Generation and Reactivity of Unsaturated Iridium and Rhodium Dimethyl Complexes

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Protonation ($HBF_4 \cdot OEt_2$) of MMe_3P_3 ($M = Rh$, Ir; $P = PMe_2Ph$) in CH_2Cl_2 gives CH_4 and *cis,mer-* $\text{MMe}_2\text{P}_3\text{BF}_4$, characterized as a fluxional (methyl site exchange) molecular $\eta^1\text{-BF}_4$ species by ¹H, ³¹P, ¹⁹F, $a = 16.708$ (6) A, $b = 9.332$ (2) A, $c = 23.214$ (8) A, $\beta = 110.05$ (2)^o, with $Z = 4$ in space group P_{21}^2/c . Ethylene replaces BF_4^- to give (at 25 °C) $[Im\Theta_2(C_2H_4)P_3]BF_4$ and (at 50 °C) $[Ir(C_2H_4)_2P_3]BF_4$ and C_2H_6 . Carbon monoxide replaces BF₄ to give first $[IrMe₂(CO)P₃]BF₄$ and then $IrMe[C(O)Me](CO)P₃$ ⁺. Ethylene reacts (25 °C) with $\text{RhMe}_{2}\text{P}_{3}\text{BF}_{4}$ to immediately give ethane and $[\text{RhP}_{4}]\text{BF}_{4}$. Carbon monoxide replaces BF_{4}^{-} to give first $RhMe_2(\text{CO})P_3{}^+$ and then acetone and $[RhP_4]BF_4$. Ethylene also promotes acetone elimination from $\mathrm{RhMe}_2(\mathrm{CO})\mathrm{P}_3^*$, to again generate $\mathrm{RhP_4^*}$. All of the above are characterized by multinuclear NMR and vibrational spectroscopy and, for [IrMe₂(C₂H₄)P₃BF₄, X-ray diffraction. Crystal data (-155 °C): *a* $= 13.373$ (4) Å, $b = 12.242$ (3) Å, $c = 18.139$ (b) Å, with $Z = 4$ in space group $P_{1}^{2}c$. and ¹¹B NMR studies and (for M = Ir) X-ray diffraction. Crystal data for IrMe₂P₃BF₄-C₆H₆ (-100 °C):

Introduction

The creation of an open coordination site in a transition-metal complex has been a longstanding objective. This may be achieved by thermal or photoinduced dissociation of a Lewis base or by reductive elimination or by electrophilic (e.g., H^+ , R^+ , Ag^+ , Tl^+) abstraction of what is formally an anionic species (e.g., halide, H^- , or R^-). The electrophilic abstraction, since it requires a counterion, raises the question of the identity of the ideal noncoordinating anion;' only when this problem is solved will the generation of authentic unsaturation be realized. A related factor is the search for the ideal solvent, being one that will dissolve the ionic reactants and products yet will not coordinate to the transition-metal orbital liberated in the electrophilic abstraction. Acetonitrile has proven itself unsuitable in this regard,²⁻⁴ while $\mathrm{CH}_2\mathrm{Cl}_2$ may have its successes.^{5,6} There are, however, claims of coordination of $\mathrm{CH}_2\mathrm{Cl}_2$,⁷⁻⁹ and halocarbons have been demonstrated to coordinate through halogen lone pairs.¹⁰⁻¹³ Related complications arose when $[\tilde{Ph}_3C]BF_4$ was employed in an attempt to abstract H⁻ from $\text{C}_p\text{Re}(\text{NO})(\text{CO})$ H: triphenylmethane occupies a coordination site on rhenium.14

These problems notwithstanding, electrophilic abstraction has been productive of much substitution chemistry in the organometallic field. The work of the group of Beck has been outstanding in this regard, with their focus being the species $M(CO)_{5}X$ (M = Mn, Re) and CpM'L₃X (M'

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 $=$ Mo, W), where X = H, Me.^{15,16} Both Beck¹⁶ and Cutler¹³ have used the resulting "incipient Lewis acids" to attack metal acyl complexes, yielding unusual μ -acyl species. Recently, Jordan¹⁷ and others¹⁸ have used such electrophilic abstraction to generate reactive early-transition-metal alkyl cations capable of olefin polymerization. With the exception of such reports, however, there has been little work creating unsaturation in the presence of reactive (e.g., migration-prone) coligands such **as** H or CH,. We have previously been successful generating reactive unsaturated hydride species via protonation of polyhydride complexes $6,19,20$ and now turn our attention toward the production of unsaturated alkyl complexes by protonation of polyalkyl compounds. This approach allows us ready access to the area of unsaturated alkyl complexes relevant to olefin polymerization or hydroformylation. We report here on the structure and comparative reactivity of the complexes $MMe₂P₃BF₄$ (M = Rh, Ir; P = PMe₂Ph) formed by stoichiometric protonation of $fac\text{-}MMe₃P₃$.

Experimental Section

All manipulations were carried out with use of standard Schlenk and glovebox procedures under prepurified nitrogen or vacuum. Solvents (THF, benzene, toluene, pentane) were dried and deoxygenated by Na/K benzophenone and vacuum transferred prior to use. CD_2Cl_2 and CH_2Cl_2 were distilled from P_2O_5 . MeMgCl, MeLi, and $\overline{HBF}_4 \cdot OEt_2$ were purchased from Aldrich. The gases C2H4 (CP grade, Matheson), 13CzH4 (99% 13C, Cambridge Isotope Laboratories), CO (Ultra High Purity, Air Products), and *13C0* (99% 13C, Mound Research) were used as received. 'H, 31P, and 13 C NMR spectra were recorded on a Nicolet NT-360 spectrometer at 360, 146, and 100 MHz, respectively. $^{11}\rm{B}$ and $^{19}\rm{F}$ NMR spectra were also recorded on a Nicolet NT-360 spectrometer at 116 and 339 MHz. Two-dimensional NMR experiments were performed 339 MHz. Two-dimensional NMR experiments were performed on a Bruker *AM-500* spectrometer. Infrared spectra were recorded on a **Perkin-Elmer 283** spectrophotometer. Where noted, precise

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quantities of gases were determined with use of a standard calibrated gas manifold. The products reported here have been identified and characterized by X-ray diffraction and by 'H, 31P, and 13C NMR spectra. Since the solid samples dissolved completely in the NMR solvents employed, and since they are interrelated by a cycle of chemical transformations, we do not offer supplemental proof of composition by elemental analysis.

