.

Contribution from the Department of Chemistry, The Pennsylvania State University, University Park, Pennsylvania 16802

Molecular Dynamics of Mixed-Metal Clusters. Carbon- 13 and Hydrogen- 1 NMR Studies of $H_2Feku_3(CO)_{13}$, $H_2FeRu_2Os(CO)_{13}$, and $H_2FeRuOs_2(CO)_{13}$

WAYNE L. GLADFELTER and GREGORY L. GEOFFROY*

Received April 1 I, 1980

Variable-temperature ¹H and ¹³C NMR spectra of $H_2Feku_3(CO)_{13}$, $H_2Feku_2Os(CO)_{13}$, and $H_2Feku_0S_2(CO)_{13}$ have been measured and interpreted in view of three distinct fluxional processes. In the lowest temperature process the bridging and terminal carbonyls bound to Fe undergo localized exchange. The second process involves migration of the carbonyls around the face of the tetrahedral cluster that contains the bridging carbonyls. The highest temperature process involves a subtle reorganization of the metal framework. This latter process accounts for interconversion of the two enantiomers of the C_1 isomer of H₂FeRu₂Os(CO)₁₃ and H₂FeRuQs₂(CO)₁₃ and allows for the $C_s \rightleftharpoons C_1$ isomerization. Line-shape analysis of the ¹H NMR spectra and comparison of the ¹³C NMR spectra show that the activation barrier for each process increases as the **Os** content of the cluster increases.

Introduction

Metal clusters have been shown to undergo a wide variety of fluxional processes in which carbonyls, hydrides, and even the metals themselves undergo rearrangement.' Mixed-metal clusters are ideally suited for studies of the fluxional processes **of** clusters because of the low symmetry which is inherent within their metal framework. In such clusters, the majority of ligands are in chemically nonequivalent positions and are thus distinguishable by NMR spectroscopy.2 Furthermore, a homologous series of mixed-metal clusters allows one to study the effects of metal substitution on the fluxional processes and their activation parameters.

In a preliminary communication we reported the results of a ¹³C NMR study of $H_2FeRu_3(CO)_{13}$ and $H_2FeRuOs_2(CO)_{13}$ in which three distinct fluxional processes were observed, with the highest activation process involving a subtle rearrangement of the metal framework of each cluster. 3 The full details of the **'H** NMR spectra of these two clusters are described herein along with the ¹H and ¹³C NMR spectra of $H_2FeRu_2Os(C-$ **O)13.** This last cluster completes the series and allows an assessment of the effect of progressive substitution of Os for Ru within the series.

Experimental Section

H₂FeRu₃(CO)₁₃, H₂FeRu₂Os(CO)₁₃, and H₂FeRuOs₂(CO)₁₃ were prepared by published procedures.⁴ Enrichment of each of these compounds was accomplished by stirring hexane solutions of each under an atmosphere of 90% enriched ¹³CO (Stohler Isotope Co.). Exact enrichment conditions varied: 70% enrichment of H₂FeRu₃- $(CO)_{13}$ after being stirred at 45-50 °C for 140 h; 60% enrichment of $H_2FeRu_2Os(CO)_{13}$ after being stirred at 25 °C for 380 h; 40% enrichment of H₂FeRuOs₂(CO)₁₃ after being stirred at 25 °C for 181 h. The approximate percent enrichment was determined by mass spectrometry.

NMR spectra were recorded by using 10-mm NMR tubes containing \sim 3 mL of sample solution. Each tube was thoroughly degassed by using freeze-pump-thaw techniques and then sealed under vacuum. Samples used for the ¹³C NMR studies contained $[Cr(\text{acc})_3]$ (0.05 M) as a shiftless relaxation agent. The lock substance in each case was deuterated solvent. Spectra were recorded on a JEOL PS-100 **FT** NMR spectrometer equipped with a Nicolet 1080 computer for analysis. Heteronuclear decoupling was accomplished by using a broad-band noise decoupler. All reported **'H** and **I3C** chemical shifts are relative to Me4Si. Line-shape analyses were conducted on The Pennsylvania State University's IBM 370 computer utilizing the

- Gladfelter, W. L.; Geoffroy, G. L. Adv. Organomet. Chem. 1980, 18, in press.
- (3) Geoffroy, G. L.; Gladfelter, W. L. *J. Am. Chem. SOC.* 1977,99,6775. (4) Geoffroy, G. L.; Gladfelter, W. L. *J. Am. Chem. Soc.* 1977, 99, 7565.

program DNMR3.⁵ Infrared spectra were recorded on a Perkin-Elmer 580 grating infrared spectrophotometer, and variable-temperature IR spectra were recorded by using an infrared cell manufactured by Harrick Scientific Corp.

