$= 2.47$ Å, $d(Hg,Cl) = 2.48$ Å, $d(P,H) = 1.40$ Å, and $\theta(P,Hg,P) =$ θ (Cl,Hg,Cl) = θ (H,P,H) = 110°. All further calculations on geometrically distorted molecules $(\theta(P,Hg,P)$ and $\theta(C),Hg,C)$ being the only distorted structural parameters) were performed without further changes of the H_{ii} 's.

Coupling constants $J(Hg, P)$ were calculated by using the formalism of Pople and Santry.³³

$$
J(\mathbf{A}, \mathbf{B}) = \frac{32}{3} h \gamma_A \gamma_B |\Psi_{s,A}(0)|^2 |\Psi_{s,B}(0)|^2 \pi_{AB}
$$

$$
\pi_{AB} = +4 \sum_{i}^{\infty} \sum_{j}^{\text{unocc}} (\frac{3}{2} \Delta E_{ij})^{-1} C_{i,A} C_{j,A} C_{i,B} C_{j,B}
$$

 $|\Psi_{s,M}(0)|^2$ is the s density of the valence *s* orbital centered on A or B, γ is a gyromagnetic ratio, ³ ΔE_{ij} are triplet excitation energies, and C_iA (C_{*IA*}, etc.) is the coefficient of an atomic s orbital centered on
A (or **B**) in the *i*th MO (*i*th MO). The coefficients were produced A (or B) in the *i*th MO (*j*th MO). The coefficients were produced by the EHMO calculations, ${}^3\Delta E_{ij}$ was taken as the difference between

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- We are aware of the structure of $[Hg(CIO_4)_2(P(C_6H_{11})_3)_2]$,³⁴ with $\theta(P,Hg,P) = 170.7^{\circ}$, $\theta(O,Hg,O) = 137.9^{\circ}$, a calculated $J(Hg,P)$ of 3400 Hg, and an observed $J(Hg, P)$ of 3755 Hz. For calculation of $J(Hg, P)$ we assume that we may consider this structure a case of four-coordinate Hg2+. Note, however, that the **Hg-O** distances are 2.93 and 3.08 **A** and that there are two further perchlorate oxygens O' at 3.27 and 3.23 Å with an angle $\theta(O',Hg,O') = 141.3^\circ$ and angles $\theta(O,Hg,O') \approx 40^\circ$. Thus an alternative interpretation of this structure would be in terms of a distorted octahedron.
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Notes

the eigenvalues E_i and E_j , and the *s* density at the nucleus was that of the neutral atom for phosphorus and that of $d^9s^1p^1$ mercury(I).

Multiple Linear Regression. The data in Table **I1** were used with unit weight to determine the coefficients *a', b,* and *c* in the expression $J(Hg, P) = a' + b[\theta(P, Hg, P) - \theta^{0}(P, Hg, P)] + c[\theta(X, Hg, X) - \theta^{0} - \theta^{0}(P, Hg, P)]$ (X,Hg,X) , where $\theta^0(P,Hg,P) = \sum \theta(P,Hg,P)/N$ and $\theta^0(X,Hg,X)$ = $\sum \theta(X,Hg,X)/N$, with N the number of observations. Choosing the origin of the regression at θ^0 , the "center of mass", minimizes the trace of the variance-covariance matrix. Results are (σ) in parentheses) $a' = 4149$ (117) Hz, $b = 25.1$ (5.5) Hz/deg, $c = -48.7$ (8.3) Hz/deg, $r^2 = 0.93$, and $(\sum \Delta^2 / N - 3)$ ^{1/2} = 350 Hz (standard deviation of an observation of unit weight; $\Delta = J_{\text{obsd}} - J_{\text{calcd}}$; elements of the correlation matrix are $C(a',b) = 0$, and $C(b,c) = 0.3$. Results for $\theta^0(P,Hg,P)$ $= \theta^{0}(X,Hg,X) = 109.5^{\circ}$ are $a'' = 3271$ (155) Hz, $C(a'',b) = -0.47$, and $C(a''$, $c) = 0.31$, and all others are the same as above. The values given in the Discussion are obtainable from the above results.