Protonation of fac-IrMe₃P₃. Generation of IrMe₂P₃BF₄. A 5-mm NMR tube was charged with $IrMe₃P₃²¹⁻²³$ (75 mg, 0.12mmol) and CD_2Cl_2 (0.5 mL), and against a N_2 flow, 1 equiv of $HBF₄·OEt₂$ (12 μ L, 0.12 mmol) was added. This caused immediate gas evolution and the formation of a yellow solution. This solution was thermochromic, becoming colorless below -20 °C. ¹H NMR spectroscopy indicated quantitative conversion to IrMe2P3BF4. **'H** NMR **(360** MHz, 25 "C, CD2C12): 6 0.50 (br q, $J_{\text{Me-P}} = 4 \text{ Hz}$, Ir(Me)₂, 6 H), 1.6 (br s, P-Me, 18 H), 7.2-7.5 (br m, P-Ph). ³¹P{¹H} (146 MHz, 19 °C, CD₂Cl₂): δ -29.0 (br s, 2 P), -44.0 (br s, 1 P). Low-temperature ¹H NMR (360 MHz, -80 °C, CD₂Cl): δ 0.40 (br m, Ir-Me, 3 H), 0.60 (br m, Ir-Me, 3 H), 1.25 (m, MeP, 6 H), 1.7 (m, Me-P, 6 H), 1.8 (m, Me-P, 6 H), 7.2-7.6 (m, P-Ph). Low-temperature ${}^{31}P{}^{1}H$ (146 MHz, -80 °C, CD₂Cl): (116 MHz, 24 °C , CD₂Cl₂) δ -1.37 (br s). (24 °C, toluene): δ -0.27 (br s). ¹⁹F NMR (339 MHz, 24 °C, CD₂Cl₂): δ -153 (sharp s). (br *8).* **lgF** NMR (339 MHz, 24 "C, CD2C12): **S** -153 (sharp *8).* 1v NMR (24 **"C,** toluene): *6* -170 **(8).** Low-temperature **lv NMR** (339 MHz, **-90** "C, CD2C12): *6* -153 (sharp **s).** Low-temperature ¹⁹F NMR (-90 °C, toluene): -185 (br s, half-width 6800 Hz). *⁶*-29.0 (d, **Jpp** = **15** Hz, 2 P), -44.0 (t, Jpp = 15 Hz, 1 P). "B *NMR*

Crystallography of $IrMe₂(PMe₂Ph)₃BF₄·C₆H₆$. A suitable crystal, grown by layering a 0.3 M benzene solution with pentane, was transferred to the goniostat with use of standard inert-atmosphere handling techniques and cooled to -100 °C for characterization and data collection.²⁴ When it was cooled, the crystal changed from pale yellow to nearly colorless. At approximately -120 **OC,** the crystals underwent a (destructive) phase transition. A systematic search of a limited hemisphere of reciprocal space located a set of diffraction maxima with symmetry and systematic absences corresponding to the unique monoclinic space group $P2₁/c$. Subsequent solution and refinement of the structure confirmed this choice. Data were collected $(6^{\circ} \leq 2\theta \leq 45^{\circ})$ with

^a Isotropic values for those atoms refined anisotropically are calculated by using the formula given by: Hamilton, W. C. Acta Crystallogr. **1959,** 12,609.

Figure **1. ORTEP** drawing of non-hydrogen atoms of IrMez- $(P\overline{M}e_2Ph)_3BF_4$, showing atom labeling.

use of a continuous θ -2 θ scan with fixed backgrounds and were reduced to a unique set of intensities and associated σ values in the usual manner.²⁴ An absorption correction was performed. Parameters of the crystal and the data are shown in Table I.

The structure was solved by a combination of direct methods (MULTAN78) and Fourier techniques and revealed a benzene molecule in the crystal lattice. A difference Fourier synthesis revealed the location of some, but not all, hydrogen atoms. All hydrogen atom positions were therefore calculated with use of

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Figure 2. Stereo stick-figure and space-filling drawings of $IrMe₂(PMe₂Ph)₃BF₄$, with $d(C-H)$ fixed at 1.05 Å.

idealized geometries and $d(C-H) = 0.95$ Å. These calculated positions were fixed for the final cycles of refinement, with the exception of those on the two methyl groups on the Ir. **A** final difference Fourier was featureless, with several peaks of intensity 1.6 e/ \AA^3 located near the metal position. Two peaks located near C(3) could be identified as possible hydrogen atoms, but no chemically reasonable **peaks** were found near C(2). For this reason, no attempt was made to include these methyl hydrogens.

The results of the structural study are shown in Tables I1 and 111 and Figures 1 and 2.

fac-RhMe₃(PMe₂Ph)₃. To a 40-mL THF solution of mer-
RhCl₃P₃²⁵ (3.0 g, 4.8 mmol; P = PMe₂Ph) was added 30 mL (45 mmol) of a 1.5 M MeLi/ $Et₂O$ solution. The yellow solution initially darkened to red-orange, which gradually faded to a pale yellow. The mixture was slurried for 2 h, cooled to 0° C, and carefully (dropwise) hydrolyzed with wet THF until gas evolution ceased. After hydrolysis, the solvent was removed under vacuum to give a pale yellow solid. Extraction with CH_2Cl_2 $(2 \times 25 \text{ mL})$ gave a yellow solution, which was evaporated under vacuum to yield an off-white solid. Washing this material with small **amounts** of toluene gave 2.0 g (3.6 mmol, 75% yield) of white $RhMe₃P₃$. 'H NMR (360 MHz, 25 "C, CD2C12): *6* -0.28 (m, **9 H),** 1.10 (d, J_{MeP} = 8 Hz, 18 H), 7.0–7.5 (m, Ph-P). ³¹P{¹H} NMR (146 MHz, 25° C, CD₂Cl₂): δ -8.5 (d, J_{PRh} = 80 Hz).

Protonation **of** fac-RhMe,P,. Generation **of** RhMezP3BF4. A 5-mm NMR tube was charged with $RhMe₃P₃$ (50 mg, 0.09) mmol) and CD_2Cl_2 (0.5 mL), and against a N_2 flow, 1 equiv of $HBF₄·OEt₂$ (9 μ L, 0.09 mmol) was added. A gas immediately evolved, giving a yellow solution. **'H** NMR spectroscopy indicated quantitative conversion to RhMe2P3BF4. 'H NMR (360 MHz, 25 °C, CD₂Cl₂): δ 0.30 (br s, Rh(Me)₂, 6 H), 1.4 (br s, P-Me, 18 H), 7.0-7.5 (m, P-Ph). $^{31}P(^{1}H)$ NMR (146 MHz, 25 °C, CD₂Cl₂): δ 2.5 (br s, 2 P), -16.0 (br s, 1 P). Low-temperature ¹H NMR (360) **MHz,** -70 "C, CDzCl2): **6** 0.25 (br m, Rh-Me, 3 H), 0.35 (br m, Rh-Me, 3 H), 1.4 (br m, P-Me, 12 H), 1.5 (br m, P-Me, 6 **H),** 7.0-7.5 (m, P-Ph). Low-temperature 31P(1HJ NMR **(146** MHz, -75 °C, CD₂Cl₂): δ 2.5 (d of d, $J_{\text{PRh}} = 110$ Hz, $J_{\text{PP}} = 17$ Hz), -16.0 (d of t, $J_{\text{PRh}} = 86 \text{ Hz}$, $J_{\text{PP}} = 17 \text{ Hz}$). ¹⁹F NMR (339 MHz, 20 °C, CD₂Cl₂): $\delta -172$ (s). ¹⁹F NMR (20 °C, toluene): $\delta -172$ (s). Low-temperature ¹⁹F NMR (339 MHz, -80 °C, CD₂Cl₂): δ -158 (sharp s). Low-temperature **l9F** NMR (-60 "C, toluene): **-174** (br s, half-width 680 Hz).

