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COMPLEXES OF PLATINUM(II) AND PALLADIUM(II) WITH HYBRID PHOSPHINE-PHOSPHINE OXIDE LIGANDS OF TYPE $Ph_2P(CH_2)_nP(=O)Ph_2$ (n = 1, 2, 3, OR 4) *

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Abstract

Treatment of $[PtCl_2(NCBu^1)_2]$ or $[PtCl_2(cyclo-1,5\text{-octadiene})]$ with $Ph_2P(CH_2)_n P(=O)Ph_2$ (n = 1, dppmO; n = 2, dppeO; n = 3, dpppO; n = 4, dppbO) gives complexes of the type cis- $[PtCl_2(Ph_2P(CH_2)_nP(=O)Ph_2]_2]$, independently of mole ratio. Treatment of cis- $[PtCl_2(dppmO)_2]$ with LiBr or NaI gives $[PtX_2(dppmO)_2]$ (X = Br, cis; X = I, cis / trans mixture) and on treatment with $Hg(C=CPh)_2$ in ethanol, trans- $[Pt(C=CPh)_2(dppmO)_2]$ is formed. Treatment of these ligands with $[PtMe_2(cyclo-octa-1,5\text{-diene})]$ gives exclusively cis- $[PtMe_2\{Ph_2P(CH_2)_nP(=O)Ph_2\}_2]$. Coordination of the P=O group was achieved by treating $[PtCl_2(Ph_2P-(CH_2)_nP(=O)Ph_2]_2]$ with either AgNO₃ or TIPF₆, whereupon cis- $[Pt(Ph_2P-(CH_2)_nP(=O)Ph_2-P,O]_2]^{2+}$ (n = 1, or 2) is formed: these cations were also synthesized by treating $[PtMe_2\{Ph_2P(CH_2)_nP(\approx O)Ph_2\}_2]$ with dppmO gives trans- $[PdCl_2(dppmO)_2]$ only and this reacts with AgNO₃ or TIPF₆ to form cis- $[PdCl_2(dppmO)_2]^{2+}$. The complexes were characterized by microanalysis, IR and NMR spectroscopy.

Although the coordination chemistry of phosphine oxides has been examined for 3d-metals [1], lanthanides [2], and actinides [3] and for some 4d- and 5d-metals in higher oxidation states [4,5], relatively little has been reported on phosphine oxides as ligands for the platinum metals. The weakly coordinating perchlorate ligand in complexes of the type $[M(OClO_3)(C_6F_5)(AsPh_3)_2]$ (M = Pt or Pd) has been displaced by Ph₃P=O [6,7], and complexes of the type $[Pt(X)(O=PPh_3)(PEt_3)_2]^+$ have been synthesized in a similar way [8,9]. Better defined palladium(II) complexes of the type $[PdX_2(O=PRR'_2)_2]$ (R = R' = Ph; X = Br or I. R = Et, R' = Ph; X = Cl or I) were reported as the products of heating the corresponding phosphine complexes in air [10]. A study of the extraction of Pd(NO₃)₂ from aqueous solutions, using

^{*} Dedicated to Professor G.E. Coates on the occasion of his seventieth birthday.

trialkylphosphine oxides, has been made [11] and there have been brief claims of the synthesis of paramagnetic $Pd-Ph_3P=O$ complexes [12,13]. 2-Pyridyldiphenylphosphine oxide has been shown to act as a chelate for platinum(IV) in $[PtBr_4(2-pyridyldiphenylphosphine oxide)]$ [14], which is the only structurally characterized phosphine oxide complex of a platinum metal. It seemed of interest to examine the behaviour of mixed tertiary phosphine/tertiary phosphine oxides as ligands towards platinum(II). We, of course, expected that the tertiary phosphine would bind strongly to the platinum but it was possible that the P=O might also be induced to coordinate because of the chelate effect, provided that the chelate ring was of a suitable size. Alternatively, the P=O group might interact only weakly with the platinum.