Results

The trimetallic clusters $H_2FeRu_2Os(CO)_{13}$ and H_2FeRu - $Os₂(CO)₁₃$ each exist in the two isomeric forms shown in Figure 1. These will hereafter be denoted by their C_1 and *C,* symmetry labels. The isomers of each cluster have not been separated, and indeed the 'H NMR spectra discussed below show that the isomers readily interconvert at laboratory temperatures. Analysis of the bridging carbonyl region of the IR spectrum of each of the trimetallic clusters shows that, at 25 ^oC in hexane solution, the C_1 isomer of $H_2FeRu_2Os(CO)_{13}$ is in greater abundance than the C_s isomer whereas for H_2 -FeRuOs₂(CO)₁₃ the isomers are of about equal concentration.^{4,6} Infrared spectroscopy also shows that the $C_s \rightleftharpoons C_1$ equilibrium of H_2F eRuOs₂(CO)₁₃ shifts to the left upon cooling. We would ideally like to compare the values obtained at different temperatures for the equilibrium concentrations of the two isomers by using IR and 'H NMR spectroscopy. However, the resolution of the IR spectra and the necessity of using different solvents for the two studies' coupled with the likely solvent dependence of the $C_s \rightleftarrows C_1$ equilibrium have prevented us from obtaining meaningful results.

'H NMR Spectra. The variable-temperature 'H NMR spectra of $H_2FeRu_2Os(CO)_{13}$ and $H_2FeRuOs_2(CO)_{13}$ are shown in Figures **2** and **3.** In the low-temperature limiting spectrum each cluster shows a singlet and a pair of doublets. The singlet is attributed to the two equivalent hydrogens of the *C,* isomer while the pair of doublets are assigned to the two nonequivalent hydrogens of the C_1 isomer. Pertinent ¹H NMR spectral parameters for the limiting spectra are given in Table I. The average magnitude of the chemical shift (negative) for the $H_2FeRu_3(CO)_{13}$, $H_2FeRu_2Os(CO)_{13}$, and $H_2FeRuOs_2(CO)_{13}$ series increases as the Os content of the

⁽¹⁾ Band, E.; Muetterties, E. L. *Chem. Rev.* 1978, *78,* 639.

⁽⁵⁾ Kleier, D. **A,;** Binsch, G. "DNMR3: A Computer Program for the Calculation of Complex Exchange-Broadened NMR Spectra. Modified Version for Spin Systems Exhibiting Magnetic Equivalence or Symmetry"; Quantum Chemistry Program Exchange: Indiana University, Bloomington, Indiana; Program 1065.

⁽⁶⁾ The bridging $\nu_{\rm CO}$ region of the spectra of $H_2F\in Ru_2Os(CO)_{13}$ and $H_2-F\in Ru_2Os(CO)_{13}$ are shown in Figure 4 of ref 4. For $H_2F\in Ru_2Os(CO)_{13}$ the bands at 1887 and 1861 cm⁻¹ are attributed to the C_s isome given in the text of ref **4.**

Saturated hydrocarbon solvents are required to obtain high-resolution IR spectra; the clusters are not sufficiently soluble in hydrocarbons for high-quality NMR spectra.

Figure 1. Carbonyl labeling schemes for $H_2Feku_3(CO)_{13}$ and the C_s and C_1 isomers of H₂FeRu₂Os(CO)₁₃ and H₂FeRuOs₂(CO)₁₃.

Figure 2. ¹H NMR spectra of $H_2FeRu_2Os(CO)_{13}$. Spectra at -41, -35, and -15 °C were recorded in an acetone- $d_6/CHCl_2F$ solvent mixture while those above -15 °C were obtained in CDCl₃. The chemical shift scale shown is for the CDCI₃ solution. The acetone $d_6/CHC1_2F$ chemical shift values are δ -18.56 d (1.65), -18.75 s, -18.83 d (1.65).

Figure 3. ¹H NMR spectra of $H_2FeRuOs_2(CO)_{13}$ recorded in CDCl₃ solution.

Scheme I

cluster increases, and for each of the C_1 isomers we attribute the upfield doublet to the hydride which bridges the M-M bond with the greatest Os content. Indeed, the data given in Table I are fully consistent with the chemical shift ordering

 > 8 $\frac{1}{8u - 0s}$ > 8 $\frac{1}{0s - 0s}$

As the temperature is raised, the resonances in the spectra of each cluster broaden as the hydrogens begin to exchange positions, coalesce, and finally sharpen to a singlet at the high-temperature limit (Figures **2** and 3). The variable temperature spectra were computer-simulated with use of the model shown in Scheme I for $H_2FeRu_2Os(CO)_{13}$. It was necessary to include both processes shown in Scheme I which involve isomerization (k_2) and exchange of the two hydrogens of the C_1 isomer (k_1) . Computer simulation using a single rate parameter (k) , which might represent either of the processes shown in Scheme I or a constant ratio of the two processes,

3. ^c This value is approximate due to overlap of several resonances in this region. *a* See Figure 1 for labeling schemes. Resonances grouped together cannot be individually assigned to the indicated carbonyls. ^b Reference

did not give a satisfactory fit to the experimental spectra. The kinetic parameters which were derived from the line-shape analyses are given in Table I, and a comparison of the calculated and experimental spectra of H_2 FeRuOs₂(CO)₁₃ is shown in Figure 3. For both $H_2FeRu_2Os(CO)_{13}$ and H_2Fe - $RuOs₂(CO)₁₃$, the ¹H NMR spectra indicate that the isomerization process occurs with a lower activation energy than does hydrogen exchange localized on the C_1 isomer.