Reaction of $IrMe₂P₃BF₄$ **with** $C₂H₄$ **.** A 5-mm medium-walled NMR tube containing 70 mg (0.1 mmol) of $IrMe₂P₃BF₄$ (generated in situ) in $\mathrm{CD}_2\mathrm{Cl}_2$ was twice freeze–pump–thaw degassed and then pressurized with 2 atm of C2H4 at **25** "C. When the mixture **was** warmed to 25 °C, it turned yellow and then bleached. ¹H NMR spectroscopy indicated quantitative formation of cis,mer- $IrMe₂(C₂H₄)P₃BF₄$. Standing overnight results in formation of large colorless crystals in high yield. In solution, [IrMe₂- $(C_2H_4)P_3$]BF₄ loses C_2H_4 in the absence of excess ethylene, regenerating IrMe₂P₃BF₄. ¹H NMR (360 MHz, 0 °C, CD₂Cl₂): δ -0.14 (br d of t, $J_{\text{ImMe-P}} = J_{\text{ImMe-P}} = 7$ Hz, 3 H), 0.85 (d of t, $J_{\text{ImMe-P}}$ $= J_{ImMe-P} = 8 Hz, 3 H$, 1.60 (overlapping m, P-Me, 18 H), 2.85 (s, **CzH4,** 4 H), 7.0-7.6 (m, P-Ph). 31P(1H) NMR (146 MHz, -20 1 P). ${}^{13}C_1{}^{1}H$ } NMR for $[IrMe_2({}^{13}C_2H_4)P_3]BF_4$ (146 MHz, 20 °C, CD₂Cl₂): δ 64 (br s, IrC₂H₄). $^{\circ}$ C, CD₂Cl): δ -41.5 (d, $J_{PP} = 15$ Hz, 2 P), -57.0 (t, $J_{PP} = 15$ Hz,

Crystallography of $[IrMe₂(C₂H₄)(PMe₂Ph)₃]BF₄$ **.** A suitable crystal, grown as described above, was transferred to the

⁽²⁵⁾ Chatt, J.; Johnson. N. P.; Shaw, B. **I,.** *J. Chem. SOC.* **1964,** 2508.

Figure 3. NOESY ¹H NMR spectrum $(500 \text{ MHz}, 25 \text{ °C}, CD_2Cl_2)$ of $\text{Tr}\text{Me}_2(\text{C}_2\text{H}_4)(\text{PMe}_2\text{Ph})_3$ ⁺ in the presence of C_2H_4 . Cross peaks (off-diagonal) show site exchange of free with coordinated C_2H_4 as well as between the two IrMe sites (\ddagger indicates Et₂O, and * indicates silicone grease).

goniostat with use of standard inert-atmosphere handling techniques and cooled to **-155** "C for characterization (Table I) and data collection.

A systematic search of a limited hemisphere of reciprocal space located a set of diffraction maxima with symmetry and systematic absences corresponding to one of the orthorhombic space groups *Pmcn* and $P2_1$ cn. Subsequent solution and refinement of the structure confirmed the noncentrosymmetric choice, $P2_1$ cn.

Data were collected $(6^{\circ} \leq 2\theta \leq 45^{\circ})$, with use of a continuous $0-2\theta$ scan with fixed backgrounds, and reduced to a unique set of intensities and associated σ values in the usual manner.²⁴ The structure was solved by a combination of direct methods **(MULTAMB)** and Fourier techniques. A difference Fourier synthesis revealed the location of some, but not all, hydrogen atoms. All hydrogen atom positions were therefore calculated by using idealized geometries and d (C-H) = 0.95 Å. These calculated positions were fixed for the final cycles of refinement.

An interesting dilemma occurs in that the molecule (including methyl and phenyl groups) possesses nearly perfect mirror symmetry. A careful examination reveals that the chosen space group is undoubtedly correct, but packing diagrams reveal an interesting

Table IV. Fractional Coordinates and Isotropic Thermal $\textbf{Parameters}^a$ for $[\textbf{IrMe}_{2}(\text{C}_{2}\text{H}_{4})(\text{PMe}_{2}\text{Ph})_{3}]\text{BF}_{4}$

| | | r arameters" for \lfloor irme ₂ (C_2 fi ₄)(r me ₂ r n) ₃]Br ₄ | | |
|-----------------|-------------------|--|-------------------|--|
| | 10 ⁴ x | 10 ⁴ | 10 ⁴ z | $10B_{\text{iso}}$, $\overline{\mathbf{A}^2}$ |
| Ir1 | $-8562b$ | $-7571(1)$ | $-9795.4(4)$ | 11 |
| P ₂ | $-8633(8)$ | $-7422(4)$ | $-8477(2)$ | 10(1) |
| C ₃ | $-8836(19)$ | $-8658(21)$ | $-7981(15)$ | 24(6) |
| C4 | $-7424(20)$ | $-7013(21)$ | $-8027(14)$ | 14(5) |
| C ₅ | $-9474(20)$ | $-6480(22)$ | $-8062(15)$ | 16(5) |
| C6 | $-9291(20)$ | $-5426(21)$ | $-7935(14)$ | 17(5) |
| C7 | $-9974(18)$ | $-4658(18)$ | $-7686(13)$ | 8(4) |
| C8 | $-10985(33)$ | $-5017(26)$ | $-7517(22)$ | 19(5) |
| C9 | $-11158(27)$ | $-6106(27)$ | $-7686(18)$ | 31 (6) |
| C10 | $-10457(20)$ | $-6817(21)$ | $-7922(15)$ | 18(5) |
| P11 | $-9753(7)$ | $-8998(6)$ | $-9880(4)$ | 15(1) |
| C12 | $-10317(28)$ | $-9189(27)$ | $-10782(18)$ | 39 (7) |
| C13 | $-9296(23)$ | $-10371(23)$ | $-9684(16)$ | 25(6) |
| C14 | $-10922(18)$ | $-8943(18)$ | $-9330(13)$ | 12(4) |
| C15 | $-11032(20)$ | $-9608(20)$ | $-8739(14)$ | 17(5) |
| C16 | $-11934(21)$ | $-9515(19)$ | $-8289(15)$ | 15(5) |
| C17 | $-12629(19)$ | $-8696(19)$ | $-8504(13)$ | 14(5) |
| C18 | $-12492(22)$ | $-8052(21)$ | $-9072(15)$ | 22(5) |
| C19 | $-11622(21)$ | $-8175(20)$ | $-9491(15)$ | 20(5) |
| P ₂₀ | $-7341(7)$ | $-6191(6)$ | $-9979(5)$ | 13(1) |
| C ₂₁ | $-6049(25)$ | $-6528(24)$ | $-9795(17)$ | 33(6) |
| C ₂₂ | $-7166(27)$ | $-5678(25)$ | $-10897(18)$ | 35(6) |
| C ₂₃ | $-7538(22)$ | $-4880(21)$ | $-9480(15)$ | 21(5) |
| C ₂₄ | $-8286(19)$ | $-4193(18)$ | $-9655(13)$ | 20(5) |
| C ₂₅ | $-8415(31)$ | $-3204(19)$ | $-9280(14)$ | 32(6) |
| C ₂₆ | $-7791(21)$ | $-2947(21)$ | $-8738(16)$ | 24(5) |
| C27 | $-6980(22)$ | $-3595(22)$ | $-8537(15)$ | 26(5) |
| C ₂₈ | $-6852(19)$ | $-4572(19)$ | $-8944(14)$ | 15(5) |
| C ₂₉ | $-9817(32)$ | $-6338(30)$ | $-9672(22)$ | 41(7) |
| C ₃₀ | $-9594(21)$ | $-6502(22)$ | $-10423(15)$ | 20(5) |
| C31 | $-8130(17)$ | $-8082(18)$ | $-10881(13)$ | 11(4) |
| C32 | $-7422(26)$ | $-8733(23)$ | -9557 (17) | 24 (6) |
| B33 | 5447 (29) | 3681 (30) | 2083 (20) | 29(7) |
| F34 | 6234 (13) | 3675 (18) | 2511 (18) | 90 |
| F35 | 5244 (14) | 2701 (14) | 1767(11) | 46 |
| F36 | 4677 (19) | 4242 (28) | 2275 (17) | 107 |
| F37 | 5704 (37) | 4331 (30) | 1532 (20) | 155 |
| | | | | |

^{*a*} See footnote *a* of Table II. ^{*b*} Not varied.