Results and discussion

We here report on a study of the four hybrid phosphine-phosphine oxides of type $Ph_2P(CH_2)_nP(=O)Ph_2$, n = 1, 2, 3, or 4 as ligands towards some platinum(II) moieties and the synthesis of one platinum complex. Recently a useful synthesis of such compounds has been reported which involves treatment of the diphosphines $Ph_2P(CH_2)_nPPh_2$ with benzyl bromide in benzene to give the monoquaternary salt, $[Ph_2P(CH_2)_nP(CH_2Ph)Ph_2]Br$, which when treated with aqueous alkali gives $Ph_2P(CH_2)_nP(CH_2Ph)Ph_2$ and toluene [15]. In our hands this method gave good results for n = 2, 3, or 4 but a poor yield for n = 1. However, we devised a good method of synthesising $Ph_2PCH_2P(=O)Ph_2$ (dppmO) which was to treat the commercially available diphenyl(methyl)phosphine oxide with n-butyllithium in THF and treat the resultant carbanion with diphenylchlorophosphine. This gave dppmO in over 60% yield: details are in the Experimental.

Treatment of either $[PtCl_2(NCBu^t)_2]$ or $[PtCl_2(COD)]$ with dppmO (2/1, 1/1, 1/2 mole ratios) in CDCl₃, gave only *cis*- $[PtCl_2(dppmO)_2]$ (1a). The reactions were



Complex	Analyses (Found (calcd.)(%)) ^b			ν (P=O) (cm ⁻¹) ^c	Other bands (cm ⁻¹)		
	СНИ		Halogen				
$1a \cdot 0.2 \text{ CH}_2\text{Cl}_2$	55.1(55.5)	4.25(4.1)	7.2(7.2)	1180m	320, 293, v(Pt-Cl)		
1b · 1 CHCl ₃	47.9(48.0)	3.7(3.8)	21.7 (20.3) ^d	1185m			
lc	48.0(48.0)	3.6(3.7)	20.2(20.3)	1195m			
1d · 1 C ₆ H ₆	63.1(63.1)	4.9(5.1)		1180m			
2b	64.4(64.4)	4.5(4.6)		1195s	2100m, v(C≡C)		
2c	58.6(58.5)	4.7(4.5)	3.6(3.4)	1190s	265, v(Pt-Cl)		
1e	56.6(57.0)	4.3(4.4)	6.7(6.5)	1195s	312, 290, v(Pt-Cl)		
1f	61.8(61.5)	5.2(5.2)		1200s			
1g	56.7(57.8)	4.5(4.7)	8.35(6.3)	1185s	314, 291, v(Pt-Cl)		
1ĥ	62.1(61.2)	5.6(5.4)		1185br			
1i	see text			1185m			
2d	60.6(61.4)	4.5(4.5)	7.2(7.25)	1195m	343s, v(Pd-Cl)		
3a	49.5(50.2)	3.5(3.7)		1130s, 1120s			
3b	53.0(53.6)	4.0(4.0)	2.65(2.5) ^e	1108s, 1128s			
3d	46.5(47.5)	3.5(3.7)	16.9(17.4)				

 TABLE 1

 INFRARED ^a AND ANALYTICAL DATA

^{*a*} As Nujol mull. ^{*b*} The presence of some solvent of crystallization in some of the complexes was confirmed by ¹H NMR spectroscopy. ^{*c*} m = medium, s = strong, br = broad. ^{*d*} Halogen (total). ^{*e*} % nitrogen.

monitored by ³¹P-{¹H} NMR spectroscopy and no evidence for chelation of the dppmO was found under these conditions. The product was isolated by precipitation with diethyl ether as a white solid. Its structure la follows from (i) microanalytical data (Table 1) (ii) two strong IR absorbtion bands, at 293 and 320 cm⁻¹, assigned to ν (Pt–Cl) and absent from the spectra of the corresponding bromide or iodide complexes and a band at 1180 cm⁻¹, due to uncoordinated P=O, (iii) the ³¹P-{¹H} NMR spectrum (Table 2), which shows ¹J(Pt-P) 3710 Hz, typical of P trans to Cl, and a resonance due to P^{v} at 25.4 ppm, shifted very little from the value for free dppmO (29.7 ppm). Interestingly, for 1a the value of ${}^{3}J(Pt-P)$ is 103 Hz, whereas with $[PtBr_4(2-pyridylphosphine oxide)]$, $Pt-P^V$ coupling was not observed, even though the phosphine oxide is coordinated [14]. The ¹H-{³¹P} NMR spectrum shows a singlet for the methylene protons, with platinum satellites, ${}^{3}J(Pt-H)$ 25.2 Hz. In the ¹H NMR spectrum, there is coupling to both P and P=O (Table 3). The corresponding dibromide is similar, see Tables and Experimental, but although the iodide was isolated as the cis-isomer, in solution it partially isomerized to the trans-isomer, as evidenced by ${}^{31}P{}^{1}H$, ${}^{1}H$ and ${}^{1}H{}^{31}P{}$ NMR spectroscopy (Tables 2 and 3): equilibrium is reached after 3 h, in CDCl₂ at 20°C, and the ratio of trans/cis is 7/3.

