium bond distances range from 2.888 (1) to 2.909 (1) **A** and are only





**Table V.** Interatomic Angles (Deg) with Esd's for  $(\mu$ -H $)(\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NCHO $)$ Os<sub>3</sub>(CO)<sub>9</sub> [P(CH<sub>3</sub>)<sub>2</sub>C<sub>6</sub>H<sub>5</sub>] (V)



slightly longer than the osmium-osmium bond distance **(2.877**  (1)  $\hat{A}$ ) found in  $Os_3(CO)_{12}$ .<sup>13</sup> The formamido ligand is bonded in a "diaxial" coordination arrangement (i.e., in two coordination positions perpendicular to the Os<sub>3</sub> plane) and bridges the  $Os(1)$ - $Os(3)$  edge of the cluster. The oxygen atom  $O(11)$ 

(13) Churchill, **M.** R.; DeBoer, B. G. *Inorg. Chem.* **1977.16, 878.** 

<sup>(12) (</sup>a) Sahajpal, A.; Robinson, S. D. *Inorg. Chem.* 1979, 18, 3572. (b)<br>Robinson, S. D.; Sahajpal, A. *Ibid.* 1977, 16, 2722. (c) Brown, L. D.;<br>Robinson, S. D.; Sahajpal, A.; Ibers, J. A. *Ibid.* 1977, 16, 2728.

**Table VI.** Interatomic Distances **(A)** with **Esd's** for  $(\mu$ -H $)(\mu$ -p-CH<sub>3</sub>C<sub>6</sub> H<sub>4</sub>NHCO $)$ Os<sub>3</sub> (CO)<sub>9</sub> [P(CH<sub>3</sub>),C<sub>6</sub>H<sub>5</sub>] (VI)

$Os(1)-Os(2)$	2.945(1)	$C(15)-C(16)$	1.393 (10)
$Os(1) - Os(3)$	2.859 (1)	$C(16) - C(11)$	1.342(9)
$Os(2)-Os(3)$	2.893 (1)	$C(14) - C(17)$	1.515 (12)
$Os(1)$ -P	2.359 (2)	P-C(27)	1.828 (9)
Os(1) – C(1)	1.827(8)	P–C(28)	1.842(8)
Os(1) – C(2)	1.867 (9)	$P - C(21)$	1.816(8)
$Os(1)-O(10)$	2.141(4)	$C(21) - C(22)$	1.376(9)
$Os(2) - C(3)$	1.890 (9)	$C(22) - C(23)$	1.446 (11)
Os(2) – C(4)	1.925 (9)	$C(23) - C(24)$	1.375 (12)
$Os(2) - C(5)$	1.866 (10)	$C(24)-C(25)$	1.331 (11)
Os(2) – C(10)	2.121(7)	$C(25)-C(26)$	1.407 (11)
Os(3) – C(6)	1.899 (10)	$C(26)-C(21)$	1.402 (10)
$Os(3)-C(7)$	1.859 (11)	$C(1)-O(1)$	1.197(8)
Os(3) – C(8)	1.948 (9)	$C(2)-O(2)$	1.192(9)
Os(3) – C(9)	1.898 (9)	$C(3)-O(3)$	1.153(9)
$C(10)-O(10)$	1.254 (9)	$C(4)-O(4)$	1.160 (9)
$C(10)-N$	1.345(8)	$C(5)-O(5)$	1.193 (11)
$N-C(11)$	1.442(8)	$C(6)-O(6)$	1.187 (10)
$C(11) - C(12)$	1.370(11)	$C(7)-O(7)$	1.199 (11)
$C(12) - C(13)$	1.403(12)	$C(8)-O(8)$	1.164 (9)
$C(13) - C(14)$	1.371 (10)	$C(9)-O(9)$	1.162 (9)
$C(14) - C(15)$	1.360 (10)		

**Table MI.** Interatomic Angles (Deg) with **Esd's** for  $(\mu$ -H)( $\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NHCO)Os<sub>3</sub>(CO)<sub>9</sub>[P(CH<sub>3</sub>)<sub>2</sub>C<sub>6</sub>H<sub>5</sub>](VI)



is bonded solely to Os(3) while the nitrogen atom is bonded solely to  $Os(1)$ . The  $C(17)-O(11)$  and  $C(17)-N$  bond lengths at 1.27 (1) and 1.33 (1) **A,** respectively, are intermediate between single- and double-bond lengths and indicate significant electron delocalization across the N-C-0 unit. **A** hydride ligand indicated by its <sup>1</sup>H NMR resonance  $\delta$  –11.35 was not observed crystallographically but is believed to occupy a bridging position across the  $Os(1)-Os(3)$  bond. This is sup-



**Figure 1.** ORTEP diagram of  $(\mu$ -H) $(\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>HNCHO)Os<sub>3</sub>(CO)<sub>10</sub> (III) showing 50% probability ellipsoids. The hydrogen atom H<sub>1</sub> is shown in an idealized position.