Table V. Selected Bond Distances **(A)** and Angles (deg) for

| $\frac{1}{2}$ | | | | | | |
|---------------|-------------|----------------------------------|------------|--|--|--|
| $Ir1-P2$ | 2.400 (5) | $Ir1-C32$ | 2.13(3) | | | |
| $Ir1-P11$ | 2.368 (8) | F34-B33 | 1.31(4) | | | |
| Ir1–P20 | 2.373(8) | F ₃₅ -B ₃₃ | 1.36(4) | | | |
| $Ir1-C29$ | 2.27(4) | F36–B33 | 1.29(4) | | | |
| $Ir1-C30$ | 2.216(27) | F37-B33 | 1.32(5) | | | |
| $Ir1-C31$ | 2.146 (23) | $C29-C30$ | 1.41(5) | | | |
| P2-Ir1-P11 | 95.39 (28) | C29-Ir1-C30 | 36.6 (11) | | | |
| $P2-Ir1-P20$ | 96.5(3) | C29-Ir1-C31 | 118.9 (12) | | | |
| P2-Ir1-C29 | 79.8 (10) | $C29-Ir1-C32$ | 162.6 (11) | | | |
| $P2-Ir1-C30$ | 116.3(8) | C30–Ir1–C31 | 82.4 (10) | | | |
| P2–Ir1–C31 | 161.3 (6) | C30–Ir1–C32 | 160.8 (11) | | | |
| P2–Ir1–C32 | 82.9(9) | C31-Ir1-C32 | 78.4 (11) | | | |
| P11–Ir1–P20 | 168.11 (23) | Ir1–C29–C30 | 69.7 (19) | | | |
| $P11-Ir1-C29$ | 90.0(10) | Ir1–C30–C29 . | 73.7 (19) | | | |
| $P11-Ir1-C30$ | 89.1 (7) | F34–B33–F35 | 114.(3) | | | |
| P11-Ir1-C31 | 84.6 (6) | F34-B33-F36 | 119. (3) | | | |
| P11-Ir1-C32 | 90.1(8) | F34-B33-F37 | 104. (4) | | | |
| P20–Ir1–C29 | 92.8 (10) | F35-B33-F36 | 115. (3) | | | |
| P20–Ir1–C30 | 86.3 (7) | F35-B33-F37 | 105. (3) | | | |
| P20–Ir1–C31 | 83.9 (6) | F36-B33-F37 | 95. (4) | | | |
| P20–Ir1–C32 | 90.6 (9) | | | | | |

pseudosymmetry, with the molecules aligned along the diagonals of the proper cell. It is possible that the crystal lies near a phase transition to a different space group. The BF_4 ions, however, are well-behaved and do not lie on the pseudosymmetry present.

The results of the structure determination are shown in Tables IV and **V** and Figure 4.

Reaction of $[IrMe₂(¹³C₂H₄)P₃]BF₄ with Ethylene. A 5-mm$ medium-walled NMR tube containing 0.046 mmol of IrMe₂-(¹³C₂H₄)P₃⁺ (generated in situ) in CD₂Cl₂ under 2 atm of ¹³C₂H₄ was heated in a 50 °C oil bath for 12 h. ¹H NMR spectroscopy revealed the formation of Ir(¹³C₂H₄)₂P₃]BF₄⁶ and ¹²C₂H₆

Figure 4. ORTEP drawing of non-hydrogen atoms in IrMe₂- $(C_2H_4)(PMe_2Ph)_3^+$, showing atom labeling.

ppm). The ${}^{13}C{}_{1}{}^{1}H{}_{1}$ NMR spectrum confirmed that no other labeled products were present.

Reaction of $IrMe₂P₃BF₄$ **with CO (1 atm):** $[IrMe₂$ $(CO)P_3$]BF₄. To a degassed 10-mL CH_2Cl_2 solution of $IrMe₂P₃BF₄$ (150 mg, 0.21 mmol), generated in situ, was added excess CO, but at 1 atm. The yellow solution immediately bleached and was stirred for **2** h. Removal of solvent and excess CO under vacuum afforded a white powder. This powder was washed with THF (only slight solubility) and **collected** by fitration to yield 150 mg (95% yield) of cis, mer -[IrMe₂(CO)P₃]BF₄. ¹H NMR (360 MHz, 25 °C, CD₂Cl₂): δ 0.22 (d of t, $J_{\text{IMe-P}} = J_{\text{IMe-P}}$ = 9 Hz, 3 H), 0.26 (d of t, $J_{I \text{rMe-P}} = 4.5$ Hz, $J_{I \text{rMe-P}} = 9$ Hz, 3 H), 1.35 (d, $J_{\text{MeP}} = 9$ Hz, 6 H), 1.60 (vt, $J_{\text{MeP}} = 4$ Hz, 6 H), 1.70 (vt, J_{MeP} = 4 Hz, 6 H), 7.0–7.65 (m, P–Ph). ³¹P{¹H} NMR (146 MHz, 25° C, CD₂Cl₂): δ -40.4 (d, $J_{PP} = 15.5$ Hz, 2 P), -52.4 (t, $J_{PP} =$ 15.5 Hz, 1 P). IR (Nujol mull, ν_{CO}): 2020 cm⁻¹. ¹³C{¹H} NMR for IrMe₂(¹³CO)P₃⁺ (100 MHz, 25 °C, CD₂Cl₂): *δ* 172 (d of t, $J_{\rm ^{13}CO-P}$ $= J_{13_{\rm CO-P'}} = 7$ Hz).

Reaction of $[IrMe₂(CO)P₃]BF₄$ with Excess CO: [IrMe-(COMe)P3]BF4. **A** 5-mm medium-walled NMR tube containing 30 mg (0.04 mmol) of the BF_4^- salt of $IrMe₂(CO)P_3^+$ in CD_2Cl_2 was degassed and then pressurized $({\sim}3$ atm) with excess CO. After 4 h, 'H NMR spectroscopy of this sealed tube indicates a 20% conversion to IrMe(COMe)(CO) P_3^+ . No further conversion occurs even after 12 h. ¹H NMR (360 MHz, 20 °C, CD₂Cl₂): δ 0.26 (d of t, $J_{\text{Im}} = J_{\text{Im}} = 7$ Hz, 3 H), 1.55 (d, $J_{\text{Me-P}} = 9$ Hz, 6 H), 1.57 (vt, $J_{\text{MeP}} = 4$ Hz, 6 H), 1.84 $^{\circ}$ C, CD₂Cl₂): δ -36.0 (d, J_{PP} = 16.8 Hz, 2 P), -50.5 (t, J_{PP} = 16.8 (s, COMe, 3 H), 7.0-8.0 (m, P-Ph). 31P(1H) NMR (146 MHz, 22 Hz, 1 P). IR (Nujol mull): 2015 s, 1600 s cm⁻¹. ¹³C(¹H) NMR for IrMe(¹³COMe)(¹³CO)P₃⁺ (100 MHz, 22 °C, CD₂Cl₂): δ 172 (d 231 (d of m, J_{13} C_{OM} $_{2}$ ¹³C_O = 33 Hz). of t of d, $J_{^{13}CO-^{13}COMe}$ = 33 Hz, $J_{^{13}CO-P}$ = 8.5 Hz, $J_{^{13}CO-P}$ = 4 Hz),

Reaction of $\widehat{RhMe}_2P_3BF_4$ with C_2H_4 . A 5-mm mediumwalled NMR tube containing 60 mg (0.11 mmol) of $RhMe₂P₃BF₄$ (generated in situ) in CD_2Cl_2 (0.5 mL) was degassed and then pressurized with 2 atm of C_2H_4 . After the tube was sealed and the mixture warmed to 25 "C, the yellow solution darkened to yellow-orange. 'H NMR spectroscopy revealed the presence of ethane (0.85 ppm), ethylene (5.25 ppm), $\text{[Rh(PMe}_{2}\text{Ph})_{4}\text{]BF}_{4}^{26}$ and a small amount of PMezPh.

