Some Complexes of Platinum, Palladium, Rhodium, and Iridium formed from the Bulky Ligand (Phenylethynyl)di-t-butylphosphine †

By H. David Empsall, Eileen M. Hyde, Eric Mentzer, and Bernard L. Shaw, * School of Chemistry, The University, Leeds LS2 9JT

The new phosphine PBut₂(C \equiv CPh) (L) gives complexes of the type trans-[MX₂L₂] (M = Pd or Pt, X = Cl or I) which exist as conformers due to restricted rotation around the phosphorus-metal bonds. The complexes are very resistant to nucleophilic attack but trans-[PtCl₂L₂] can be protonated, e.g., in CF₃CO₂H, [PtCl₂L{PBut₂(CH=CPh)}] + forms reversibly and with dry hydrogen chloride [PtCl₂{PBu^t2[CH=C(Cl)Ph]}2] is obtained. The complexes *cis*-[PtMe₂L₂] and *trans*-[PtCl(Me)L₂] are prepared by displacing cyclo-octa-1,5-diene (cod) from [PtMe₂(cod)], *etc.*, but with 2 mol of L [PtCl₂(cod)] readily gives [PtCl₂(cod)L] in which only one of the cod double bonds is coordinated. The complexes trans- $[MCI(CO)L_2]$ (M = Rh or Ir) have also been prepared and the iridium complex converted into [IrHCl₂(CO)L₂] with HCl. With trans-[MCI(CO)L₂] one rotamer is much preferred over the other two. Treatment of $H_2 IrCl_6$ with L gives five-co-ordinate [IrHCl_2L_2] which appears to be trigonal bipyramidal. whereas $RhCl_3$ gives $[RhHCl_2L_2]$ which is square pyramidal. Treatment of *trans*- $[MCl(CO)L_2]$ (M = Rh or Ir)

with sodium propan-2-olate in propan-2-ol gives the chelate complexes [M(CO){(PBut₂[CH=C(O)Ph]{L].

With sodium methoxide in methanol both acetylenic bonds are attacked giving [Ir(CO){PBut₂[CH=C(O)Ph]}-{PBut₂[CH=C(OMe)Ph]}]. Infrared, ¹H, ^{\$1}P, and some ¹³C n.m.r. data are given.

TRANSITION-METAL COMPLEXES with tertiary di-t-butylphosphines, $PBu_{2}^{t}R$ (R = alkyl or aryl), frequently show unusual chemistry and n.m.r. phenomena including low co-ordination numbers,^{1,2} unusual valency states,³ stable multihydrides,⁴ metallation reactions,⁵ very large chelate rings,⁶ or rotamers caused by restricted rotation about phosphorus-metal bonds.⁷ It therefore was of interest to study ligands of the type PBu^t₂(C=CR) for two main reasons. (i) There is a very large difference in the steric requirements of the t-butyl and the linear acetylenic group: one would therefore expect unusually large energy barriers to rotation about the phosphorusmetal bonds. Some of the chemical phenomena, e.g. metallation and large-ring formation, seem to be dependent on the size of these energy barriers. (ii) The acetylenic bonds of the phosphines might undergo reactions after co-ordination to give other unusual products.

† No reprints available.

¹ S. Otsuka and T. Yoshida, J. Organometallic Chem., 1973,

- 56, 413.
 ² A. Musco, W. Koran, A. Silvani, and M. W. Anker, J.C.S. Chem. Comm., 1973, 938.
 ³ D. Emergell, F. M. Hyde, and B. L. Shaw, J.C.S. Dalton, ³ H. D. Empsall, E. M. Hyde, and B. L. Shaw, J.C.S. Dalton,
- 1975, 1690. ⁴ B. L. Shaw and R. E. Stainbank, J.C.S. Dalton, 1972, 2108.

We chose to study as ligand (phenylethynyl)di-tbutylphosphine which was prepared from phenylethynyllithium and chlorodi-t-butylphosphine. This solid phosphine was characterized by its methiodide salt and by microanalyses and i.r. and n.m.r. spectroscopy (see Experimental section). The ¹³C n.m.r. data on this new phosphine in CDCl₃ solution are, with the assignment shown in (I): 30.3(d), ${}^{2}J(PH)$ 14, b; 33.3(d), ${}^{1}J(PH)$



18, a; 88.9(d), ²/(PH) 24 Hz, d; 106.6(s), c; 124.3(s), p; 128.8(s), o; and 131.9(s) p.p.m., m; the resonance for C^1 was probably obscured by the peak at δ 131.9 p.p.m. Shifts are relative to SiMe₄.

Platinum and Palladium Complexes.-Treatment of trans-[PtCl₂(NCBu^t)₂] or trans-[PdCl₂(NCPh)₂] with 2

⁵ H. D. Empsall, E. M. Hyde, D. Pawson, and B. L. Shaw, J.C.S. Dalton, 1977, 1292 and refs. therein.
⁶ A. Pryde, B. L. Shaw, and B. Weeks, J.C.S. Dalton, 1976,

- 322. ⁷ B. E. Mann, C. Masters, B. L. Shaw, and R. E. Stainbank,

TABLE 1

Yields, colours, m.p.s, and analytical data (calculated values in parentheses) for some complexes of PBut₂(C=CPh), L