We have also prepared some organometallics containing dppmO ligands. Treatment of *cis*-[PtCl₂(dppmO-*P*)₂] with Hg(C=CPh)₂ in hot ethanol gave *trans*-[Pt(C=CPh)₂(dppmO)₂] (**2b**) in 65% yield. This formulation follows from (i) the analytical data (Table 1) and the *trans*-configuration from the single band due to ν (C=C) at 2100 cm⁻¹ (Table 1) (ii) the ³¹P-{¹H} NMR spectrum, in which ¹J(Pt-P) is 2600 Hz, typical of *P trans* to *P* and (iii) the ¹H NMR spectrum, in which the methylene protons show a doublet of triplets with Pt satellites. The doublet (*J*(P-H) 12 Hz) is due to coupling to P=O and the triplet is a "virtual triplet" due

Complex	δ(P)	$\delta(PO)$	$^{1}J(PtP)$	$^{3}J(PtPO)$	N ^b	Solvent ^c
1a	1.5	25.4	3710	103	12	Α
1b	1.4	25.2	3664	95	17	Α
f 1c	-2.3	26.6	3470	102	10	Α
2a	-1.8	27.1	2497	49	n.r.	Α
1d	13.6	25.4	1890	39	10	В
2b	5.5	24.9	2600	95	12	С
2c	18.3	25.5	3110	90	13	С
le	10.0	32.9	3660	~ 0	57	С
1f	19.1	28.9	1880	~ 0	49	С
lg	70.5	32.6	3675	~ 0	~ 0	С
1h	16.6	30.6	1882	~ 0	~ 0	С
1i	17.4	31.5	1890	~ 0	~ 0	С
2a	11.8	24.4			~ 0	Α
3a	36.0	59.2			d	D
3b	3.3	64.7	3877	72		С
3c	3.6	66.7	3850	76		С
3d	1.9	50.8	3655	~ 0	~ 0	C

^{*a*} Chemical shifts (δ) in ppm ; ± 0.1 ppm relative to 85% H₃PO₄ (positive shift to high frequency; coupling constants (*J*) in Hz (± 2 Hz). Spectra were recorded at 25°C, unless stated otherwise. ^{*b*} N = |²J(PPO)+⁴J(PP'O)|. ^cA, CDCl₃; B, C₆D₆; C, CD₂Cl₂; D, (CD₃)₂CO. ^{*d*}²J(PPO) 5Hz.

to the mutually *trans* and strongly coupled Pt-bonded P-atoms (see Table 3). In this and other *trans*-isomers the methylene protons absorb at lower field than the methylenes of the *cis*-isomers.

TABLE 3 ¹H AND ¹H-{³¹P} NMR ^a DATA

Com- plex	$\delta(CH_2)$	$^{3}J(PtCH_{2})$	$^{2}J(PCH_{2})$	$J\{P(=O)CH_2\}$	δ(CH ₃)	$^{2}J(\text{PtCH}_{3})$	N ^b	Sol- vent
la	3.78	25.2	~ 6	~12				A
1b	3.82	24.6	12.0	12.0				Α
1c)	3.88	24.9	~12	~ 12				Α
2a)	4.49	26.1	~ 10 ^d	11.7				Α
1d	3.54	16.1	7.8	12.0	0.51	69	14.0	В
2b	4.36	24.0	8.0 ^d	12.0				Α
2c	4.22	18.1	7.8 ^d	11.0	-0.29	40	14.2	Α
le	2.64	n.r. ^e	n.r.	n.r.				Α
1f	2.82	n.r.	n.r.	n.r.	1.33	68	13.8	В
1g	2.3–1.9m ^f							
1h	1.6-1.4m					65	13.8	В
li	1.6-1.4m				0.43	65	13.2	А
2d	3.88	_	7.8 ^d	11.3				
3a	4.90		9.8	13.9				С
3c	4.76	28.8	9.8	13.4				С
3d	3.23m	-	-	-				С