Acknowledgment. R.W.K. and M.P. thank the Schweizerischen Nationalfonds zur Forderung der Wissenschaftlichen Forschung for financial support and Mr. L. Zoller for computational assistance.

Registry No. I, 27902-66-5; **11,** 14057-00-2; **111,** 80063-20-3; $Hg(ac)_2(cis-Ph_2PCH=CHPPh_2), 80063-21-4; HgCl_2(cis \texttt{Ph}_2 \texttt{PCH=CHPPh}_2$), 80063-22-5; $\texttt{Hg(SCN)}_2 (cis\text{-}Ph_2 \text{PCH}$ = 22-3; Hg((EtO)₂PO₂)₂(cis-Ph₂PCH=CHPPh₂), 80083-23-4; Hg₁₂- $(cis-Ph₂PCH=CHPPh₂), 80083-24-5; Hg(ac)₂(PPh₃)₂, 66119-73-1;$ $HgCl₂(PPh₃)₂$, 14494-85-0; $HgBr₂(PPh₃)₂$, 14586-76-6; Hg-CHPPh₂), 80106-35-0; $Hg(CN)_{2}(cis-Ph_{2}PCH=CHPPh_{2}),$ 80083- $(SCN)_2(PPh_3)_2$, 27290-69-3; HgI₂(PPh₃)₂, 14494-95-2.

Supplementary Material Available: Listings of structure factor amplitudes and of anisotropic thermal parameters (51 pages). Ordering information is given on any current masthead page.

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Controlling the Number of Metal Sites to Which Ph₂PCH₂CH₂PPhCH₂CH₂PPh₂ Coordinates in Tungsten **Carbonyls'**

Richard L. Keiter,*za James W. Brodack, Rodney D. Borger, and Lew W. Cary^{2b}

Receioed *May* 6. 1981

Polydentate phosphorus ligands have been used extensively in coordination chemistry for the past 20 years but in a narrowly focused manner that places principal emphasis on their chelating properties. **A** rich and relatively unexplored aspect of these ligands is their ability to bind in many arrangements other than the familiar fully chelated mode in monometallic species. This area of chemistry has been largely ignored because selective syntheses that eliminate tedious separations have not been available. In this work we have chosen to prepare complexes of $Ph_2PCH_2CH_2PPhCH_2CH_2PPh_2$, triphos, to illustrate the utility of vinyl-addition reactions in controlling ligand coordination.

triphos was first synthesized in 1962 ³ Interest in the ligand lagged through the 1960s because its preparation was not attractive. It became widely used, however, following the discovery of high-yield syntheses by King,⁴ Issleib,⁵ and Meek.⁶

At present it is routinely used in complexation studies such as those aimed at elucidating the reactions in homogeneous hydroformylation,⁷ Fischer-Tropsch synthesis, $\frac{8}{3}$ and catalytic hydrogenation.⁹

The ligating possibilities of triphos have been previously outlined, and specific examples of these species have been identified.¹⁰ In order to synthesize such complexes routinely, however, it is necessary to find a method of controlling the number of metal sites to which the polyphosphine coordinates. Substitution reactions are in general inadequate because they too often lead to mixtures of isomers or mixtures of mono- and bimetallic products that may not be easily separated. In this work we report a method of obtaining the desired products that is based upon building the complex of interest from judiciously selected coordinated fragments. The approach has been used previously in the preparation of monodentate complexes of diphos, $Ph_2PCH_2CH_2PPh_2$, such as (OC) , $M(diphos)$ and $(OC)₄M(diphos)₂$ (M = Cr, Mo, W).¹¹

-
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⁽¹⁾ Presented at the 1980 Biennial Inorganic Chemistry Symposium at Guelph, Canada.

^{(2) (}a) Eastern Illinois University. (b) Presently at Nicolet Technology Corp., Mt. View, CA 94041. (3) W. Hewertson and H. R. Watson, J. Chem. *SOC.,* 1490 (1962).