	37:11			Microanalytical data (%)			
Complex	(%)	Colour	M.p. $(\theta_c/^\circ C)$	С	A		
(1) $[PtCl_{a}L_{a}]$	86	Yellow	243	51.1(50.65)	6.1(6.1)	9.5(9.35)	
$\langle 2 \rangle$ [PdCl.L.]	98	Yellow	218 - 221	57.35(57.35)	6.8(6.9)	10.55(10.6)	
(3) [PtI.L.]	94	Orange	286 - 288	41.1(40.8)	4.9(4.9)	27.1(26.95) ª	
(4) [PdI ₂ L ₂]	90	Purple	253 - 260	45.4(45.05)	5.45(5.45)	$30.1(29.75)^{a}$	
(5) [Pd ₂ Cl ₄ L ₂]	79	Orange	220 - 224	45.8(45.35)	5.5(5.45)	17.1(16.25)	
(6) [PtCl ₂ {PBu ^t ₂ [CH=C(Cl)Ph]} ₂]	75	White	187 - 192	46.5(46.2)	5.8(5.8)	16.8(17.05)	
$(7) [PtMe_2L_2]$	78	White	137—142 ^b	57.4(56.9)	7.4(7.3)	0(0)	
(8) $[PtCl(Me)L_2]$	76	White	198 - 202	53.7(53.7)	6.7(6.7)	5.0(4.8)	
(9) [PtCl ₂ (cod)L]	46	White	145 - 150	47.1(46.45)	5.8(5.7)	11.8(11.4)	
(10) [IrCl(CO)L ₂]	78	Yellow	202 - 205	52.9(52.95)	6.1(6.2)	5.1(4.75)	
(11) $[IrHCl_2(CO)L_2]$	38	White	182 - 184	50.0(50.5)	6.1(6.05)	9.3(0.05)	
(12) $[RhCl(CO)L_2]$	62	Yellow	194 - 197	60.2(60.15)	6.8(7.05)	5.6(5.4)	
(13) $[IrHCl_2L_2]$	15	\mathbf{Red}	160 - 165	50.7(50.8)	6.2(6.25)	9.7(9.35)	
(14) [RhHCl ₂ L ₂]	45	\mathbf{Red}	140 - 145	57.8(57.6)	6.9(7.1)	10.5(10.6)	
(15) $[Rh_2Cl_4L_2]$	47	Orange	133 - 140	45.6(45.75)	5.7(5.5)	17.3(16.9)	
(16) $[Ir(CO){PBut_2[CH=C(O)Ph]}L]$	94	Yellow	195 - 199	54.1(54.3)	6.4(6.5)	0(0)	
(17) $[Rh^{1}(CO){PBut}_{2}[CH=C(O)Ph]]L]$	95	Yellow	174 - 178	61.7(61.85)	7.4(7.4)	0(0)	
(18) $[Ir(CO){PBu_2[CH=C(O)Ph]}-{PBu_{0}[CH=C(OMe)Ph]}]$	90	Yellow	155 - 159	52.4(53.6)	6.6(6.75)	0.5(0)	
	۵ Iodi	ine analysis.	^b Gas evolved	on heating.			

TABLE 2

Spectral data

		1 H I	N.m.r."					
	But		<u>۸</u>	31P N	.m.r.¢			
	$\delta + 0.2$	$I^{b} + 0.5$		$\overline{\delta + 0.1}$	I(PtP) +		$1.r. (cm^{-1})$	a
	p.p.m.	Η̈́z	Other resonances	p.p.m.	Hz	ν (C \equiv C)	ν (M–Cl)	Other bands
Platinum	and palla	dium				,	· · ·	
(1) "	1.62(t)	15.4		26.5	2 628	2 196s	338s	
(2) [·]	1.61(t)	15.5		35.1(br) 41.7(br)	$\frac{2607}{f}$	2 189s	351s	
(3)	1.64(t)	15.4		13.2 26.0	$2 \ 477 \\ 2 \ 451$	2 178s		
(4) (5)	$1.62(t) \\ 1.62(d)$	$\begin{array}{c} 15.6 \\ 16.8 \end{array}$		52.1	g	2 183s 2 160s		
(6) (7) i	1.62(t) 1.70(d)	$13.7 \\ 13.2 $	$\begin{array}{c} 6.18(t) \ {}^{k} \ (7.2, \ 30.0) \\ 1.47(b) \ {}^{k} \ (, \ 70.5) \end{array}$	$\begin{array}{c} 29.1 \\ 28.5 \end{array}$	2 585 1 858	2 178m	334s	$1.569m[\nu(C=C)]$
$(8)^{i}$ $(9)^{i}$	1.87(t) 1.62(d)	$\begin{array}{c} 15.0 \\ 16.3 \end{array}$	1.49(t) k (14.0, 80.0) 2.75, 3.45, 4.95, 5.56 m	$\begin{array}{c} 36.3 \\ 28.2 \end{array}$	$\begin{array}{c}3 274 \\ 3 342 \end{array}$	2 184s 2 167s	258s 320s, 292s,	
Iridium a	and rhodiu	m					274s	
	δ	J *		δ	J	v(C≡C)	ν(M−Cl)	$\nu(C\equiv O)$
(10)	1.58(t)	15.1		f		2 181	313s	1 942s
(11)	1.60(t) 1.71(t)	$15.8 \\ 16.2$	-16.2(t) (Ir-H) ${}^{2}I(PH)$ 12.6	f		2 184	252s, 313s	2 035s
(12)	1.58(t)	12.8	J (= ==) ====	f		2.183m	308s	1 954s
(13)	$\begin{cases} 1.44(d) \\ 1.70(d) \end{cases}$	$15.7 \\ 15.8$	-31.3(t) (Ir-H) ${}^{2}I(PH)$ 21.6	15.0 ⁿ		2 178m	268s, 304s	
(14) ⁱ	1.79(t)	14.8	-29.8(dt), ¹ $J(RhH)$ 27.0 ² $I(PH)$ 30.6	47.3	¹ J(RhP) 105	2 193 m	295s	
			. .,				ν (C=C)	
(16)	f		f	$\{ 43.9 \} \\ 78.6 \}$	² J(PP) 293	2 178 m	1 525m	1 922a
(17)	f		f	42.8	¹ J(RhP) 127 ² J(PP) 284	2 171 m	1 519m	1 932s
(18)	f		f	86.0 45.2, 76.7	${}^{1}J({ m RhP})$ 137 ${}^{2}J({ m PP})$ 292		1 526s	1 923s