^{*a*} Spectra measured at 100 MHz. Chemical shifts (δ) in ppm (\pm 0.01) to high frequency of SiMe₄ and coupling constants (*J*) in Hz. (\pm 0.1) except for ²*J*(*P*tCH₃) (\pm 1 Hz). ^{*b*} *N* = |³*J*(PCH₃)_{*trans*} + ³*J*(PCH₃)_{*cis*}|. ^{*c*} A = CDCl₃; B = C₆D₆; C = (CD₃)₂CO. ^{*d*} |²*J*(PCH₂)+⁴*J*(PCH₂)|. ^{*e*} n.r. = not resolved. ^{*f*} m = multiplet.

TABLE 2

³¹P-{¹H} NMR DATA^a

The action of dppmO on $[PtMe_2(COD)]$ was studied by ³¹P-{¹H} NMR spectroscopy in CDCl₃ or C₆D₆: in both cases only *cis*- $[PtMe_2(dppmO-P)_2]$ (**1d**) was formed. This complex was prepared and isolated from toluene solution: characterizing data are in the Tables. When treated with dry HCl (1 mol) **Id** was converted into the *trans*-mono-methyl complex (**2c**), for which characterizing data are in the Tables. We hoped that the dimethyl complex (**1d**) would undergo oxidative addition reactions but when a solution of this complex (0.067 *M*) in dichloromethane was treated with MeI (0.27 *M*), no reaction occurred, even after 2 weeks at room temperature.

The homologous ligands dppeO and dpppO were also studied. These with $[PtCl_2(NCPh)_2](dppeO)$ or $[PtCl_2(COD)](dpppO)$ gave exclusively the *cis*-monodentate complexes 1e and 1g, respectively. Similarly the dimethylplatinum complexes 1f, 1h and 1i were prepared by treating $[PtMe_2(COD)]$ with dppeO, dpppO, or dppbO, respectively. These various complexes were more difficult to purify than the dppmO analogues and 1i was only characterized by IR and NMR spectroscopy and not be elemental analysis.

Attempts to effect P=O coordination to platinum(II) or palladium(II) treating the halo complexes of type 1 with large coordinating anions such as BPh₄ or BF₄, failed. However, by treating the chloro complexes of type 1 with silver or thallium salts to remove chloride, P=O coordination was achieved. Thus treatment of a solution of the palladium complex 2d with silver nitrate gave a precipitate of silver chloride and



addition of ammonium hexafluorophosphate then gave the dppmO-PO chelate 3a. The corresponding platinum bis-chelate was isolated as the nitrate salt 3b and as the hexafluorophosphate salt 3c, although the nitrate salt 3b gave satisfactory elemental analytical data the carbon analysis for the hexafluorophosphate complex 3c was low and the fluorine analysis high (Table 1), however, the ³¹P-{¹H} NMR data indicate the cations to have the *cis*-structure shown in 3b and 3c in each case. The bis-chelated dppeO salt 3d was also prepared. For the complexes 3b, 3c and 3d the values of ¹J(PtP) are large, 3877, 3850 and 3655 Hz, respectively, and what one would expect for P *trans* to a hard donor such as O=P. The chemical shift of the coordinated P=O phosphorus is shifted downfield by ca. 35-40 ppm from the shift of the uncoordinated phosphorus. No coupling was observed between Pt and P=O for the dppeO-PO chelate 3d, showing that ²J(Pt-PO) is very small and suggesting that coupling between Pt and P in the dppmO chelate 3c (76 Hz) or 3b (72 Hz) is mainly a 3-bond coupling. As might be expected 3c reacted with Et₄NI in CH₂Cl₂

to give $[PtI_2(dppmO-P)_2]$ (NMR evidence) although this could not be isolated pure.