**Figure 2.** ORTEP diagram of  $(\mu-H)(\mu-p-CH_3C_6H_4NCHO)Os_3$ -**(C0)9[P(CH3)2C6H5] (V)** showing *50%* probability ellipsoids.

ported by a bending away of the carbonyl ligands  $C(6)-O(6)$ and C(10)-O(10) as characterized by the large angles **Os-**  (3)-Os(1)-C(6) (120.0 (3)<sup>o</sup>) and Os(1)-Os(3)-C(10) (117.8 (3)<sup>o</sup>) from the Os(1)-Os(3) bond. The complex contains ten linear terminal carbonyl ligands. **Six** lie essentially in the plane of the Os<sub>3</sub> triangle while four are perpendicular to it.

Complex I11 reacts with dimethlphenylphosphine in refluxing heptane to give the monosubstituted product  $(\mu-H)$ -**(p-pCH3C6H4NCHO)Os3(C0)9[P(CH3)2C6H5]** (V) in **51%**  yield. This complex was also analyzed crystallographically, and an ORTEP diagram of its molecular structure is shown in Figure 2. Interatomic distances and angles are listed in Tables IV and **V.** Like 111, V also contains a triangular cluster of osmium atoms. The osmium-osmium bond distances are slightly longer than those found in 111. The increase is similar to the 0.02 **A** increase in metal-metal bond lengths, which was observed when a phosphite ligand was substituted for a carbonyl ligand in  $\mathrm{Os}_3(\mathrm{CO})_{12}$ .<sup>14</sup> The p-tolylformamido ligand shows a diaxial coordination and bridges an edge of the cluster. The formamido hydrogen and hydride ligands were not observed crystallographically but were confirmed spectroscopically. The hydride ligand probably bridges the  $\overline{Os}(1)-\overline{Os}(2)$ bond. The dimethylphenylphosphine ligand occupies an equatorial coordination site on Os(2) approximately trans to

**<sup>(14)</sup> Eknfield, R. E.; Johnson, B. F.** *G.;* Raitbby, **P. R.; Sheldrick,** *G.* **M.** *Acra Crystallogr., Sect. B* **1978,** *B34, 666.* 





<sup>a</sup> For all three compounds.

the  $Os(2)-Os(3)$  bond. There are nine linear terminal carbonyl ligands.

**The Carboxamido Complexes.** The major product (isolated in 32% yield) obtained from the reaction of I1 with p-tolyl isocyanate has the formula  $(\mu$ -H) $(\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NHCO)- $Os_3(CO)_9[P(CH_3)_2C_6H_5]$  (VI). This complex was characterized spectroscopically and by an X-ray crystallographic analysis. The infrared spectrum of VI shows an absorption at 3440 cm<sup>-1</sup>, which can be attributed to an N-H stretching vibration while a broad resonance at  $\delta$  7.23 in the <sup>1</sup>H NMR spectrum can be assigned to an N-H hydrogen atom. A hydride resonance was observed at  $-13.92$  ppm.

An **ORTEP** diagram of VI is shown in Figure 3. Interatomic distances and angles are given in Tables VI and VII. This complex also contains a triangular cluster of three osmium atoms. There are nine linear terminal carbonyl ligands. **A**  dimethylphenylphosphine ligand is coordinated in an equatorial coordination site to  $Os(1)$ . The most interesting ligand is a carboxamido ligand which bridges the **Os(** 1)-Os(2) edge of the cluster in a diaxial coordination arrangement. It is bonded to the cluster through the carbon atom  $C(10)$  to  $Os(2)$  and the oxygen atom  $O(10)$  to  $Os(1)$ . The  $C(10)-O(10)$  distance at 1.254 (8) **A** and the C(lO)-N distance at 1.345 (8) **A**  indicate multiple-bond character in the form of a delocalization through the N-C-O unit. A hydrogen arom is believed to be bonded to the nitrogen atom, but this was not observed crystallographically. The hydride ligand was not observed crystallographically, but it is believed to bridge the **Os(** 1)- Os(2) edge of the cluster as in I11 and V. V was also obtained in the reaction of II with  $p$ -tolyl isocyanate. It was difficult to separate pure V from VI, but, spectroscopically, the ratio of V/VI was approximately 1/3.