There are five possible nonchelated triphos complexes of pentacarbonyltungsten **(I-V,** Chart **I).** This report outlines the syntheses of these five complexes and demonstrates the versatility of addition reactions in the selective coordination of polyphosphines.

Experimental Section

Physical **Measurements.** 31P NMR and infrared spectra were obtained from chloroform solutions as described elsewhere.¹¹ Chemical shifts are reported with positive values downfield from the 85% phosphoric acid reference.

Materials. Diphenylvinylphosphine, phenyldivinylphosphine, diphenylphosphine, phenylphosphine, and tungsten hexacarbonyl were purchased from Pressure Chemical Co. and used without further purification. All reactions were carried out under a nitrogen atmosphere.

Preparations. The complexes $(OC)_5 WPPh_2CH=CH_2$ and $(OC)_5 WPPh_2H$ were prepared as described previously.¹¹ (OC) _sWPPh $(CH=CH₂)₂$ (86%) was prepared by THF displacement from (OC) , W(THF)¹² and purified by molecular distillation (60 °C) at 0.02 torr for 2 days).

 (vs) $(E + 1)$ **(OC)₅WPPh(CH==CH₂)₂: IR 1983 (w) (B₁), 2075 (s)** $(A_1^{(2)}), 1941$ cm⁻¹; ³¹P NMR δ 2.27 (J_{WP} = 236.3 Hz).

C, 36.91; H, 2.44; P, 6.46. Anal. Calcd for $C_{15}H_{11}O_5PW$: C, 37.07; H, 2.28; P, 6.37. Found:

(1.0 **g,** 2.1 mmol) and **2,2'-azobis(isobutyronitri1e)** (AIBN) (0.1 **g)** was added PPh₂H (0.75 mL). The mixture was heated for 24 h at 75 °C. Excess phosphine was removed by high vacuum. The oily mass eventually crystallized from CH₂Cl₂/CH₃OH (40%): mp 69–74

^oC; IR 1983 (w) (B₁), 2074 (s) (A₁⁽²⁾), 1940 (vs) (E + A₁⁽¹⁾) cm⁻¹. **(OC), WPPh(CH₂CH₂PPh₂)₂ (I).** To (OC), WPPh(CH=CH₂)₂

Anal. Calcd for $C_{39}H_{33}P_3O_5W$: C, 54.56; H, 3.87; P, 10.82. Found: C, 54.40; H, 3.80; P, 10.99.

(OC), WPPh₂CH₂CH₂PPhCH₂CH₂PPh₂ (II). A mixture of $(OC)_5WPhP(CH=CH_2)_2$ (1.2 mmol), PPh₂H (1.0 mmol), and AIBN (0.1 g) was heated for 24 h at 75 °C. Excess $PhP(CH=CH₂)₂$ was removed under high vacuum after which (OC)₅WPPh₂H (1.0 mmol) and AIBN (0.1 **g)** were added. The new mixture was heated for 24 h at 75 °C. The oil that resulted was dissolved in a minimum of $CH₂Cl₂$ and an equal volume of CH₃OH. Refrigeration at 6 °C led to oil formation. The process was repeated twice more to give an oil, which was identified by ³¹P NMR (20%): IR 1982 (w) (B_1) , 2074 (s) $(A_1^{(2)})$, 1941 (vs) $(E + A_1^{(1)})$ cm⁻¹

(2.0 **g,** 3.7 mmol) and AIBN (0.1 **g)** was added PPhH2 (3.7 mmol). The mixture was heated for 24 h at 75 °C. Unreacted PPhH₂ was removed by high vacuum. Attempts to crystallize the product from $CH₃OH/CH₂Cl₂$ gave an oil, which was chromatographed (silica gel) $PhP|CH_2CH_2PPh_2W(CO)_{3}]_2$ (III). To (OC), WPPh₂(CH=CH₂)

Figure 1. ³¹P{H} NMR spectrum of (OC) , WPPh $(CH_2CH_2PPh_2)$.

with 80% petroleum ether/20% ethyl acetate. An oil was obtained which crystallized after a period of 2 months (58%): mp 80-85 °C; IR 1983 (w) (B₁), 2074 (s) $(A_1^{(2)}, 1941$ (vs) $(E + A_1^{(1)})$ cm⁻¹.