"At 35 °C and 60 MHz in CDCl₃ unless stated otherwise; d = doublet, t = triplet. ${}^{b}|{}^{3}J(PH) + {}^{5}J(PH)|$. At ambient temperature and 36.43 MHz in CDCl₃ unless otherwise stated. Shifts relative to 85% H₃PO₄; br = broad. d As Nujol mulls; w = weak, m = medium, s = strong. The numbers refer to the complexes in Table 1. See text. Not recorded. CH=C, ${}^{2}J(PH)$ and ${}^{3}J(PH)$ in parentheses. Hydrogen-1 n.m.r. in C₆H₆ or C₆H₆-C₆D₆. J(PH). ${}^{k}\delta(Me)$. In C₆H₆-C₆D₆. CH=C, of cod. In C₆H₆-C₆D₆.

mol equivalents of $\operatorname{PBut}_2(C\equiv CPh)$ (L) gave trans-[MCl₂L₂] (M = Pt or Pd) (see Experimental section and Table 1). Both complexes have the trans configuration from the i.r. and ¹H n.m.r. data (Table 2). The ³¹P n.m.r. spectrum of trans-[PtCl₂L₂] at 300 K in CDCl₃ shows two approximately equally intense peaks with satellites at δ 26.5 [¹J(PtP) 2 628] and 33.2 p.p.m. [¹J(PtP) 2 607 Hz]. These species are very probably due to the conformers (rotamers) (II) and (III) (although we cannot say which is which). As the temperature is



increased these peaks broaden reversibly, e.g. at 328 K two broad singlets are observed. This broadening is caused by some rotation about the platinum-phosphorus bonds. The ³¹P spectrum of trans-[PdCl₂L₂] at 300 K consists of two broad singlets at δ 35.1 and 41.7 p.p.m. which become very sharp at 233 K (8 33.4 and 40.7 p.p.m.) but collapse into one broad singlet at 328 K $(\delta 39.4 \text{ p.p.m.})$. The effects of temperature changes are reversible. The complex trans- $[PtCl_2L_2]$ was very resistant to attack by iodide ion, e.g. it was recovered unchanged after treatment with sodium iodide in refluxing acetone for 17 h. However, after reflux for 3 h with NaI in 2-methoxyethanol the di-iodo-complex was formed in 94% yield. The ³¹P n.m.r. spectrum of this di-iodide at 300 K shows the presence of two conformers (II) and (III) [8 13.2, ¹*J*(PtP) 2 477; 26.0 p.p.m., ¹J(PtP) 2 451 Hz]. Metathesis of trans-[PdCl₂L₂] with NaI in acetone at 20 °C gave red-brown trans-[PdI₂L₂]. Treatment of trans-[PdCl₂(PBu^t₂Ph)₂] with NaI in acetone under similar conditions gives the bridged complex [Pd₂I₄(PBu^t₂Ph)₂] with loss of phosphine, *i.e.* PBu^t₂(C \equiv CPh) probably has a smaller steric requirement with respect to an iodide ligand than does PBu^t₂Ph. The chloro-bridged binuclear complex $[\mathrm{Pd}_2\mathrm{Cl}_4\{\mathrm{PBu^t}_2\text{-}$ (C≡CPh)₂]² was readily prepared by mixing equimolar amounts of Na₂[PdCl₄] and L. The bridging system is split by pyridine to give [PdCl₂L(py)] which must have the two chlorines mutually trans since it shows just one strong band, at 351 cm⁻¹, in the far-i.r. spectrum.