Reaction **of** RhMeP3BF4 with CO in **THF. A** 5-mm NMR tube containing 75 mg (0.14 mmol) of $RhMe₂P₃BF₄$ (generated in situ) in THF (0.6 mL) was degassed and then pressurized with excess CO at -196 "C. The tube was sealed and warmed to **25** "C; the yellow solution pales and then deposits colorless crystals. The tube was kept at -20 °C overnight to afford additional crystals along with a white powder. 'H NMR spectroscopy confirms that both the crystals and powder are **cis,mer-[RhMez(CO)P31BF,.** Evacuating a solution of $\mathrm{RhMe}_2(\mathrm{CO})\mathrm{P_3^+}$ regenerates $\mathrm{RhMe}_2\mathrm{P_3BF_4};$ in solution under N_2 , slow decarbonylation also occurs. ¹H NMR (360 MHz, 20 °C, CD₂Cl₂): δ 0.20 (m, RhMe₂, 6 H), 1.20 (d, J_{MeP} 6 H), 7.0-7.5 (m, P-Ph). 31P{1H) NMR (146 MHz, **20** "C, **CD2C12):** = 9 Hz, 6 H), 1.40 **(vt,** JMep = 4 Hz, 6 **H),** 1.60 (vt, JMep = 4 Hz,

 $=76$ Hz, $J_{PP} = 23$ Hz, 1 P). IR (Nujol mull): 2055 cm⁻¹. ¹³C(¹H) NMR for $\text{[RhMe}_{2}^{(13}\text{CO)}\text{P}_{3}\text{]}BF_{4}$ (100 MHz, 20 °C, CD₂Cl₂): δ 188 δ -1.0 (d of d, J_{PRh} = 98 Hz, J_{PP} = 23 Hz, 2 P) -17.0 (d of t, J_{PRh} (d of d of t, $J_{\text{13CO-Rh}} = 45 \text{ Hz}$, $J_{\text{13CO-P}} = J_{\text{13CO-P'}} = 9 \text{ Hz}$).

Reaction of $\overrightarrow{[Rm}Me_2(CO)P_3]\overrightarrow{BF}_4$ with CO in CH_2Cl_2 . A 5-mm medium-walled NMR tube containing 30 mg (0.45 mmol) of $RhMe_2(CO)P_3^+$ in CD_2Cl_2 was degassed and then pressurized with excess CO. After 2 h, the colorless solution had turned yellow-orange and 'H NMR spectroscopy revealed the presence of only acetone (2.00 ppm) and $[Rh(PMe₂Ph)₄]BF₄$. Identification as acetone was confirmed by rerecording the 'H NMR spectrum after addition of acetone.

Reaction of $\left[\text{RhMe}_2(\text{CO})\text{P}_3\right]\text{BF}_4$ **with** C_2H_4 **.** A 5-mm medium-walled NMR tube containing 28 mg (0.04 mmol) of $RhMe₂(CO)P₃⁺$ was degassed and then pressurized with 2 atm of ethylene. Monitoring ('H NMR) this reaction over a **24-h** period revealed the gradual disappearance of signals for $RhMe₂(CO)P₃⁺$ with the appearance of acetone and $[\bar{R}h(PMe₂Ph)₄]\bar{B}F₄$ (also identified by ${}^{31}P$ NMR spectroscopy and X-ray diffraction²⁶).

Results

Synthesis of $\mathbf{MMe}_{3}(\mathbf{PMe}_{2}\mathbf{Ph})_{3}$ **(** $\mathbf{M} = \mathbf{Ir}, \mathbf{Rh}$ **).** The polyalkyl complex $IrM\acute{e}_3P_3$ (P = PEt₃, PEt₂Ph) was first prepared by Chatt and Shaw²¹ by refluxing a solution of $mer-IrCl₃P₃$ with an excess of MeMgCl in benzene. This method produces the facial trimethyl isomer in about **70%** yield. It is reported^{25,27} that attempts to produce the Rh analogue with this same procedure failed to produce any RhMe₃L₃ (L = PR₃). Wilkinson et al.²⁸ subsequently were successful in preparing $RhMe₃(PMe₃)₃$ in low yield by the reaction of $Rh_2(O_2CCH_3)_4$ with $MgMe_2$ in the presence of excess PMe₃ at low temperatures. In contrast to earlier reports, we did obtain \overline{RhMe}_3P_3 ($P \equiv PMe_2Ph$) from the reaction of $mer-RhCl₃P₃$ with MeMgCl in benzene, although in low (28%) yield. The major product in this reaction is the easily separated complex *cis,mer-* $RhClMe₂P₃$. $RhMe₃P₃$ can, however, be prepared in good yield from the reaction of mer-RhCl₃P₃²⁵ with excess MeLi in THF. The facial stereochemistry of this product is easily deduced from the single resonance in the $^{31}P(^{1}H)$ NMR spectrum and the second-order multiplet in the 'H NMR spectrum due to the magnetic inequivalence of the rhodium methyl groups.

Protonation of $MMe₃P₃$ ($M = Ir$, Rh). Generation of MMe₂P₃BF₄ and Structure of IrMe₂P₃BF₄. Treatment of fac-IrMe₃P₃ with equimolar $HBF_4 \cdot OEt_2$ in CD_2Cl_2 or C_6D_6 at room temperature gives immediate gas evolution $(CH₄$ by ¹H NMR). The ³¹P^{{1}H} NMR spectrum of the resulting complex at **20** "C consists of two broad resonances, which sharpen to an AB_2 pattern upon cooling to -80 °C. The ¹H NMR spectrum $(CD₂Cl₂)$ is also temperature-dependent with a single broad IrMe resonance at 0.50 ppm $(20 °C)$ splitting into two separate methyl signals at -80 °C. Similar behavior is observed in both the ³¹P and ¹H NMR spectra of RhMe₂P₃BF₄ produced by stoichiometric protonation of $RhMe₃P₃$ in CD₂Cl₂. Remarkably, both complexes show good solubility in nonpolar solvents such as benzene and toluene.

These observations do not discriminate between a 16e, five-coordinate $IrMe₂P₃⁺$ structure and an 18e six-coordinate structure with an agostic methyl group or with coordinated CH_2Cl_2 . Moreover, these data and the benzene solubility, in particular, suggest possible coordination of BF₄. When pentane is layered into a concentrated benzene solution of $IrMe₂P₃BF₄$, there occurs significant conversion to new materials (by 31P NMR) con-

⁽²⁶⁾ Lmdquist, E. **G.;** Streib, W. E.; Caulton, K. G. *Inorg. Chim. Acta* **1989,** *159,* **23.**

⁽²⁷⁾ Chatt, J.; Underhill, A. E. J. Chem. Soc. 1963, 2088.