Anal. Calcd for $C_{44}H_{33}P_3W_2O_{10}$: C, 44.68; H, 2.79; P, 7.86. Found: C, 44.81; H, 2.78; P, 7.56.

(OC), WPPh₂CH₂CH₂PPh[W(CO)₅](CH=CH₂). To a mixture of (OC) ₅WPPh $(CH=CH_2)_2$ (2.67 g, 5.45 mmol) and $KO(t-Bu)$ (0.1) g) in THF (150 mL) was added dropwise $(OC)_5 WPPh_2H$ (1.31 g, 2.57 mmol) dissolved in THF (50 mL) over a 1-h period. The residue, obtained after solvent removal, was crystallized from CH_2Cl_2/CH_3OH (56%). The first fraction was contaminated with the trimetallic complex (V): mp 123-125 °C; IR 1983 (w) (B_1) , 2074 (s) $(A_1^{(2)})$, 1941 (vs) $(A_1^{(1)} + E)$ cm⁻¹. Absorption bands are somewhat broadened because of inequivalent $W(CO)$ ₅ groups.

Anal. Calcd for $C_{32}H_{22}O_{10}W_2P_2$: C, 38.57; H, 2.21; P, 6.22. Found: C, 38.30; H, 2.09; P, 6.08.

(OC)_sWPPh₂CH₂CH₂PPh[W(CO)_s]CH₂CH₂PPh₂ (IV). To a mixture of $(OC)_5WPPh_2CH_2CH_2PPh[W(CO)_5]$ $(CH=CH_2)$ (1.16 **g,** 1.16 mmol) and KO(t-Bu) (0.1 **g)** in THF (150 mL) was added $PPh₂H$ (1.12 mmol). The mixture was heated under reflux for 1.5 h, after which solvent and excess $PPh₂H$ were removed to yield an oily mass. The product was crystallized from CH_2Cl_2/CH_3OH (40%): mp 139-141 °C; IR 1983 (w) (B₁), 2073 (s) $(A_1^{(2)})$, 1941 (vs) $(A_1^{(1)}$
+ **E**) cm⁻¹.

Anal. Calcd for $C_{44}H_{33}P_3W_2O_{10}$: C, 44.68; H, 2.79; P, 7.86. Found: C, 44.61; H, 2.80; P, 8.11.

 $(OC)_5 WPPh[CH_2CH_2PPh_2W(CO)_5]_2$ **(V).** To a mixture of (OC) , WPPh₂H (2.12 g, 4.15 mmol) and $KO(t-Bu)$ (0.2 g) in THF (25 mL) was added (over a 1-h period) (OC) , WPPh $(CH=CH₂)₂$ (1.03 **g,** 2.12 mmol) in THF (25 mL). The white product was crystallized from CH_2Cl_2/CH_3OH (79%): mp 160-162 °C; IR (signals broad) 1985 (w) (B₁), 2076 (s) $(A_1^{(2)})$, 1941 (vs) $(A_1^{(1)} +$ $E)$ cm⁻¹

Anal. Calcd for $C_{49}H_{33}O_{15}W_3P_3$: C, 39.07; H, 2.21; P, 6.17. Found: C, 38.92; H, 2.49; P, 5.92.

Results and Discussion

The complex $(OC)_5WPPh(CH=CH_2)_2$ is a nonviscous liquid, which can be obtained in high purity by molecular distillation. It is stable in air at room temperature and can be stored for long periods of time if it is protected from light. It is an excellent starting material for the production of I, undergoing free radical induced addition of diphenylphosphine.¹³

$$
(OC)5WPPh(CH=CH2)2 + 2PPh2H $\frac{AIBN}{75 \text{ °C}}$
Ph₂PCH₂CH₂PPh[W(CO)₅]CH₂CH₂PPh₂
$$

The structure of I is unambiguously established by its ${}^{31}P$ spectrum (Figure 1), which shows a downfield triplet (7.3 ppm) and an upfield doublet (-12.5 ppm) of appropriate intensities $(^3J_{\text{PP}} = 33.8 \text{ Hz})$. ¹⁸³W satellites are observed for the coordinated phosphorus atom $(J_{WP} = 236.7 \text{ Hz})$. In solution, I is slowly oxidized to a monoxide which shows 31P chemical

⁽¹²⁾ W. Strohmeier and F. Müller, Chem. Ber., **102**, 3608 (1969).