The line-shape analysis requires a knowledge of the relative concentrations of the C_1 and C_s isomers at each temperature. These values were obtained by integration of spectra measured at temperatures below exchange followed by extrapolation to higher temperatures. The $C_s \rightleftharpoons C_1$ equilibrium was observed to shift to the left upon cooling of each cluster, in complete accord with the IR results mentioned above, and the ΔH and ΔS values obtained from measurement of the equilibrium constants at various temperatures are included in Table I.

Solubility problems prevented measurement of the spectra of $H_2FeRu_2Os(CO)_{13}$ in the same solvent over the entire temperature range; CDCl₃ was employed above -8 °C whereas an acetone- $d_6/CHCl_2F$ mixture was used at lower temperatures. Slight differences in the chemical shifts were noted in the two different solvents, and we were thus unable to model the exchange processes of this cluster over the entire temperature range. The ΔG^* values listed in Table I were obtained by simulating only the three lowest temperature spectra, and the rate constants obtained at -15 °C were used for the ΔG^* calculation.

Low-Temperature Limiting 13C NMR Spectra. The lowtemperature limiting ¹³C NMR spectra for $H_2FeRu_3(CO)_{13}$, $H_2FeRu_2Os(CO)_{13}$, and $H_2FeRuOs_2(CO)_{13}$ are shown in Figure 4. Figure 1 gives the labeling schemes used for assigning the observed resonances to specific carbonyls.

 H_2 **FeRu₃(CO)₁₃ and** H_2 **FeRuOs₂(CO)₁₃. The resonances in** the low-temperature limiting spectra of $H_2FeRu_3(CO)_{13}$ and $H_2FeRuOs_2(CO)_{13}$ were assigned earlier in ref 3, and the detailed rationale need not be repeated here. The specific assignments are summarized in Table 11. Basically, three important factors were considered in developing these assignments. First, carbonyl ligands which are located trans to the hydride ligands show stronger ${}^{13}C-{}^{1}H$ coupling than do carbonyls cis to the hydrides. Comparison of the 13 C and ${}^{13}C{}^{1}H$ } spectra thus allows a determination of which resonances are due to CO's trans to the hydrides and which are not. Second, it was found that the terminal carbonyls bound to different metals group together in characteristic chemical shift regions. The chemical shift decreases relative to $Me₄Si$ upon descending the triad: Fe $(204-211 \text{ ppm})$ > Ru $(184-189 \text{ nm})$ ppm) > Os (168-177 ppm). Third, for $H_2FeRuOs_2(CO)_{13}$ it was necessary to rely on information gained from its 'H NMR spectrum which showed the *C,* isomer to be more abundant than the C_1 isomer; thus the more intense resonances in the 13C NMR spectra are assigned to the carbonyls on the *C,* isomer.

 $H_2Feku_2Os(CO)_{13}$. The assignment of the resonances in the ¹³C NMR spectrum of $H_2FeRu_2Os(CO)_{13}$ is not as unambiguous as that for $H_2FeRuOs_2(CO)_{13}$, primarily because

Figure 4. Low-temperature limiting 13 C NMR spectra: (a) H_2 Fe- $Ru_3(CO)_{13}$ (-95 °C), (b) $H_2FeRu_2Os(CO)_{13}$ (-90 °C), and (c) $H_2FeRuOs_2(CO)_{13}$ (-60 °C).

neither the C_s nor the C_1 isomer is dominant at low temperature and we cannot argue **on** the basis of relative intensity. Furthermore, the peak widths are broader in the $^{13}C^{1}H$ spectra of $H_2FeRu_2Os(CO)_{13}$ (Figure 5) making the comparison of the 13 C and 13 C 11 H 13 spectra difficult. The increased broadening is likely due to increased $^{13}C^{-13}C$ coupling since the $H_2FeRu_2Os(CO)_{13}$ cluster was enriched to 60% ¹³CO compared to 40% enrichment for $H_2FeRuOs_2(CO)_{13}$.