We attempted to effect both nucleophilic and electrophilic attack on the acetylenic groups of the co-ordinated phosphines in trans-[PtCl₂L₂]. However, essentially quantitative recoveries of this complex were obtained after the following treatments: (*i*) sodium acetate in refluxing methanol for 1 d; (*ii*) Na[O₂CMe] in refluxing methanol-benzene for 1 d; (*iii*) sodium methoxide (10 mol) in refluxing methanol for 3 h; and (*iv*) sodium propan-2-olate (10 mol) in refluxing propan-2-ol for 15 min. Treatment of the complex with sodium propan-2-olate (10 mol) for 24 h gave a mixture which could not be separated. The i.r. spectrum showed a band at 2 184m cm⁻¹ due to ν (C=C) and one at 1 534s cm⁻¹ assigned to ν (C=C), indicating some attack on the acetylene bond(s). The complex trans- $[PtCl_2L_2]$ is converted into [PtCl₂{PBu^t₂[CH=C(Cl)Ph]}₂] by a solution of hydrogen chloride in dry diethyl ether. The ¹H n.m.r. spectrum shows a 'virtually 'coupled 1:2:1 triplet due to the t-butyl hydrogens at δ 1.62 p.p.m. [$|^{3}J(PH)$ + $^{5}J(PH)$ 7.2 Hz] with platinum satellites [J(PtH) 30.0 Hz] due to the two vinylic hydrogens. The ³¹P n.m.r. and i.r. spectra data (Table 2) also confirm the trans configuration of the phosphines and chlorines. We suggest that addition of hydrogen chloride occurs via initial electrophilic attack at the *a*-carbon atom of the alkyne followed by quenching of the carbonium ion by chloride. This would be entirely analogous to the addition of hydrohalogenic acids to diphenylacetylene⁸ or propyne.⁹ The ¹H n.m.r. spectrum of trans-[PtCl₂L₂] in deuteriochloroform-trifluoroacetic acid (1:1) shows two doublets for the t-butyls, indicating inequivalent phosphine ligands, and a triplet at δ 6.60 p.p.m., with an intensity corresponding to one hydrogen [J(PH) 6.0 Hz], with platinum satellites [J(PtH) 27.6 Hz]. The ³¹P n.m.r. spectrum shows an AB + ABX pattern (Table 2), the value of $J(P_A P_B)$ 375 Hz indicating trans-phosphorus nuclei. We therefore suggest that the cation trans- $[PtCl_{2}L{PBu_{2}(CH=CPh)}]^{+}$ is formed in CF₃CO₂H. Attempts to precipitate this cation as the tetraphenylborate salt were unsuccessful, the i.r. spectrum indicating that much of the complex trans- $[PtCl_2L_2]$ was present in the precipitate. When trans-[PtCl₂L₂] was heated under reflux in CF₃CO₂H for 4 h and the solution evaporated to dryness, the unchanged dichloro-complex was recovered. Treatment of the free phosphine PBu^t₂(C=CPh) with a solution of hydrogen chloride in diethyl ether gives [PBu^t₂(C≡CPh)H]Cl there being no evidence of addition to the triple bond. In contrast to the platinum complex, the palladium complex in CDCl₃-CF₃CO₂H (1:1) shows two broad resonances in the t-butyl region, and no signal due to vinyl hydrogens was observed.

Nucleophilic attack at the triple bond of the palladium complex was also attempted but decomposition to very dark products occurred on treatments with sodium propan-2-olate in propan-2-ol for 15 min or with sodium hydroxide in refluxing ethanol for 5 min.

We also attempted to replace the chloride ligands or to attack the triple bonds of *trans*-[MCl₂{PBu^t₂(C≡CPh)}₂] by methyl groups using methyl-lithium. The dichloroplatinum complex was heated with an excess of LiMe (5 mol) in benzene at 80 °C for 2 h. The resulting white solid product was a mixture which could not be separated; however, it had only 2% chlorine by microanalysis so it probably contained some of the required

⁹ H. Hunziker, R. Meyer, and H. H. Gunthard, *Helv. Chim.* Acta, 1966, **49**, 497.

⁸ K. Griesbaum, W. Naegele, and G. G. Wanless, J. Amer. Chem. Soc., 1965, 87, 3151.

methyl complex(es). Treatment of trans-[PdCl₂L₂] with LiMe (2 mol) in benzene gave an intense green solution and what appeared to be green crystals. Attempts to isolate solids from such reactions led to decomposition, e.g. the green crystals immediately decomposed on washing with dry air-free diethyl ether. A large excess $(\times 10)$ of LiMe gave deep blue crystals and an intense green solution, but attempts to isolate these deep blue crystals led to decomposition. We successfully prepared the complexes cis-[PtMe₂L₂], however, by displacing cyclo-octa-1,5-diene (cod) from [PtMe₂(cod)] with the ligand L (2 mol) in benzene. This formulation follows from the microanalytical and ¹H and ²¹P n.m.r. data (Tables 1 and 2).

The displacement of cod by $PBu_{2}^{t}(C=CPh)$ (2 mol) in benzene was followed by observing the disappearance of the t-butyl ¹H n.m.r. doublet of free PBu^t₂(C=CPh) and the appearance of the doublet of the product (the solutions were ca. 10⁻¹ mol dm⁻³). Reaction is complete in ca. 15 min at 34 °C. No intermediate could be detected by ¹H n.m.r. spectroscopy. Treatment of $[PtMe_2(cod)]$ with only 1 equivalent of L gives a 50% yield of cis-[PtMe₂L₂] and unchanged [PtMe₂(cod)], which stays in solution. The product cis-[PtMe₂L₂] is one of the few examples of a complex containing two phosphines with di-t-butyl substituents which are of cis configuration. Complexes of the type trans-[MX₂- $(PBu_{2}^{t}H)_{2}$ (M = Pd or Pt; X = Cl or Br) are isomerized photochemically to the corresponding cis complexes ¹⁰ and the chelating ligand $C_6H_4(CH_2PBut_2)_2$ -1,2 forms chelate complexes with Pt^{II}.¹¹

The complex cis-[PtCl(Me)(cod)] reacts with 2 mol equivalents of L to give trans-[PtCl(Me)L₂] as white prisms. The formulation is based on the analytical and spectroscopic data, e.g. a strong i.r. absorption peak at 258 cm⁻¹ is typical of v(Pt-Cl) (trans-methyl) and triplets in the ¹H n.m.r. spectrum due to the Bu^t and PtMe groups also indicate the *trans* configuration. The ³¹P n.m.r. spectrum shows a peak at 8 36.3 p.p.m. [¹*J*(PtP) 3 274 Hz].