⁽²⁸⁾ Andersen, R. A.; Jones, R. A.; Wilkinson, G. *J.* Chem. *Soc., Dalton Trans.* **1978, 446.**

current with deposition of single crystals of $IrMe₂P₃BF₄$. The X-ray diffraction study (Figure 1) reveals this solid to be comprised of the molecular species $cis,mer-IrMe₂$ - $(PMe₂Ph)₃BF₄$, where the octahedral Ir(III) has inequivalent methyl groups, one of which is trans to η^1 -coordinated BF_{4} .

All Ir-P distances are statistically identical, but the two mutually trans phosphines bend away from P18 toward the methyl of C2. That methyl group has the longer of the two Ir-C bonds (by some 4σ), the shorter Ir-C3 being trans to the weakly bonded η^1 -BF₄. There is evidence of graphitic (face-to-face) stacking of the phenyl rings on P18 and P27, and the rotational conformation of the bonds from iridium to P9 and P27 is such as to interleave the attached four P-methyl groups with the $IrMe₂$ group (Figure 2).

There is evidence for distortion of BF_4 by coordination. The $(\mu-F)-B$ distance is longer than the B-F(terminal) bonds by some 4σ , and the angles (μ -F)-B-F are consistently smaller $(106.6 (12)-108.6 (12)°)$ than the angles between the terminal fluorines $(109.5 (13)-113.0 (13)°)$. The angle at the bridging fluoride is distinctly nonlinear (159.5 (9) ^o), being bent away from the bulky phosphines and toward C2.

Reactivity of $IrMe₂P₃BF₄$ **with** $C₂H₄$ **.** We sought to explore the synthetic utility of $Me_2Ir\bar{P}_3BF_4$ and for this purpose employed the unactivated olefin C_2H_4 . The complex in either benzene or CH_2Cl_2 reacts rapidly with C_2H_4 to produce $[IrMe₂(C₂H₄)P₃]BF₄$. The ¹H and ³¹P NMR spectra in $\mathit{CD}_2\mathit{Cl}_2$ at 0 and -20 °C, respectively, are consistent with a cis,mer geometry in the cation but, at 25 $^{\circ}$ C and **500** MHz, the 'H NMR spectrum shows broad lines for each Ir-Me group and for coordinated and free C_2H_4 (added to prevent conversion to $Me₂IrP₃BF₄$). NOESY two-dimensional 'H NMR spectra studies show cross peaks (Figure 3) linking free and coordinated ethylene, as well as linking the two inequivalent Ir-Me peaks. These results are consistent with the equilibrium in eq 1, which represents a single (dissociative) mechanism to accomplish both site exchanges, via participation by the 16-electron transient $IrMe₂P₃⁺$.

In an attempt to promote the insertion of ethylene into the cis Ir–Me bond, a CD_2Cl_2 solution containing IrMe₂- $(^{13}C_2H_4)P_3$ ⁺ under 2 atm of $^{13}C_2H_4$ was held at 50 °C for 12 h. 'H NMR spectroscopy indicated the presence of ethane and the formation of the bis(olefin) complex [Ir- $(^{13}C_2H_4)_2P_3|BF_4.$ ⁶ $~^{13}C(^{1}H)$ NMR spectroscopy confirmed the presence of the bis(olefin) complex and revealed (besides ${}^{13}C_2H_4$) no other ${}^{13}C$ -containing products.

An X-ray diffraction study of $[IrMe₂(C₂H₄)$ - $(PMe₂Ph)₃BF₄$ reveals it to be comprised of noninteracting BF₄⁻ anions and cis,mer cations (Figure 4). The coordination geometry is approximately octahedral, with some distortions related to the larger (C_2H_4) and smaller (CH_3) ligands. Thus, the cis P-Ir-P angles are 95-97° and the cis $H_3C-Ir-CH_3$ and $H_3C-Ir-P2$ angles are 78.4 and 82.9", respectively. Atoms P11 and P20 likewise bend toward methyl carbon C31, as seen in $IrMe₂P₃BF₄$. The Ir-P2 bond, which is trans to methyl carbon C31, is elongated by 3σ (difference) relative to the other two Ir-P bonds. Similarly, the **1r-C** distances to ethylene (2.22 (3) and 2.27 (4) **A)** are considerably longer than those (2.14

(2)-2.17 (2) **A)** in the closely related Ir(1) complex Ir- $(C_2H_4)_2(PMe_2Ph)_3^+$, where there is no methyl group trans to ethylene. Thus, there is structural evidence for a trans bond weakening, which manifests itself in the solution ligand dissociation described by eq 1.

Reactivity of $IrMe₂P₃BF₄$ **with CO.** $IrMe₂P₃BF₄$ **re**acts rapidly with carbon monoxide in either benzene or methylene chloride to produce the dialkyl carbonyl complex cis,mer -[IrMe₂(CO)P₃]BF₄. ¹H and ³¹P NMR spectroscopy as well as ¹³C^{{1}H} NMR spectroscopy (obtained with Ir $\text{Me}_2(\text{^{13}CO})\text{P}_3^+$, confirm this static octahedral arrangement. The solution infrared spectrum is also consistent with this formulation, showing a single strong absorbance at 2062 cm⁻¹. Unlike IrMe₂(C₂H₄)P₃⁺, IrMe₂- $(CO)P_3$ ⁺ is resistant to ligand (carbonyl) loss even under vacuum.

Placing IrMe₂(CO)P₃⁺ under additional CO pressure (>2 atm) gives partial conversion to the acyl complex mer- $[IrMe(COME)(CO)P₃]BF₄ (A)$. The ¹H NMR spectrum

shows only one IrMe resonance, **as** well **as** a singlet at 1.84 ppm for the acyl methyl. The solid-state infrared **spectrum** of this equilibrium mixture shows, in addition to absorbances for $IrMe₂(CO)P₃⁺, bands at 2015 and 1600 cm⁻¹ for$ the carbonyl and acyl moieties, respectively. Reacting IrMe₂P₃BF₄ with excess ¹³CO produces IrMe₂(¹³CO)P₃⁺ and the doubly labeled cation \mathbf{IrM} e $(^{13}\text{COM}$ e $)^{13}\text{CO}$) $\mathrm{P_{3}}^{+}$. The $^{13}C(^{1}H)$ NMR spectrum reveals a characteristic downfield shift for the acyl group (231 vs 172 ppm for Ir- $13CO$) with strong (33 Hz) coupling to the trans $13C$ labeled carbonyl. In the 'H NMR spectrum, the acyl methyl is split into a doublet **(5** Hz) by the 13C label. The acyl complex spontaneously decarbonylates in the absence of free CO to reform the dimethyl carbonyl complex. Attempts to drive this carbonyl insertion further with use of added ligands other than CO, such as $CH₃CN$ and PMe2Ph, were unsuccessful.