As with $H_2F\in Ru_3(CO)_{13}$ and $H_2F\in RuO_{52}(CO)_{13}$, the carbonyl resonances in $H_2FeRu_2Os(CO)_{13}$ are grouped by metals, with the Os carbonyls in the 170-177-ppm range, the Ru carbonyls in the 185-193-ppm range, and the Fe carbonyls above 204 ppm. The resonances at 221, 232, and 213 ppm are respectively assigned to the bridging carbonyls a, i, and j. These assignments are made primarily by a comparison to the spectra of $H_2FeRu_3(CO)_{13}$ and $H_2FeRuOs_2(CO)_{13}$: carbonyls a (227 ppm) of the C_s isomer of $H_2FeRu_2Os(CO)_{13}$ both bridge Fe-Ru bonds and hence are in similar chemical environments as a in $H_2Feku_3(CO)_{13}$ (229 ppm); carbonyl i of $H_2FeRu_2Os(CO)_{13}$ (232 ppm) and i of $H_2FeRuOs_2(CO)_{13}$ (229 ppm) also both bridge Fe-Ru bonds and hence have similar chemical shifts; carbonyl j of H₂FeRu₂Os(CO)₁₃ (213 ppm) and j of $H_2FeRuOs_2(CO)_{13}$ (211 ppm) both bridge Fe-Os bonds. **As** with the terminal carbonyls, the bridging carbonyls that are associated with Os occur upfield of those associated with only Fe or Ru.

Figure 5. ¹³C NMR spectra of $H_2FeRu_2Os(CO)_{13}$. The resonances at 201.1 and 200.6 ppm which average to the 192.2-ppm resonance and which are marked with a dagger (t) are due to $Ru₂O_s(CO)₁₂$ impurity which arises from decomposition of $H_2FeRu_2Os(CO)_{12}$ during the enrichment process. This was confirmed by isolating $Ru₂Os(CO)₁₂$ and determining its spectrum. The numbered resonances in the -90 OC spectrum are attributed to carbonyls which cannot be distinguished: (1) b, c, k, 1; (2) see text; **(3)** *e,* **p, q.**

Resonances attributed to the terminal Fe carbonyls of $H_2FeRu_2Os(CO)_{13}$ (b, c, k, l) are at 211 and 204 ppm. These resonances cannot be specifically assigned but each is apparently the sum of two accidentally equivalent resonances, one from each isomer. Their chemical shifts compare well to those of $H_2Feku_3(CO)_{13}$ (211 and 203 ppm) and H_2Feku - $\mathrm{Os}_2(CO)_{13}$ (C₅, 211 and 204 ppm; C₁, 210 and 201 ppm).

Five nonequivalent carbonyls are bound to Os in the two isomers $(C_s, d$ and e; C_1 , p, q, and t), and four resonances are clearly observed in the $170-177$ -ppm range. The large resonance at 173 ppm apparently arises from two signals. The peak at 174 ppm is the only one obviously affected by 'H coupling, indicating it is either e on the C_s isomer or t on the C_1 isomer. At -20 °C, before the onset of isomerization, this is one of the three remaining Os carbonyl resonances (Figure *5).* On the basis of the exchange processes that occur in $H_2FeRu_3(CO)_{13}$ and $H_2FeRuOs_2(CO)_{13}$ (vide infra), these three carbonyls would be d and e on the *C,* isomer and t on the C_1 isomer. It is also apparent that at this temperature the 174-ppm resonance has increased in intensity relative to the other Os carbonyls. Recalling that the $C_s \rightleftharpoons C_1$ equilibrium shifts to the right with increasing temperature, this carbonyl must be bound to the C_1 isomer and is therefore attributed to carbonyl t. At this temperature, the 170- and 173-ppm resonances show a 1:2 intensity ratio and are assigned to carbonyls d and e, respectively. The second peak at 173 ppm and the peak at 177 ppm must be assigned to carbonyls p and 9.

We are unable to definitively assign the resonances due the Ru bound carbonyls. However, five resonances remain in the Ru carbonyl region in the -20 "C spectrum (Figure *5),* and on the basis of the fluxional processes discussed below, these should correspond to carbonyls h on the *C,* isomer and *0,* r, s, and u on the C_1 isomer. Of these five resonances, the peak at 188 ppm decreases in intensity relative to the others as the temperature is raised, indicating that it is due to carbonyl h on the C, isomer.

Carbonyl-Exchange Processes. H₂FeRu₃(CO)₁₃. The variable-temperature ¹³C NMR spectra of H_2 FeRu₃(CO)₁₃ were shown in Figure 1 of ref 3, and the corresponding spectral changes were discussed therein. Here we simply summarize the main features of those results. **As** the temperature is raised from the low-temperature limiting spectrum at -95 °C, resonances due to carbonyls a-c collapse, indicating the occurrence of bridge-terminal interchange localized on Fe as the first process. Before these peaks disappear completely, a second fluxional process begins at -65 °C and averages resonances due to carbonyls a-d and h

The most reasonable mechanism that would average these carbonyls involves their movement in a cyclic process about the $Fe-Ru_1-Ru_2$ plane.