Treatment of cis-[PtCl₂(cod)] with L (2 mol) in benzene gives $cis - [PtCl_2(cod)L]$ as white prisms. The formulation is based on the analytical and spectroscopic data (Tables 1 and 2). The ¹H n.m.r. spectrum shows a t-butyl doublet, two broad resonances at δ 2.75 and 3.45 p.p.m. due to eight methylene hydrogens on the cod, and resonances at 4.95 and 5.56 p.p.m. each of intensity two corresponding to the vinylic hydrogens. We suggest that only one of the cod double bonds is coordinated and that the resonance at δ 5.56 p.p.m. corresponds to the hydrogens on an unco-ordinated double bond. The ³¹P n.m.r. spectrum shows a peak at δ 28.2 p.p.m. [${}^{1}J(PtP)$ 3 343 Hz]. The i.r. spectrum shows a peak at 2 167 cm⁻¹ due to v(C=C) and peaks at 320, 292, and 274 cm⁻¹ due to v(Pt-Cl). The complex is non-conducting in nitrobenzene. When treated with iodomethane in acetone, yellow cis-[PtI₂(cod)] precipitated and the phosphonium salt [PMeBu^t₂(C=CPh)]I was isolated from the mother liquor. The phosphine ligand is thus labile.

Iridium and Rhodium Complexes.--We attempted to prepare [IrHCl₂(CO)L₂] of configuration (IV) by adding the phosphine (2 equivalents) to a solution prepared by the action of carbon monoxide on a boiling ethanolic solution of hexachloroiridic(IV) acid.¹² The solid product was a mixture of the required product contaminated with trans-[IrCl(CO)L₂]. This iridium(I) complex was readily prepared by treating the mixture with triethylamine in ethanol, and characterized in the usual way (Tables 1 and 2). As expected this complex consisted of rotamers due to restricted rotation about the iridiumphosphorus bonds, e.g. the ³¹P n.m.r. spectrum at 300 K consisted of two broad resonances and at 223 K of six sharp resonances, two singlets and one AB pattern corresponding to (II; X = Cl, Y = CO), (II; X = CO, Y = Cl), and (III; X = Cl, Y = CO). For the AB



pattern J(PP) 334 Hz. One of the singlets is very strong, one very weak, and the AB pattern is of intermediate intensity. It is likely that the preferred conformation is (II; X = Cl, Y = CO) since an X-ray structure of trans-[IrCl(CO){ $Bu_2^tPC \equiv C(CH_2)_5C \equiv CPBu_2^t$ }] ¹³ shows that only the conformer with $C \equiv O$ and $C \equiv C$ adjacent is present in the solid state, and the ³¹P n.m.r. spectrum shows that virtually all the complex is present as one conformer. The complex trans-[IrCl(CO)L₂] was recovered unchanged after treatment with sodium tetrahydroborate in ethanol.

The complex $[IrHCl_2(CO)L_2]$, configuration (IV), was prepared by treating the iridium(I) complex with hydrogen chloride. The i.r. spectrum showed bands at 2 261, 2 184, 2 035, 313, and 252 cm⁻¹ due to v(Ir-H), ν (C=C), ν (C=O), and ν (Ir-Cl) (two bands), respectively. As expected, the ¹H n.m.r. pattern showed a 1:2:1triplet hydride pattern and two t-butyl 1:2:1 triplets.

The complex *trans*-[RhCl(CO)L₂] was readily prepared from $[Rh_2Cl_2(CO)_4]$ and the phosphine. As with the iridium complex, this rhodium complex consisted of conformers in solution with two A2X and one ABX pattern in the ³¹P n.m.r. spectrum (Table 2) at 253 K. As before, one conformer is present in only small amounts, *i.e.* (II; X = CO, Y = CI), and one is preferred (II; X = CI, Y = CO).

Treatment of H_2 IrCl₆ with 4 mol equivalents of PBut₂(C=CPh) in refluxing propan-2-ol gives the five-12 A. J. Deeming and B. L. Shaw, J. Chem. Soc. (A), 1968,

1887. ¹³ H. D. Empsall, E. Mentzer, D. Pawson, B. L. Shaw, R. 1977 311. Mason, and G. A. Williams, J.C.S. Chem. Comm., 1977, 311.

¹⁰ A. Bright, B. E. Mann, C. Masters, B. L. Shaw, R. M. Slade, and R. E. Stainbank, *J. Chem. Soc.* (A), 1971, 1826.
¹¹ C. J. Moulton and B. L. Shaw, *J.C.S. Chem. Comm.*, 1976, 2017.

^{365.}

2289

co-ordinate $[IrHCl_2L_2]$ as a red solid. The ¹H n.m.r. spectrum shows two doublets due to the t-butyls and a high-field triplet at $\delta - 31.3$ p.p.m., with a large coupling constant of 21.6 Hz due to IrH. The ³¹P n.m.r. spectrum is a singlet. The i.r. spectrum shows a band at 2 178 cm⁻¹ due to v(C=C), two bands due to v(Ir-Cl) (at 304 and 268 cm⁻¹), but no band due to v(Ir-H). In iridium hydrides with very bulky tertiary phosphines a band due to v(Ir-H) is often very weak or not observed.³ We formulate this iridium(III) hydride complex as trigonal-bipyramidal (V) with the phosphines in the