Reactivity **of** RhMe2P3BF4. Hoping that the insertion of ethylene into a Rh-C bond would be more favorable than with iridium, we investigated the reactivity of $RhMe₂P₃BF₄$ with ethylene. $RhMe₂P₃BF₄$ in $CH₂Cl₂$ reacts with excess ethylene at room temperature to produce ethane and $[Rh(PMe₂Ph)₄]BF₄$. We have previously determined²⁶ that the RhP_4 ⁺ product in this reaction results from phosphine redistribution of the unsaturated transient species RhP_3^+ . Monitoring (by ¹H NMR) the reaction between $RhMe₂P₃BF₄$ and ethylene at low temperature shows no evidence for the formation of the compound $[RhMe₂(C₂H₄)P₃]BF₄.$

In THF, $RhMe₂P₃BF₄$ reacts with carbon monoxide to deposit colorless crystals of cis,mer - [RhMe₂(CO)P₃]BF₄. This complex possesses spectral properties similar to those of the iridium analogue. Unlike the iridium compound, $RhMe₂(CO)P₃⁺$ in solution, under vacuum, loses CO to regenerate RhMe₂P₃BF₄. Tetrahydrofuran is thus an especially favorable solvent for preparing $[RhMe₂(CO)P₃]$ -BF4, since the compound precipitates spontaneously from this solvent. RhMe₂(CO)P₃⁺ dissolved in CH₂Cl₂ reacts with added CO to form acetone and a Rh-containing product (eq 2). The rhodium-containing final product is, as found in the reaction of ethylene with $RhMe₂P₃BF₄$, the

$$
cis, mer-RhMe_2(CO)P_3^+ \xrightarrow{L = CO, C_2H_4} (CH_3)_2CO + RhP_4^+ + ... (2)
$$

redistribution product of the RhP_3 ⁺ species: RhP_4 ^{+,26} Nucleophiles other than CO will also promote the formation of acetone and $RhP₄⁺$. When $RhMe₂(CO)P₃⁺$ in CD_2Cl_2 is exposed to an ethylene atmosphere, slow $(t_{1/2})$ $= 12$ h) reductive elimination of acetone and the formation of $RhP₄⁺$ are observed.

Discussion

The overwhelming majority of apparent 16-electron complexes of the "middle" transition elements contain potential π -donor ligands (halogen, OR, SR, NO, etc.). It is thus quite exceptional to find isolable unsaturated complexes containing only phosphine and alkyl or hydride ligands. The 16-valence-electron complexes $MMe₂P₃⁺$ (M = Rh, Ir), produced by protonation of MMe_3P_3 with HBF_4 , are in accord with this general situation. In the solid state, BF_4 is η^1 -coordinated, forming cis, mer-IrMe₂(FBF₃)P₃ and affording the Ir(III) d^6 center a preferred octahedral environment. The distortion of $BF₄$ upon coordination is similar to that found in $Cu(PPh_3)_3FBF_3$ and trans-IrH- $(PPh₃)₂Cl(CO)(FBF₃)$ with an elongation of the $(\mu$ -F)-B distance over that of the B-F(termina1) bonds. The B-F-Ir angle of 159.5" is, however, markedly more linear than the M-F-B angle found for trans-IrH(PPh₃)₂Cl(CO)- $(FBF₃)$ (125.7°),¹⁶ probably as a result of steric repulsions between $FBF₃$ and the three phosphine ligands. The Ir-F distance of 2.39 *8,* is over 0.1 **8,** longer than in IrH- (PPh_3) , Cl(CO)(FBF₃), showing the weakness of this interaction.

In CH₂Cl₂ solvent, the ¹⁹F NMR spectrum of $IrMe₂P₃BF₄$ shows a single line at the chemical shift of free BF_4 . This chemical shift is unchanged down to -90 °C. These observations suggest essentially complete separation of cation and anion in CH_2Cl_2 ; we cannot state whether the cation is 18-electron $Ir\tilde{M}e_2(CH_2Cl_2)P_3^+$ or 16-electron IrMe₂ P_3 ⁺. One additional subtle feature demands interpretation: the ³¹P{¹H} resonance of IrMe₂P₃BF₄ in CH₂Cl₂ shows well defined P/P coupling at -90 °C, but this is unresolved (due to broad lines) at 19 °C. Similarly, the Me-P ¹H NMR signals are broad at 25 $^{\circ}$ C. We interpret this as indicating the coexistence of two distinct species at 19 °C; these interconvert at an intermediate exchange rate. One of these could be a small fraction of the η^1 -BF₄ species seen in the solid state.

In contrast, in benzene or toluene solvent, there is spectral evidence (in addition to solubility behavior) for retention of η^1 -BF₄. First, the ³¹P chemical shifts of Ir $Me₂P₃BF₄$ in toluene are 6-10 ppm downfield of their values in CH_2Cl_2 . When the temperature in toluene is lowered, a broadening of the 19F signal is observed for IrMe₂P₃BF₄. At -90° C, the resonance has shifted upfield by **15** ppm to -185 ppm and has a width at half-height of 6800 Hz. For comparison, Beck has obtained low-temperature ¹⁹F NMR data (CD_2Cl_2) in which separate resonances for the bridging and terminal fluorides are observed.¹⁵ For instance, the complex $Cp(CO)_{3}MoFBF_{3}$ at -80 **"C** shows a doublet at -156 ppm for the terminal fluorides and a quartet at -372 ppm for the bridging fluoride. However, in CpFe(CO)₂($\overline{\eta}$ ¹-BF₄), bridging and terminal F differ by only 13 ppm.¹³ While separate ¹⁹F signals for $IrMe₂P₃FBF₃$ in toluene are not frozen out at -90 °C, a definite BF₄ interaction is inferred from the extremely broad nature of the 19F resonance. Although this broadening is consistent with a terminal-to-bridging fluoride-exchange process that has a low energy barrier, the fact that this average chemical shift is altered upon cooling necessitates the simultaneous occurrence of an equilibrium process. An equilibrium between a coordinated BF_4 and a solvent-separated BF_4 would account for this observation.

The proposed coordination of a ligand as poor as CH_2Cl_2 demonstrates the high Lewis acidity of a 16-electron complex devoid of π -donor ligands. Nevertheless, authentic 16-electron IrMe₂ P_3 ⁺ is of central importance as a transient in the fluxionality (eq 1) (unusual for a $d⁶$ octahedron) and reactivity of $IrMe₂P₃BF₄$.

The coordinated BF_4^- is easily replaced by ethylene, forming **cis,mer-[IrMe2(C2H4)P3JBF4.** A NOESY two-dimensional NMR experiment on this complex reveals methyl site exchange by an ethylene dissociation process and again implicates the fluxional $IrMe₂P₃$ + species. The fluxional behavior of $[IrMe_2(C_2H_4)P_3]BF_4$ is again rare for a $d⁶$ octahedral compound and is in contrast to that for the recently reported isoelectronic, stereochemically rigid complex **cis,mer-[IrH2(C2H4)(PMe2Ph),]BF,.** The Beck group has similarly reported the displacement of coordinated BF_4 by ethylene with the formation of the thermally unstable complex *trans*-HIr(PPh₃)₂(CO)(C₂H₄)Cl⁺. The solid-state X-ray structure determination of $IrMe₂$ - $(C_2H_4)P_3^+$ reveals an octahedral structure with ethylene coplanar with the two methyl groups and the unique phosphine phosphorus. The BF_4 is well separated and tetrahedral. The C=C(ethylene) distance of 1.41 **8,** compares with other transition-metal-ethylene complexes; no exceptionally short C-C distance is present, as might be expected for such a weakly held ethylene ligand. Overall, the structure resembles that of $IrMe₂(FBF₃)P₃$ with an ethylene in place of η ¹-BF₄. The trans phosphine ligands deviate from a linear arrangement (168") in an effort to minimize steric interactions with the remaining phosphine ligand. As with $IrMe₂(FBF₃)P₃$, the Ir-C bond for the methyl trans to the weakly coordinated ligand $(\eta^1-BF_4$ and C_2H_4) shows a markedly shortened length as compared to that for the other methyl trans to phosphine.