The final fluxional process begins at -45 °C and simultaneously averages the three remaining carbonyls e-g with a-d and h. We have proposed³ that this final exchange process occurs by a shift in the metal framework in which the Fe atom moves closer to Ru_3 and away from Ru_1 with a concomitant shift of the bridging carbonyls and one bridging hydrogen. This generates a structure completely identical with the initial structure but in which the $Fe-Ru_2-Ru_3$ triangle now possesses the bridging carbonyls

A cyclic process, identical with that described above, can now occur around the Fe-Ru₂-Ru₃ plane. A third equivalent shift would allow a cyclic process around the Fe-Ru₁-Ru₂ plane. These equivalent shifts and the resultant cyclic movements will have the effect of averaging all the carbonyls in the molecule. An alternate final exchange process, consistent with the observed NMR changes, would involve localized scrambling on Ru_1 coupled with a cyclic process around the $Ru_1-Ru_2-Ru_3$ triangle. Although these two possible mechanisms cannot be distinguished for $H_2FeRu_3(CO)_{13}$, the NMR data discussed below for $H_2FeRuOs_2(CO)_{13}$ unambiguously show that the exchange in the latter cluster occurs by the intrametallic rearrangement process.³

 H_2 FeRuOs₂(CO)₁₃. The variable-temperature ¹³C NMR spectra of $H_2FeRuOs_2(CO)_{13}$ are shown in Figure 6. As the temperature is raised, three basic carbonyl-exchange processes appear. As for $H_2FeRu_3(CO)_{13}$, the bridging and terminal iron carbonyls exchange in the lowest temperature process. These iron carbonyl resonances are fully broadened by -40 \degree C in the C_1 isomer, while those in C_s are not broadened until -20 °C. The second process, exactly analogous to that in $H_2FeRu_3(CO)_{13}$, averages carbonyls i-m, p, q, and presumably n of C_1 and carbonyls a-c, f, and g of C_s through a cyclic movement of the carbonyls around the plane that originally contained the bridging carbonyls. Again, C_1 begins this process about 40 °C before C_s .

The third fluxional process can be rationalized as a shift in the metal framework similar to that discussed above for

Molecular Dynamics of Mixed-Metal Clusters

Figure 6. ¹³C NMR spectra of H_2 FeRuOs₂(CO)₁₃.

Scheme Il

 $H_2FeRu_3(CO)_{13}$. In the C_1 isomer this process begins at 10 "C at which point the resonances due to carbonyls *0,* s, and presumably r begin to average. The resonances of carbonyls t and u do not collapse, but importantly they do become equivalent. This is readily explained by a rearrangement of the metal framework and associated ligands in which the Fe atom moves closer to Os_2 and away from Os_1 . This, in effect, generates the enantiomer of the first structure and represents a racemization process

The Fe-Ru-Os₂ triangle now possesses the bridging carbonyls. **A** cyclic process around this triangle averages *0,* s, and r with i-I. Coupling of the intrametallic rearrangement with the cyclic processes has the net effect of averaging carbonyls i-s. Carbonyls t and u, however, do not enter into this exchange. **As** illustrated in Scheme 11, this rearrangement has the effect of showing an average environment to t and u, and the observation that t and **u** become equivalent provides strong support for this mechanism.

The second effect of the intrametallic rearrangement in C_1 occurs when the Fe atom moves away from Ru and generates **Scheme I11**

the Fe- Os_1 - Os_2 triangle with the bridging CO's. This movement results in isomerization to the C_s isomer and is the final averaging process. It begins at 40 \degree C and results in the averaging of all carbonyls on both isomers. Importantly, these **13C** NMR spectra clearly indicate that racemization occurs prior to isomerization in the averaging of all carbonyls on both isomers.

 H_2 FeRu₂Os(CO)₁₃. The variable-temperature ¹³C NMR spectra of $H_2FeRu_2Os(CO)_{13}$ are shown in Figure 5. As in $H_2FeRuOs_2(CO)_{13}$, the two isomeric forms act separately in each of the exchange processes. At -70 °C, the Fe carbonyls $a-c$ have coalesced on the C_s isomer. The analogous process on the C_1 isomer occurs at -40 °C. Since it was not possible to definitively assign all the resonances in the low-temperature limiting spectrum to specific carbonyls, it is difficult to unambiguously distinguish the next processes to occur. However, the lack of changes in the Os-CO region between -40 and -70 OC and the lack of appearance of an averaged Fe-CO resonance are consistent with exchange of the carbonyls in a cyclic path around the Fe-Ru₁-Ru₂ face of the C_s isomer. Above -40 "C, peaks due to carbonyl m, n, and p coalesce with the iron carbonyl resonances. This process must involve the movement of carbonyls around the $Fe-Ru$ -Os face of the C_1 isomer. At 0 **"C,** there begins a general decrease in intensity of all resonances to give coalescence and what appears to be a single resonance growing in at 193 ppm in the **70** "C spectrum.