configuration (VIII) was formed. The i.r. spectrum shows bands at 2 178, 1 922, and 1 525 cm⁻¹ due to v(C=C), v(C=O), and v(C=C) respectively. The ³¹P n.m.r. spectrum shows an AB pattern at δ 43.9 and 78.6 p.p.m. [J(PP) 293 Hz] showing *trans*-phosphines, and the ¹H n.m.r. spectrum four equally intense peaks centred at δ 1.46 p.p.m. due to the t-butyl hydrogens (separations *ca.* 2, 12.5, and 2 Hz) and a doublet of doublets for the vinyl hydrogen at δ 4.8 p.p.m. [J 3.0 and 5.0 Hz]. A rhodium complex was similarly prepared and characterized (see Table 2 for ³¹P n.m.r. and i.r. data). The vinyl



trigonal plane and *cis*-chlorides. In agreement with this formulation, the red complex (V) reacts readily with carbon monoxide to give the colourless complex (IV). We have made several other co-ordinatively unsaturated iridium(III) hydrides [IrHCl₂(PR₃)₂] (PR₃ is a very bulky tertiary phosphine such as PMeBu^t₂) but these have all had configuration (VI) and react with CO to give a sixco-ordinate carbonyl hydride of configuration (VII). Presumably the relatively small steric requirements of C=CPh allows the P-Ir-P angle to be considerably less than 180° as in (V), whereas ligands such as PEtBu^t₂ are preferentially *trans*, as in (VI).

When RhCl₃·3H₂O is treated with 4 mol equivalents of PBut₂(C=CPh) at 20 °C the red five-co-ordinate rhodium(III) hydride [RhHCl₂ L_2] is formed. The ¹H and ³¹P n.m.r. and i.r. evidence (Table 2) indicates that this complex probably has the square-pyramidal structure (VI) as found previously for complexes of the type $[RhHCl_2(PR_3)_2]$.¹⁴ It is not clear why the configuration should be different from that of the corresponding iridium complex described above. The complex $[RhHCl_{2}{PBu^{t}(C=CPh)}_{2}]$ slowly decomposes in chloroform solution to give an orange insoluble complex which appears to be a binuclear complex of Rh^{II} of type $[Rh_2Cl_4L_2]$. This complex was too insoluble for molecular-weight determinations. The ¹H n.m.r. spectrum of a deuteriochloroform solution of [RhHCl₂L₂] slowly changes at 34 °C. After 20 min a doublet at 8 1.42 p.p.m. (112.4 Hz) is just visible: this corresponds to the free phosphine L. However, the resolution soon deteriorates, which we interpret as being due to the formation of paramagnetic species [*i.e.* the rhodium(II) complex]. When $[RhHCl_2L_2]$ in benzene is treated with CO it rapidly gives trans-[RhCl(CO)L₂].

When the complex *trans*-[IrCl(CO)L₂] was treated with a solution of sodium propan-2-olate in propan-2-ol the chelate complex $[Ir(CO){PBut_2[CH=C(O)Ph]}L]$ of

¹⁴ C. Masters and B. L. Shaw, J. Chem. Soc. (A), 1971, 3679.

hydrogen formed a multiplet at δ 4.67 p.p.m. (CDCl₃). It is not clear whether small amounts of water in the



propan-2-ol-sodium propan-2-olate mixture were responsible for this hydration reaction or whether it involved breakdown of the propan-2-ol (or isopropoxide) possibly as shown in the Scheme. We have found for example



that the phosphine oxide $PBu_2^t(C\equiv CEt)O$ with sodium propan-2-olate gives $PBu_2^t[CH=C(OPr^i)Et]O$.¹⁵ Carty et al.¹⁶ showed that cis-[PdCl₂{PPh₂(C=CCF₃)}₂] reacts with water to give a variety of products including $[Pd{PPh_2[CH=C(O)CF_3]}_2]$ and suggested that the initial step is nucleophilic attack by water on the β -carbon followed by protonation and intramolecular hydrogenchloride elimination.

Treatment of trans-[IrCl(CO)L₂] with sodium methoxide in refluxing methanol gave [Ir(CO){PBut₂[CH=C(O)-Ph]}{PBut₂[CH=C(OMe)Ph]}] (trans-phosphorus donor atoms). This product was not obtained analytically pure, but the i.r. and ³¹P n.m.r. data (Table 2) and

¹⁵ D. Pawson and B. L. Shaw, unpublished work.

¹⁶ A. J. Carty, S. Jacobson, R. T. Simpson, and N. J. Taylor, J. Amer. Chem. Soc., 1975, **97**, 7254.

¹H n.m.r. data (below) agree with the formulation: four peaks due to the t-butyl hydrogens centred at δ 1.46 p.p.m., a singlet at 3.25 p.p.m. due to OMe, and sets of resonances due to the two vinylic protons at 4.83 [J(PH) 3.6 and 7.2] and 5.22 p.p.m. [J(PH) 2.4 and 6 Hz]. As was the case with reaction in propan-2-ol, the chelate ring might have been formed by a mechanism analogous to that shown in the Scheme or because of trace amounts of water in the sodium methoxide-methanol solution.

EXPERIMENTAL

All the operations involving $\mathrm{PBut}_2(C\equiv CPh)$ were carried out under argon. The analytical and spectroscopic data are recorded in the Tables.