Given the cis ethylene/methyl relationship in [IrMe,- $(C_2H_4)P_3]BF_4$, the possibility of ethylene insertion was anticipated. Heating IrMe₂(13C₂H₄)P₃⁺ under a ¹³C₂H₄ atmosphere overnight results in the formation of $[Ir(13 C_2H_4$)₂P₃]BF₄ and ethane without the appearance of any possible ethylene insertion products (e.g. $\rm CH_{3}^{13}CH_{2}^{13}CH_{2}CH_{3}$, $^{13}CH_{2}="13}CHCH_{3}$.

Carbon monoxide will also displace η^1 -BF₄, forming cis, mer-[IrMe₂(CO)P₃]BF₄. This complex, unlike IrMe₂- $(FBF_3)P_3$ and $[IrMe₂(C₂H₄)P_3]BF_4$, is stereochemically rigid and shows no tendency for ligand (CO) dissociation. In the presence of excess CO, Ir \overline{Me}_2 (CO) P_3^+ is partially converted to the acyl complex mer-[IrMe(COMe)- $(CO)P_3|BF_4$. Even under 2 atm of CO, insertion to produce the acyl complex only proceeds to ca. 20% completion. Deinsertion of CO occurs in solution under vacuum or on standing under N_2 for a period of 2 days and regenerates the dimethyl cation.

The stereochemistry of the acyl complex A, particularly the trans arrangement of the acyl and entering CO ligands, suggests that the C*O that adds to $IrMe₂(CO)P₃⁺ enters$ between P' and Me' (Scheme I), either in the 18-electron carbonyl complex (path a) or in an acyl precursor (path b). That this is the correct stereochemistry of nucleophilic attack is also supported by the fact that, for rhodium, C*O above can be replaced by ethylene and still result in acetone formation.

Although factors that control acyl-alkyl equilibria in $iridium(III)$ complexes are not well understood, Bennett²⁹

has found that the equilibrium between the five-coordinate acyl complex $Ir(Cl)(COR)(PMe₂Ph)₃$ ⁺ and the six-coordinate alkyl complex $mer-Ir(CI)(R)(CO)(PMe₃Ph)₃$ ⁺ is entirely on the side of the alkyl complex. Our work shows that replacement of chloride with methyl in this complex shifts the equilibrium toward the acyl.

In THF, $RhMe₂P₃BF₄$ reacts with CO to rapidly deposit the THF-insoluble complex *cis,mer*-[RhMe₂(CO)P₃]BF₄. When additional CO is added to CH_2Cl_2 solutions of this dimethyl carbonyl cation, the elimination of acetone occurs with the formation of $Rh(PMe₂Ph)₄$ ⁺. Although not observed, it is quite reasonable that a cis methyl acyl complex, similar to that observed for Ir, is the intermediate which proceeds to eliminate acetone under mild conditions. 30 The formation of acetone and not ethane establishes that CO insertion, followed by nucleophile-promoted reductive elimination of acetone, is a faster, more favorable process than simple reductive elimination of ethane. Such behavior finds precedent in the reactivity of RuR₂- $(CO)_{2}P_{2}.^{31}$

The project described here was initiated to produce unsaturated metal complexes containing ligands anticipated to be reactive toward unsaturated nucleophiles (e.g., CO and C_2H_4). The results reported here reveal rather dramatically the differences in the reactivity and stability of analogous rhodium and iridium complexes. As is generally observed, the reactions of the 5d metal are slower than those of a 4d analogue (e.g., eq 3, where $T \leq 25$ °C Exercise the stations of the 5d metal are slower
of a 4d analogue (e.g., eq 3, where $T \le 25$ °C
MMe₂P₃BF₄ + C₂H₄ $\frac{T}{T}$ C₂H₆ + ... (3)

$$
MMe2P3BF4 + C2H4 \xrightarrow{T} C₂H₆ + ... (3)
$$

for rhodium but $T \ge 50$ °C for iridium). The products are also different. For iridium, the 18-electron trigonal-bipyramidal Ir(C_2H_4)₂ P_3 ⁺ is obtained. In contrast, the functional unit $Rh(PMe₂Ph)₃$ ⁺ apparently fails to bind ethylene well, and this 14-electron species undergoes a complex phosphine ligand redistribution to generate Rh- $(PMe₂Ph)₄$ ⁺. It is remarkable that ethylene is capable of *inducing* reductive elimination of ethane from $RhMe₂P₃BF₄$ but unable to *bind* to $RhP₃⁺$.

When the nucleophile is CO, the reactivity is equally distinct for the two metals. For iridium, carbonylation proceeds through a CO adduct to give a carbonyl-methyl-acetyl species that does not reductively eliminate. For rhodium, the reaction proceeds without *detectable* intermediates to eliminate acetone. However, the excess CO (like C_2H_4) is incapable of stabilizing RhP_3^+ , and the same redistribution observed above takes place, to again yield RhP_4^+ . $Rh(PMe_2Ph)_3Cl$, upon treatment with $LiBF_4$ in $CH₂Cl₂$, also produces major amounts of [Rh- $(PMe_2Ph)_4|BF_4$. We will report elsewhere on our attempts to inhibit phosphine redistribution using a tridentate phosphine.

All of the above reactions may be classified as nucleophile-induced reductive eliminations. Such reductive eliminations, being the product-releasing step, are integral to several catalytic reactions. In the case of ethylene hydrogenation, we earlier reported 6 that elimination of ethane from $HIrEtP_3^+$ is ethylene-promoted. Here, we observe nucleophile-promoted carbon-carbon reductive eliminations. In the case of CO and rhodium, the nucleophile becomes part of the eliminated molecule but is unable to stabilize the Rh product. The concept of nucleophile promotion of reductive elimination is even effective when that nucleophile does not bind to the products (eq **4)** and In the case of CO and rhodium, the nucleophile
les part of the eliminated molecule but is unable to
ize the Rh product. The concept of nucleophile
otion of reductive elimination is even effective when
ucleophile does not

$$
\mathrm{RhMe}_2(\mathrm{CO})\mathrm{P}_3^+\xrightarrow{\mathrm{C}_2\mathrm{H}_4}\mathrm{Me}_2\mathrm{CO} + \mathrm{Rh}\mathrm{P}_4^+ + \dots \quad (4)
$$

is thus clearly a kinetic phenomenon. These reductive eliminations are found to be more kinetically facile for rhodium than for iridium, with the corollary benefit that intermediates are readily detected for the sluggish iridium analogues. The availability of unsaturated polyalkyl complexes (or their operational equivalent) by the acidolysis methodology employed here is thus useful in the generation, characterization, and study of reactive organometallic intermediates.

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Registry No. fac-IrMe₃P₃, 15927-48-7; IrMe₂P₃BF₄, 127973-**72-2; IrMe2P3BF4.C6H6, 127973-73-3; mer-RhC13P3, 14882-42-9;** $fac-RhMe₃P₃$, 128051-02-5; $RhMe₂P₃BF₄$, 123143-41-9; $C₂H₄$, **74-85-1; cis,mer-IrMe2(C2H4)P3BF,, 127973-75-5; CO, 630-08-0; cis,mer-[IrMez(CO)P3]BF4, 127973-77-7; [IrMe(COMe)P3]BF4, 127973-79-9; &,mer-[RhMe2(CO)P3]BF4, 127973-81-3.**

Supplementary Material Available: Listings of anisotropic thermal parameters for IrMe₂(PMe₂Ph)₃BF₄.C₆H₆ and [IrMe₂-**(C2H4)(PMe2Ph)3]BF4 (2 pages); listings of observed and calculated structure factors (16 pages). Ordering information is given on any current masthead page.**

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⁽³⁰⁾ The intramolecularity of the elimination process has not **been proven here.**

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