⁽¹³⁾ D. W. Meek, D. L. Dubois, and J. Tiethof, Adu. *Chem.* Ser., **No. 150, 335 (1976). 2,2'-Azobis(isobutyronitrile),** AIBN, is a convenient free radical source for these reactions just **as** it is for the syntheses of **un-**

Table I. "P NMR Data of Pentacarbonyltungsten Complexes of Ph₂ PCH₂ CH₂ PPhCH₂ CH₂ PPh₂

complex	δ WPPh	δ WPPh ₂	δ PPh	δ PPh.	$J_{\rm PP}$ Hz	$J_{\rm WP}$ Hz
I	7.3			-12.5	33.8	236.7
$_{II}$		12.5	-16.6	-12.8	29.4 ^a 31.1	239.4
Ш		13.6	-17.3		30.7	239.9
IV	6.8	12.7		-12.1	36.5^{b}	238.6 ^c
					32.7	239.9 ^d
V	7.5	13.5			34.7	239.4^{c} 242.0 ^d
I (oxide)	7.4			-12.6	43.2^e	g
				31.1 ^f	34.2	
II (oxide)			40.7^{f}			g
ll (oxide)				32.7^{f}		g
III (oxide)			45.6^{f}		46.4^e	g
IV (oxide)		12.4		29.6^{f}	46.3^{e}	g
					31.9	

 $J_{\rm PPh}$, W-PPh. $J_{\rm PPhW-PPh}$, c $J_{\rm PPhW}$. d $J_{\rm PPh}$ _y f_c g Herewhed. $e_{\delta \text{PPO}}$. $f_{\delta \text{PO}}$. g Unresolved.

Figure 2. ³¹P{H} NMR spectrum of $(OC)_5WPPh_2CH_2$ - $CH₂PPhCH₂CH₂PPh₂$.

shifts at -12.6 , 7.4, and 31.1 ppm for uncoordinated phosphorus, phosphorus coordinated to tungsten, and phosphorus coordinated to oxygen, respectively. Phosphorus-phosphorus coupling between coordinated phosphorus atoms is 43.2 Hz and between coordinated and uncoordinated atoms is 34.2 Hz (Table **I).** phorus, phosphorus coordinated to tungsten,
coordinated to oxygen, respectively. Phosph
coupling between coordinated phosphorus and
and between coordinated and uncoordinated
(Table I).
Distanting II, a structural isomer of

Obtaining **11,** a structural isomer of I, was synthetically **DIDE TT** AIBN

PhP(CH=CHZ)(CH2CH2PPh2) AIBN PhP(CH=CH2)(CH2CH2PPh2) + (0C)SWPPhZH - (OC) *5* WPPh2CH2CHZPPhCH2CH2PPh2 **I1**

difficult and was best accomplished by a two-step reaction involving $Ph_2PCH_2CH_2PPh(CH=CH_2)$ as an unisolated intermediate. A downfield doublet $(12.5$ ppm, $^{3}J_{PP} = 29.4$ Hz) flanked by ¹⁸³W satellites $(J_{WP} = 239.4 \text{ Hz})$, an upfield doublet $(-12.8$ ppm, $J_{\text{PP}} = 31.1$ Hz), and a doublet of doublets (-16.6) ppm) (Figure 2) clearly identifies the complex and dramatically distinguishes it from isomer **I.** Isomer I1 is somewhat more air sensitive than isomer **I,** and mixtures of $(OC)_5WPPh_2CH_2PPh(O)CH_2CH_2PPh_2$ ($\delta_{P=Q}$ 40.7) and $\overline{(OC)}$ ₃WPPh₂CH₂CH₂PPhCH₂CH₂PPh₂(O) $(\delta_{P\rightarrow Q}$ 32.7) were observed spectroscopically but not isolated.