(Phenylethynyl)di-t-butylphosphine.n-Butyl-lithium (0.153 mol) in hexane (82 cm³) was added dropwise over 0.5 h to a well stirred solution of ethynylbenzene (15.6 g, 16.8 cm³, 0.153 mol) in diethyl ether (50 cm³). The resulting white suspension was heated under reflux for 2 h then added over 10 min to a cooled (-10 °C) solution of chlorodi-tbutylphosphine (18.4 g, 0.102 mol) in diethyl ether (70 cm³). The mixture was then heated under reflux for 20 h and hydrolyzed with water. The product was isolated by distillation, b.p. 188 °C (bath temperature) (27 mmHg),* white prisms, m.p. 46–49 °C, yield 80% (Found: C, 77.6; H, 9.2. $C_{20}H_{15}P$ requires C, 78.0; H, 9.4%), ν (C=C) in Nujol at 2 181s cm⁻¹. N.m.r. spectra in CDCl₂: ³¹P. δ 12.6 p.p.m.; ¹H, δ 1.28(d) p.p.m. [³J(PH) 12.5 Hz], But. With methyl iodide in acetone the compound readily gave the methiodide salt as needles, m.p. 150-155 °C, yield 69% (Found: C, 52.3; H, 6.8; I, 32.4. C21-H₁₈IP requires C, 52.6; H, 6.75; I, 32.7%).

trans-Dichlorobis[(phenylethynyl)di-t-butylphosphine]platinum(II).—A suspension of dichlorobis(t-butyl cyanide)platinum(II) (1.26 g, 2.67 mmol) in toluene (10 cm³) was treated with PBut₂(C \equiv CPh) (1.39 g, 5.60 mmol) and the mixture was stirred for 5 min at 20 °C. The resulting clear solution was diluted with light petroleum (b.p. 80—100 °C) to give the product (1.75 g, 2.30 mmol) as yellow prisms. trans-Dichlorobis[(phenylethynyl)di-t-butylphosphine]-

palladium(II) was prepared similarly from dichlorobis-(benzonitrile)palladium(II) as orange prisms from toluenelight petroleum (b.p. 80—100 °C).

trans-Iodobis[(phenylethynyl)di-t-butylphosphine]plati-

num(II).—A solution of trans-[PtCl₂{PBut₂(C≡CPh)}₂] (0.08 g, 0.11 mmol) in 2-methoxyethanol (5 cm³) containing sodium iodide (0.174 g, 1.1 mmol) was heated under reflux for 3.5 h. The resulting yellow solution was evaporated and the residual oil extracted with toluene. The extract was diluted with light petroleum (b.p. 80—100 °C) to give the product (0.094 g, 0.10 mmol) as yellow prisms.

trans-Iodobis[(phenylethynyl)di-f-butylphosphine]palladium(II).—A solution of trans-[PdCl₂{PBut₂(C≡CPh)}₂] (0.24 g, 0.35 mmol) in acetone (7 cm³) containing NaI (0.54 g, 3.54 mmol) was stirred for 16 h at 20 °C. The mixture was then cooled to ca. -15 °C giving the product as red-brown prisms. These were washed with water and dried, yield 0.27 g (0.32 mmol).

 $Di-\mu-chloro-bis\{chloro[(phenylethynyl)di-t-butylphosphine]-palladium(II)\}.$ A solution of $PBut_2(C\equiv CPh)$ (0.45 g, 1.81 mmol) in methanol (5 cm³) was added to a solution of

* Throughout this paper: 1 mmHg \approx 13.6 \times 9.8 Pa.

disodium tetrachloropalladate(II) (0.64 g, 30.1% Pd, 1.81 mmol) in methanol (8 cm³). The resulting suspension was stirred for 16 h at 20 °C. The *product* (0.61 g, 1.45 mmol) was filtered off as orange prisms.

trans-Dichlorobis[(2-chloro-2-phenylvinyl)di-t-butylphosphine]platinum(II).—A solution of trans-[PtCl₂{PBut₂-(C=CPh)₂] (0.10 g, 0.13 mmol) in dichloromethane (5 cm³) was treated with a solution of hydrogen chloride (13.2 mmol) in dry diethyl ether (3 cm³). The suspension was stirred for 16 h at 20 °C. The resulting clear solution was cooled to ca. 15 °C giving the product (0.083 g, 0.10 mmol) as white prisms.

cis-Dimethylbis[(phenylethynyl)di-t-butylphosphine]platinum(II).—(Phenylethynyl)di-t-butylphosphine (0.098 g, 0.40 mmol) was added to a solution of cis-(cyclo-octa-1,5-diene)dimethylplatinum(II) (0.064 g, 0.20 mmol) in benzene (2 cm³). The mixture was stirred for 15 min at 20 °C to give a clear solution, and the solvent was removed

under reduced pressure. The resulting oil was treated with light petroleum (b.p. 60—80 °C) and the *product* (0.11 g, 0.15 mmol) formed as white prisms. trans-Chloro(methyl)bis[(phenylethynyl)di-t-butylphosphine]platinum(II).—A solution of cis-chloro(cyclo-octa-1,5-diene)methylplatinum(II) (0.082 g, 0.23 mmol) in benzene (1 cm³) was treated with PBu^t₂(C≡CPh) (0.12 g, 0.46 mmol). The solvent was removed under reduced

0.46 mmol). The solvent was removed under reduced pressure. The resulting oil was treated with light petroleum (b.p. 60-80 °C) and the *product* (0.13 g, 0.17 mmol) formed as white prisms.