A minor product (1% yield) was also isolated from the reaction of I with  $p\text{-}CH_3C_6H_4NCO$ . Its infrared spectrum showed an N-H stretching vibration at  $3430 \text{ cm}^{-1}$  and a carbonyl absorption pattern which resembled that of 111. Its <sup>1</sup>H NMR spectrum showed a broad resonance at  $\delta$  7.36 (N-H) and a high-field singlet at  $-14.07$  ppm which can be attributed to a bridging hydride ligand. The mass spectrum showed a parent ion at *m/e* 991 **(1920s)** and ions corresponding to the loss of each of nine carbonyl ligands. Accordingly, we formulate this product as  $(\mu$ -H) $(\mu$ - $p$ -CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NHCO)Os<sub>3</sub>(CO)<sub>10</sub>

#### Scheme I





Figure 3. ORTEP diagram of  $(\mu$ -H)( $\mu$ -p-CH<sub>3</sub>C<sub>6</sub>H<sub>4</sub>NHCO)Os<sub>3</sub>- $(CO)_9[P(CH_3)_2C_6H_5]$  (VI) showing 50% probability ellipsoids.

(VII), which we believe is a decacarbonyl analogue of **VI.**  Allowing that VI1 might have been formed from I11 or vice versa, we thermolyzed pure samples of I11 and VI1 independently. I11 could not be converted into VI1 and vice versa even under conditions more severe than those of their formation. Similar results were obtained for V and VI.

### Discussion

Mononuclear transition metal hydride complexes react with isocyanates by hydrogen transfer, producing either chelating formamido complexes<sup>12a</sup> or C-coordinated carboxamido complexes.<sup>15</sup> The latter products are formed in base-promoted reactions and are believed to proceed via a proton abstraction from the metal hydride and readdition to a metal isocyanate adduct. **l5** 

The reaction of I with aryl isocyanates is exemplified in Scheme I. Hydride transfer to either the carbon or nitrogen atom of the isocyanate can occur, although transfer to the carbon is by far the preferred route. Both the formamido and carboxamido ligands bridge an edge of the cluster. Since I11 and VI1 cannot be interconverted, it appears that they must have been formed in independent, competing reactions. While we have shown previously that intramolecular and dissociative hydrogen-transfer processes can produce different products,<sup>3a</sup> this does not appear to be an explanation of the difference in this case since the same products in the same ratios were obtained in both hexane and triethylamine/hexane solvents. Whether or not an adduct between I and the isocyanate

**(15)** Angelici. R. J. *Acc.* Chem. Res. **1972, 5, 335** and references therein.

molecule is formed prior to hydrogen transfer remains to be established. Although we have not observed any such adducts, we believe the reactions most probably do proceed via such intermediates.6 Carboxamido complexes similar to VI1 have been formed through the reaction of primary amines with  $Os_3(CO)_{12}.^{16}$ 

The molecular structure of  $II^{17}$  is similar to that of  $(H)_{2}$ - $Os_3(CO)_9P(C_6H_5)_3^{18}$  and  $(H)_2Os_3(CO)_9(CN-t-C_4H_9),^{19}$  having the phosphine ligand in an equatorial coordination site with the bridging hydride ligands along the adjacent "unsaturated" osmium-osmium bond. **In** contrast to the reaction of I, the reaction of II with  $p$ -tolyl isocyanate proceeds predominantly with transfer of hydrogen to the *nitrogen atom* with formation of the carboxamido complex VI. The arrangement of ligands in the product suggests that the isocyanate molecule has entered the cluster at the hydrogen-bridged osmium-osmium bond, the presumed site of highest reactivity.<sup>20</sup> Why hydrogen transfer from I1 occurs preferentially to the nitrogen atom while the opposite was true for I has not been revealed by this work. Steric bulk introduced by the phosphine and an increase in electron density on the metal atoms caused by the more highly donating phosphine ligand are reasonable explanations; however, in either case the result is clear. *Changing the ligand structure of the cluster has caused a pronounced change in the regioselectivity of the hydrogen transfer.* Such effects could play an important role in product control in some cases of cluster catalysis.

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**Supplementary Material Available:** Final fractional atomic coordinates, thermal parameters, and structure factor amplitudes for all three structures (75 pages). Ordering information is given **on** any current masthead page.

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