Discussion

CO-Exchange Mechanisms. Three distinctly different COexchange processes are resolved for these mixed-metal clusters. The first process to occur at the lowest temperatures is exchange of the bridging and terminal carbonyls bound to Fe. A reasonable mechanism for this exchange is shown in Scheme I11 and involves opening of one of the carbonyl bridges, a subsequent trigonal twist of the resultant $Fe(CO)_3$ unit, and finally re-formation of the CO bridge. A consequence of this mechanism is that the bridging CO's should decrease in intensity less rapidly than the terminal CO's since the twist must occur twice in order for exchange **of** both bridging CO's to occur. This suggestion is most strongly supported by the H₂FeRu₃(CO)₁₃ spectra shown in Figure 1 of ref 3. At -55 "C the bridging CO resonance is clearly visible while the terminal CO resonances have collapsed into the base line.

The next exchange process to occur at slightly higher temperatures involves migration of the carbonyls around the Fe-M-M triangle which possesses the bridging carbonyls. It seems reasonable to propose that the intermediates in this cyclic movement are the tautomers which have the semibridging carbonyls bound mainly to Ru or Os atoms instead

of Fe as indicated in Scheme IV. During the occurrence of this process, the hydrogens remain in their original positions. This is consistent with the observed 'H NMR spectra of $H_2FeRu_2Os(CO)_{13}$ and $H_2FeRuOs_2(CO)_{13}$ since racemization, the net effect of hydrogen movement, does not occur in the temperature range in which the 13C NMR spectra show the cyclic fluxional processes.

A number of di- and polynuclear complexes have been shown to undergo similar cyclic exchange processes. $8-12$ Cotton and co-workers $8,9$ have suggested that coplanarity is not a mandatory requirement for the ligands that participate in such a process but only that the path be continuous and closed. This requirement is certainly fulfilled in these clusters, and furthermore the crystal structure of $H_2FeRu_3(CO)_{13}^{13}$ does indicate that seven of the eight carbonyls involved are indeed virtually coplanar. The only exception is one of the terminal carbonyls attached to Fe, but by the time the cyclic process begins to occur, this carbonyl is rapidly exchanging with the bridging CO's.

The third and final process to occur is one which involves a shift in the metal framework and which accounts for racemization, isomerization and total exchange of all the carbonyl ligands. This process and its implications are best illustrated by consideration of the drawings in Scheme **V** which depict the metal framework of $H_2FeRuOs_2(CO)_{13}$, the two bridging CO's, and the two bridging hydrides. The asymmetry of the cluster is grossly exaggerated for clarity. The process basically involves movement of the iron away from one metal and closer to another with a concomitant shift of the bridging carbonyls. It also involves a slight elongation or compression of all the M-M bonds, and it must be accompanied by a shift in position of one of the bridging hydrides.

In reference to Scheme V and by starting with the C_{1a} enantiomer, if the Fe moves away from $Os₁$ toward $Os₂$, it generates the C_{1b} enantiomer. This process leads to racemization since H_b must also shift in the process. Movement of Fe away from Ru in either of the C_1 enantiomers and toward both Os atoms leads to the C_s isomer and accounts for the isomerization process. Each time the cluster rearranges, the carbonyls execute the cyclic process about a different Fe-M-M

- (8) Cotton, F. **A,;** Hanson, B. E.; Jamerson, J. D.; Stultz, B. R. *J. Am. Chem.* **SOC. 1911,** *99,* **3293.**
- (9) Cotton, **F. A,;** Haines, R. J.; Hanson, B. E.; Sekutowski, J. C. *Inorg. Chem.* **1918,** *17,* **2010. (10)** Johnson, **B. F.** *G.;* Lewis, J.; Reichert, B. E.; Schorpp, **K.** T. *J. Chem.*
- **SOC.,** *Dalton Trans.* **1916,** 1403. **(1 1)** Tachikawa, M.; Richter, S. I.; Shapley, J. R. *J. Organomet. Chem.*
- **1911,** *128,* C9. **(12)** Martinengo, S.; Heaton, B. T.; Goodfellow, R. J.; Chini, P. *J. Chem.*
- **SOC.,** *Chem. Commun.* **1977j39. (13)** Gilmore, C. **J.;** Woodward, P. *J. Chem.* **SOC.** *A* **1971,** 3453.

face and hence involve different carbonyl ligands in that process. The cyclic processes coupled with the framework rearrangement leads to total exchange of all the carbonyl ligands.