Other reported examples of monoligated monometallic complexes are the unseparated isomers of $CH₃COFe(CO)$ -(triphos)(η^5 -C₅H₅) and of (OC)₄Fe(triphos).^{14,15} In addition, $[V(CO)_{5}(triphos)]$ ⁻ has been observed in solution by IR and $51V$ NMR spectroscopy.¹⁶

It one wishes to obtain pure monoligated monometallic isomers of triphos, clearly, substitution reactions are not desirable. Vinyl-addition reactions provide a route whereby the coordination of terminal or central phosphorus atoms is not left to chance. Thus it is possible to synthesize **I** without generating **I1** and vice versa. **In** addition, the absence of vacant coordination sites during the course of the reaction precludes the formation of phosphine-bridged polymetallic species.

Complex III was also obtained by free radical addition. A

\n(OC),
\n
$$
WPPh_2(CH=CH_2) + PPhH_2 \rightarrow (OC)
$$
,
\n $WPPh_2CH_2CH_2PH_2CH_2W(CO)$,
\nIII

downfield doublet (13.6 ppm, ${}^{3}J_{\text{PP}} = 30.7 \text{ Hz}$) with 183W satellites $(J_{\rm WP} = 239.9 \text{ Hz})$ and an upfield triplet (-17.3 ppm) establish the structure of 111. Air oxidation of **111** gave the oxide, which revealed a downfield triplet (δ_{PO} 45.6, J_{pp} = 46.4 Hz), which was assigned to the phosphoryl group. $_2CH_2CH_2PH_2CH_2W(CO)$,

III

pm, $^3J_{PP} = 30.7$ Hz) with ¹⁸³W

and an upfield triplet (-17.3 ppm)

II. Air oxidation of III gave the

nfield triplet (δ_{PO} 45.6, $J_{PP} = 46.4$

b the phosphoryl group.

of III, was prepa

Complex IV, the isomer of **111,** was prepared in two steps.

establish the structure of III. Air oxidation of III gave the
oxide, which revealed a downfield triplet (
$$
\delta_{PO}
$$
 45.6, J_{PP} = 46.4
Hz), which was assigned to the phosphoryl group.
Complex IV, the isomer of III, was prepared in two steps.
(OC),
WPPh(CH=CH₂)₂ + (OC),
WPPh₂CH₂CH₂PPh[W(CO)₅](CH=-CH₂)
(OC),
WPPh₂CH₂CH₂PPh[W(CO)₅](CH=CH₂) +
PPh₂H $\xrightarrow{KO(t-Bu)}$
(OC)₅WPPh₂CH₂CPPh[W(CO)₅](CH=CH₂) +
IPh₂H $\xrightarrow{KO(t-Bu)}$
IV

The intermediate was isolated as a white solid in reasonable yield (56%) (δ_{WPPh_2} 12.8, $J_{\text{PP}} = 35.7 \text{ Hz}$, $J_{\text{WP}} = 240.9 \text{ Hz}$; δ_{WPPh} 4.5, J_{WP} = 238.6 Hz) even though separation of it from starting materials and trimetallic complex, V, was required. Characterization of IV by 31P NMR confirms its structural arrangement. The uncoordinated phosphorus atom gives rise to a doublet $(-12.1 \text{ ppm}, {}^{3}J_{\text{PP}} = 36.5 \text{ Hz})$ as does the coordinated terminal phosphorus atom ($\delta = 12.7$, $J_{PP} = 32.7$ Hz, J_{WP} = 239.9 Hz). The coordinated central phosphorus atom appears as a doublet of doublets (6.8 ppm, $J_{WP} = 238.6 \text{ Hz}$). The oxide, occasionally found as a contaminant, gives rise to two doublets (δ_{WPPh_2} 12.4, $J_{\text{PP}} = 31.9 \text{ Hz}$; δ_{PO} 29.6, $J_{\text{PP}} = 46.3 \text{ Hz}$ Hz) and a doublet of doublets obscured by signals from its precursor. ¹⁸³W satellites were not observed because of low concentration. No attempt was made to resolve the optical isomers of **IV** or its oxide.