Dichloro(cyclo-octa-1,5-diene)[(phenylethynyl)di-t-butylphosphine]platinum(II).—A suspension of cis-dichloro(cycloocta-1,5-diene)platinum(II) (0.19 g, 0.52 mmol) in benzene (5 cm³) was treated with PBut₂(C=CPh) (0.13 g, 0.52 mmol). The mixture was stirred for 30 min at 20 °C. The resulting colourless solution was evaporated and the residual oil extracted with methanol. The product formed as white prisms (0.15 g, 0.24 mmol).

trans-(Carbonyl)chlorobis[(phenylethynyl)di-t-butylphosphine]iridium(I).-A solution of hexachloroiridic(IV) acid (1.10 g, 42.8% Ir, 2.44 mmol) in absolute ethanol (20 cm³) and rhodium trichloride trihydrate (0.007 g, 0.03 mmol) was stirred in a carbon monoxide atmosphere for 24 h under slow reflux. (Phenylethynyl)di-t-butylphosphine (1.26 g, 5.13 mmol) was then added to the resulting vellow solution. The suspension was then stirred for 2 h at 20 °C. Filtration gave a cream solid (1.54 g) which was found to be a ca. 1:1mixture of trans-[IrCl(CO){PBut2(C=CPh)}2] and carbonyldichlorohydridobis[(phenylethynyl)di-t-butylphosphine]iridium(III) (see text). A suspension of the mixture (1.54 g) in absolute ethanol (3 cm^3) was treated with triethylamine (0.12 g, 1.20 mmol) and the mixture was heated under reflux for 5 min then cooled to ca. -15 °C giving the product (1.43 g, 1.90 mmol) as yellow prisms.

trans-Carbonyldichlorohydridobis[(phenylethynyl)di-t-butylphosphine]iridium(III).—A solution of trans-[IrCl(CO)-{PBut₂(C \equiv CPh)₂] (0.16 g, 0.22 mmol) in dichloromethane (3 cm³) was treated with a solution of hydrogen chloride (0.24 mmol) in diethyl ether (0.75 cm³). The resulting colourless solution was diluted with absolute ethanol (3 cm³) then cooled to ca. -15 °C giving the product (0.066 g, 0.085 mmol) as white prisms.

trans-Carbonylchlorobis[(phenylethynyl)di-t-butylphos-

0.49 mmol Rh). The required *product* (0.20 g, 0.31 mmol) separated as yellow prisms.

Dichlorohydridobis[(phenylethynyl)di-t-butylphosphine]iridium(III).—A solution of H_2IrCl_6 (0.44 g, 0.97 mmol) in propan-2-ol (10 cm³) was heated for 5 min until it became brownish green. The solution was cooled and PBut₂-(C \equiv CPh) (0.98 g, 3.97 mmol) added under argon. The resulting green slurry was then heated under reflux for 5 h. The suspension obtained was filtered and the filtrate was cooled to *ca.* -15 °C giving the *product* (0.11 g, 0.15 mmol) as red prisms.

Dichlorohydridobis[(phenylethynyl)di-t-butylphosphine]rhodium(III).—(Phenylethynyl)di-t-butylphosphine (0.76 g, 3.1 mmol) was added to a solution of $RhCl_3\cdot 3H_2O$ (0.19 g, 0.77 mmol) in absolute ethanol (15 cm³). The resulting deep red solution was stirred for 20 min at 20 °C then cooled to *ca.* -15 °C giving the *product* (0.23 g, 0.35 mmol) as red prisms.

 Di_{μ} -chloro-bis{chloro[(phenylethynyl)di-t-butylphosphine]rhodium(II)}.—A solution of [RhHCl₂{PBut₂(C=CPh)₂] (0.11 g, 0.16 mmol) in chloroform (1 cm³) was stirred for 16 h at 20 °C. The product was filtered off as orange prisms (0.03 g, 0.07 mmol).

Carbonyl[2-(di-t-butylphosphino)-1-phenylethyleneoxo-

in propan-2-ol (11.4 cm³, 0.14 mol dm⁻³) under reflux for 16 h. The resulting suspension was evaporated to an oil which was then treated with water. The *product* (0.13 g, 0.18 mmol) was filtered off as yellow prisms.

Carbonyl [2-(di-t-butyl phosphino)-1-phenylethyleneoxo-

OP][(phenylethynyl)di-t-butylphosphine]rhodium(1).—A suspension of trans-[RhCl(CO){PBut}_2(C=CPh)}_2] (0.11 g, 0.16 mmol) in a solution of sodium propan-2-olate in propan-2-ol (9.4 cm³, 0.1 mol dm⁻³) was heated under reflux for 16 h. The resulting suspension was evaporated to an oil which was then treated with water. The product (0.10 g, 0.15 mmol) was filtered off as yellow prisms.

Carbonyl[2-(di-t-butylphosphino)-1-phenylethyleneoxo-OP][(2-methoxy-2-phenylvinyl)di-t-butylphosphine]iridium(I). $--The complex trans-[IrCl(CO){PBut}_2(C=CPh)}_2] (0.15 g, 0.20 mmol) was treated with a solution of sodium methoxide in methanol (2.1 cm³, 0.8 mol dm⁻³) under reflux for 22 h. The resulting suspension was then evaporated to an oil in vacuo and water was added. The product (0.14 g, 0.18 mmol) was filtered off as yellow prisms and could be recrystallized from dichloromethane-methanol.$

We thank Johnson, Matthey Ltd. for a generous loan of platinum metal salts, the S.R.C. for the award of a research assistantship (to E. M. H.), and I.C.I. Ltd. for the award of a fellowship (to H. D. E.).

[7/918 Received, 27th May, 1977]