The ¹³C and ¹H NMR spectra of $H_2FeRuOs_2(CO)_{13}$ indicate that two distinct hydrogen migration processes occur for the C_1 isomer. Only one of these involves actual exchange of the two hydrogens on the C_1 isomer, and, thus, is the only one detected in the 'H NMR experiments.

The lower temperature, and hence lower activation energy, process is that specifically depicted in Scheme V which involves migration of only one hydrogen atom. The hydrogen which bridges the $Ru-Os₂$ bond (H_b) could simply move across the $Ru-Os_1-Os_2$ triangle to bridge the $Ru-Os_1$ bond. No exchange occurs with H_a which bridges the Os-Os bond, and hence this movement would not be detected in the 'H NMR spectrum. However, this process does necessitate a concerted shift in the metal framework and is thus detected by ^{13}C NMR, and it, of course, does lead to racemization, Scheme **V.**

An estimate of ΔG^* for this process can be obtained by careful examination of the behavior of carbonyls t and u in the variable-temperature NMR spectrum, Figure 6. **As** shown in Scheme **11,** carbonyls t and u become equivalent during this exchange process, but they do not exchange with any of the other carbonyls. This system can thus be treated as an isolated two-site exchange problem in which both sites are equally populated. A reasonable estimate of ΔG^* , obtained simply from the chemical shift difference of carbonyls t and u (15 Hz; 1 00-MHz instrument) and their coalesence temperature $(10 \degree C)$, is 15 kcal/mol.

The second, higher temperature, and hence higher activation energy process that is detectable by ${}^{1}H$ NMR spectroscopy is that depicted in Scheme VI. In this process both hydrogens move through an intermediate in which they are each terminally bound to the two Os atoms. This intermediate could collapse as shown in Scheme VI to give either C_{1a} or C_{1b} . Note that this process must also be accompanied by a concomitant shift in the metal framework. In this scheme, H_a and H_b exchange each time the framework shift occurs. For H_2F e- $RuOs₂(CO)₁₃, \Delta G[*]₂₈₃ = 17.8 kcal/mol for this process, com$ pared to the ΔG^*_{283} value of 15 kcal/mol for the first process.

Effect of Variations in the Metal Framework. Although within the series $H_2FeRu_3(CO)_{13}$, $H_2FeRu_2Os(CO)_{13}$, and $H_2FeRuOs_2(CO)_{13}$ the exchange processes are identical, *the activation barrier for each process increases as the osmium content of the cluster increases.* We are not able to unambiguously rationalize this trend, but a similar increase in activation energy in moving from Ru to Os has been noted for the series of monomeric $[M(P(OR)₃)₅]$ (M = Ru, Os) complexes studied by English and co-workers.¹⁴ It is unlikely that the activation energy increase can be accounted for solely on the basis of a size increase in the metal involved in the fluxional process since Ru and Os probably have similar atomic radii in these clusters. In $Ru_3(CO)_{12}$ and $Os_3(CO)_{12}$, for example, the metal atomic radii are 1.43 and 1.44 Å, respectively.¹⁵ On

 $\overline{}$

⁽¹⁴⁾ English, **A.** D.; Ittel, **S.** D.; Tolman, C. **A,;** Meakin, P.; Jesson, J. P. *J. Am. Chem.* **SOC. 1977,** *99,* **117.**

the other hand, Os certainly possesses a greater amount of electron density than does Ru, and this could be the important factor in determining the activation energies.

An interesting comparison can be made to the homonuclear cluster $H_2Ru_4(CO)_{13}$, 4, which is isostructural with the mix-

ed-metal clusters reported herein.¹⁶ The exchange processes which this cluster undergoes have been found by Milone and co-workers" to be quite similar to those of the mixed-metal analogues. However, bridge-terminal interchange localized on Ru₁ is not the lowest temperature process in $H_2Ru_4(CO)_{13}$, but instead the cyclic exchange of CO's around the Ru_1 - Ru_2-Ru_3 face occurs rapidly even at -72 °C. It was proposed that the lower activation barrier observed for $H_2Ru_4(CO)_{13}$ compared to $H_2FeRu_3(CO)_{13}$ results from the increased stability of bridging carbonyls bound to first-row metals.¹⁷ This proposal is further supported here. In Scheme IV the intermediates **2** and **3** have one of the bridging carbonyls bridging either a Ru-Ru, a Ru-Os, or an Os-Os bond. The least favorable case would involve migration of the CO's around the Fe-Os₁-Os₂ face of the C_s isomer of $H_2FeRuOs_2(CO)_{13}$. Accordingly, we find this to have the highest activation barrier observed for the cyclic process in any of the clusters. The energy difference between the tautomers **1-3** in Scheme IV is least in the case where all the metals are Ru and is greatest when one metal is Fe and the other two are Os.