Isomers **111** and IV are particularly unusual because these are the first reported examples of two independent metal moieties attached to nonchelating triphos. The complex $Fe₂(CO)₂(C₃H₅)₂(triphos)$, previously reported, is thought to be a biligated bimetallic type in which the end and center phosphorus atoms are bound to two iron atoms which are bridged by carbonyl groups.^{10,15} Two metal atoms are also attached to triphos in the case of $Br(CO)₃Mn(triphos)Cr (CO)_{5}^{17}$ and $[(\eta^{5}-C_{5}H_{5})_{2}Mn_{2}(CO)(NO)_{2}(triphos)][PF_{6}]_{2}^{15}$ but these contain chelated manganese. triphos may also chelate through the two end phosphorus atoms as has been proposed for *trans-* $[\eta^5$ -C₅H₅V(CO)₂(triphos)].¹⁶ a and center

is which are

oms are also

((triphos)Cr-

os may also

as has been

is

y, V can be
 $\frac{KO(t-Bu)}{2}$
 $h_2W(CO)_5]_2$

Of the five complexes reported in this study, **V** can be $KO(t, R_{II})$

$$
2(OC), WPPh2H + (CO), WPPh(CH=CH2)2 \xrightarrow{\text{RO}(1-Ba)} (OC), WPPh[CH2CH2PPh2W(CO)5]
$$

synthesized in highest yield, is most readily crystallized, and is the least soluble.

⁽¹⁴⁾ R. B. King, P. N. Kapoor, and R. N. Kapoor, Inorg. Chem., **10, 1841**

^{(1 97 1).} (15) R. B. King and J. C. Cloyd, **Jr.,** Inorg. Chem., **14,** 1550 **(1975).**

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Muhlbach, B. Rausch, and D. Rehder, *Ibid.,* **205, 343 (1981). (17) M.** L. Schneider, N. J. Coville, and I. *S.* Butler, *J.* Chem. *SOC.,* Chem. *Commun.,* **799 (1972);** I. S. Butler and N. J. Coville, *J.* Organomer. Chem., **80, 235 (1974).**

The ³¹P NMR spectrum of V is a characteristic AB₂ type in appearance, much resembling that of the free ligand.¹³ The satellite spectrum can be compared to that of $Ph_2P(Se)$ -CH₂CH₂P(Se)PhCH₂CH₂PPh₂(Se), which was recently analyzed.¹⁸ It consists of the AB_2 part of an AB_2X spin system overlapping with the ABC part of an ABCX spin system. The center phosphorus resonance is found at 7.5 ppm $(^3J_{\text{pp}} = 34.7$ $Hz, J_{WP} = 239.4 Hz$, and the terminal phosphorus atoms are found at 13.5 ppm $(J_{WP} = 242.0 \text{ Hz})$.

In the course of this work we have found free radical additions to be useful and even preferred so long as at least one of the reactants is an uncoordinated phosphine. When both secondary phosphines and vinylphosphines were coordinated, the free radical method failed, but in those cases potassium tert-butoxide led to the desired addition. This observation has been noted in the preparation of diphos derivatives as well.¹⁹

The enormous potential of using addition reactions for ligation control becomes apparent in this study. Many complexes, until now considered chemical oddities because of their synthetic inaccessibility, will become commonplace and available for catalytic, mechanistic, and spectroscopic studies.