- **(15)** Churchill, M. **R.;** Hollander, F. J.; Hutchinson, J. P. *Inorg. Chem. 1977, 16, 2655.*
- **(16)** Yawney, D. B. W.; Doedens, R. J. *Inorg. Chem. 1972,11,838.* (17) Aime, *S.;* Milone, L.; Osealla, D.; Sappa, E. *Inorg.* Chim. *Acra 1978, 29.* **L211.**

Intrametallic Rearrangement. The last exchange process in $H_2FeRu_3(CO)_{13}$ and the last two processes in $H_2FeRuOs_2$ - $(CO)_{13}$ and $H_2FeRu_2Os(CO)_{13}$ involve reorganization of the metal framework. The actual magnitude of the shifts within the framework is relatively small, yet it still significantly contributes to the final averaging process. The metal contributes to the final averaging process. framework of $H_2FeRu_3(CO)_{13}$ is shown below

with the average metal-metal bond distances derived from the reported crystal structure.¹³ These values indicate that the greatest change that would occur in any one bond length during the rearrangement is 0.11 **A.** We tend to view the reorganization process as more of a breathing motion of the metal framework, but one which has coupled to it motions **of** the carbonyl and hydride ligands.

It appears that intrametallic rearrangement processes are not at all uncommon, as noted by Band and Muetterties¹ in a recent review, but they are still poorly understood. In our case we would particularly like to understand how the various motions contribute to the overall rearrangement process. For example, what percentage of the 15.6-kcal/mol activation barrier for isomerization of $H_2FeRuOs_2(CO)_{13}$ is due to reorganization of the metal framework and what percentage is due to hydrogen migration? We expect that such matters will be the subject of future publications from this and other research groups.

Acknowledgment. We appreciate the assistance of Alan Freyer during the course of the NMR experiments and Professor Lloyd Jackman for helpful discussions. This research was supported in part by the Office of Naval Research. G.L.G. gratefully acknowledges the Camille and Henry Dreyfus Foundation for a Teacher-Scholar grant and the **A.** P. Sloan Foundation for the award of a research fellowship.

Registry No. $H_2Feku_3(CO)_{13}$, 12375-24-5; $H_2FeRu_2Os(CO)_{13}$, 61863-84-1; $H_2FeRuOs_2(CO)_{13}$, 61816-94-2.

Contribution from the Department of Chemistry, Tulane University, New Orleans, Louisiana 701 18

Ligand Substitution Processes in Tetranuclear Metal Carbonyl Clusters. 2. $Co_4(CO)_9(\mu$ -CO)₃ Derivatives

DONALD J. DARENSBOURG* and MICHAEL J. INCORVIA

Received July 26, *1979*

Kinetic and mechanistic studies of carbon monoxide substitution processes in the tetranuclear metal clusters Co₄- $(CO)_{12-\pi}[P(OCH_3)_3]_n$ ($n = 0, 1, 2$) are reported. Entering ligands employed in these investigations were trimethyl phosphite, triphenylphosphine, or ¹³CO. The CO substitutional rates in $Co_4(CO)_{12}$ with P(OCH₃)₃ or P(C₆H₅)₃ were too rapid to be studied by conventional-techniques and were unaffected by the addition of the radical scavenger **2,6-di-tert-butylphenol;** nevertheless, the rate of ¹³CO incorporation into Co₄(CO)₁₂ occurs at only a moderate velocity. The exchange reaction between ¹³C-labeled carbon monoxide and $Co_4(CO)_{12}$ has allowed for a detailed analysis of the v(CO) stretching vibrations for the bridging carbonyl ligands. Although the rate of reaction of $Co_4(CO)_{11}[P(OCH_3)_3]$ with $P(OCH_3)_3$ to afford $Co_4(CO)_{10}[P(OCH_3)_3]_2$ has been shown to be strongly dependent on the concentration of trimethyl phosphite, the comparable rate for formation of the mixed-ligand complex $Co_4(CO)_{10}[P(CCH_3)_3][P(C_6H_5)_3]$ from $Co_4(CO)_{11}[P(CCH_3)_3]$ and triphenylphosphine follows a first-order dissociative path. Rate constants and activation parameters for dissociative CO loss in $Co_4(CO)_{12-n}[P(OCH_3)_3]_n$ (n = 0, 1, 2) were found not to be significantly dependent upon the value of *n*.

Introduction

Muetterties has articulated the presumption that transition-metal clusters may serve as simple homogeneous models **of** metal surfaces in both chemisorption and catalytic pro $cesses.¹⁻³$ Indeed, several recent reports have appeared of homogeneous catalysis employing metal clusters as catalysts for such important processes as the water-gas shift reaction, $4-6$

(1) Muetterties, **E. L.** Bull. **SOC.** *Chim. Belg. 1975, 84, 959.* (2) Muetterties, **E. L.** *Bull.* **SOC.** *Chim. Belg. 1976, 85,* 451.