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Registry No. I, 79919-64-5; I (oxide), 79933-12-3; **11,** 79919-65-6; **111,** 79919-66-7; **111** (oxide), 79919-67-8; IV, 79919-68-9; IV (oxide), 79919-69-0; V, 79919-70-3; (OC)₅WPPh₂CH₂CH₂PPh(O)- $\rm CH_2CH_2PPh_2$, 79919-71-4; (OC)_sWPPh₂CH₂CH₂PPhCH₂CH₂- $PPh₂(O)$, 79919-72-5; (OC), WPPh₂CH₂CH₂PPh[W(CO)₅](CH= CH₂), 79933-13-4; (OC)₅WPPh(CH=CH₂)₂, 79919-73-6; (OC) _SWPPh₂H, 18399-62-7; (OC) _SWPPh₂(CH=CH₂), 64012-10-8; PPh2H, 829-85-6.

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Kinetics and Mechanism of Oxidative Addition of Methyl Iodide to Four- and Five-Coordinate Iridium(1) Complexes

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Oxidative addition of methyl iodide to four-coordinate iridium and rhodium complexes is important in the Monsanto process' in the conversion of methanol to acetic acid, which was found to be catalyzed by iodide ions.^{2,3} We have lately found some more oxidative-addition reactions that are catalyzed by iodide ions.4

Our interest lies in the reactivities of five-coordinated d⁸ iridium complexes. Recently we presented the first evidence of dioxygen attacking a five-coordinate iridium(1) iodide complex,⁴ and we also obtained evidence for direct attack of dihydrogen on the five-coordinate complex IrH(CO)(PPh₃)₃.⁵

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Figure 1. Plots of k_{obsd} vs. [CH₃I] for different [I⁻] (in mol-dm⁻³) at 25 °C. Experimental points were fitted to the theoretical lines obtained from the spss program.

We report here the kinetic results of the iodide-catalyzed reaction

$$
[(\text{cod})\text{Ir(phen)}]^+ \text{Cl}^- + \text{CH}_3 \text{I} \xrightarrow{[\text{I}^+]} (\text{cod})\text{Ir(phen)}(\text{CH}_3)(\text{I})]^+ (1)
$$

 $(cod = cycloocta-1, 5-diene; phen = 1, 10-phenanthroline).$ The four-coordinate $[\text{Ir}(\text{cod})(\text{phen})]^+ \text{CI}^-$ in the presence of iodide establishes equilibrium 2 in methanol solution. A K_e value $[(\text{cod})\text{Ir}(\text{phen})]^+ \text{CI}^- + \text{I}^- \xrightarrow{\text{K}_e} (\text{cod})\text{Ir}(\text{phen})]$ (2) establishes equilibrium 2 in methanol solution. A *K,* value

$$
[(\text{cod})\text{Ir(phen)}]^+ \text{Cl}^- + \text{I}^- \xrightarrow{\text{K}_e} (\text{cod})\text{Ir(phen)}\text{I} \quad (2)
$$

of 227 ± 5 cm³-mol⁻¹ (at 25 °C) was determined spectrophotometrically.

The rate of reaction 1 was found to be first order in both [complex] and [methyl iodide] and increased with increasing sodium iodide concentration. Under the conditions of the kinetic measurements all the reactions went to completion. Since the concentration of methyl iodide was always in sufficiently large excess to remain essentially constant throughout the reaction, the observed rate of disappearance of Ir(cod)- $(\text{phen})X (X = Cl, I)$ in every case studied yielded linear semilog graphs. The data at 20, 30, and 35 \degree C are given in Table I.

Plots of k_{obsd} vs. [CH₃I] at different concentrations of iodide at 25 °C are shown in Figure 1. A family of straight lines through the origin for the methyl iodide concentration range used was obtained. The slopes of these plots increase with increasing [NaI]. All the data conform to the rate law given in eq 3, which can be derived from Scheme I.

$$
k_{\text{obsd}} = \left(\frac{k_1 + k_2 K_{\text{e}}[I^-]}{1 + K_{\text{e}}[I^-]}\right) [\text{CH}_3 I] \tag{3}
$$

The rate law given in eq 3 simplifies to $k_{\text{obsd}} = k_1[\text{CH}_3\text{I}]$ if the concentration of added iodide is zero. The slope of plot (a) in Figure 1 then gives the values of *k,* (Table 11), the rate

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