Treatment of the dpppO chloro complex 1g with either silver nitrate or thallium hexafluorophosphate gave only dark intractable oils, presumably because the sevenmembered ring chelate produced is not stable. Treatment of the dimethyl compound 1h with HPF₆ also led to decomposition; with 1d or 1f, treatment with HPF₆ gave 3c or 3d, respectively $\binom{31}{1}P-\binom{1}{1}H$ NMR spectroscopic evidence).

Experimental

The techniques and instrumentation used were as described in other recent papers from this laboratory [16].

Diphenylphosphino(diphenylphosphonyl)methane (dppmO). A solution of n-butyllithium in hexane (13.5 cm³, 1.7 M, 23 mmol) was added at -78° C with stirring to a solution of diphenylmethylphosphine oxide (4.9 g, 22.9 mmol). The solution was allowed to warm up to room temperature and then after 10 min recooled to -20° C and diphenylchlorophosphine (5.08 g, 23 mmol) added to it, with stirring, over 20 min. The mixture was then warmed to ca. 20° C and stirred for 2 h, after which it was hydrolysed with degassed aqueous ammonium chloride. The required product was isolated with ether and formed white prisms from propan-2-ol. Yield 6.0 g, 64%.

 $cis-[PtCl_2(dppmO-P)_2]$ (1a). A solution of dppmO (0.20 g, 0.25 mmol) in dichloromethane (5 cm³) was added to a solution of $[PtCl_2(NCBu^t)_2]$ (0.108 g, 0.25 mmol) in acetonitrile (10 cm³). The mixture was then stirred for 30 min, after which it was evaporated under reduced pressure to ca. 4 cm³ and diethyl ether added slowly; this gave the required product as white microcrystals. Yield 0.21 g, 79%.

 $cis-[PtBr_2(dppmO-P)_2]$ (1b). A mixture of $cis-[PtCl_2(dppmO)_2]$ (0.20 g, 0.188 mmol) and lithium bromide (0.3 g) in acetone (30 cm³) was stirred for 3 h and then evaporated to dryness. The required product was extracted into chloroform and formed yellow microcrystals from chloroform/propan-2-ol. Yield 0.18 g, 83%.

trans- $[PtI_2(dppmO-P)_2]$ (2a). 2a was prepared similarly and formed yellow microcrystals. Yield 83%.

trans-[Pt(C=CPh)₂(dppmO-P)₂] (2b). A mixture of Hg(C=CPh)₂ (0.075 g, 0.188 mmol) and cis-[PtCl₂(dppmO)₂] (0.20 g, 0.188 mmol) in ethanol (10 cm³) was heated under reflux for 15 min. The mixture was then filtered to remove HgCl₂ and evaporated to dryness under reduced pressure, giving the required product, which formed yellow microcrystals from dichloromethane/diethyl ether. Yield 0.146 g, 65%.

 $cis-[PtMe_2(dppmO-P)_2]$ (1d). dppmO (0.20 g, 0.50 mmol) was added to a solution of [PtMe_2(COD)] (0.083 g, 0.25 mmol) in toluene (5 cm³) at ca. 50 °C. The resultant yellow solution was slowly cooled to -20 °C, giving the required product as white prisms. Yield 0.224 g, 87%.

trans-[PtClMe(dppmO-P)₂] (2c). A solution of HCl (0.188 mmol) in methanol (1 cm³) was generated by the addition of acetyl chloride (12.9 μ l, 0.188 mmol). This solution was then added to a solution of cis-[PtMe₂(dppmO-P)₂] (0.193 g, 0.188 mmol) in dichloromethane (3 cm³). The required product was isolated and formed a white powder from dichloromethane/n-hexane. Yield 0.16 g, 81%.

 $cis-[PtCl_2(dppeO-P)_2]$ (1e). dppeO (0.87 g, 2.1 mmol) and $[PtCl_2(PhCN)_2]$ (0.47 g, 1.0 mmol) were heated together in refluxing benzene (25 cm³) for 2 h. The mixture was then evaporated to dryness and the residue triturated with diethyl

ether, to give the required product, which formed white microcrystals from chloro-form/diethyl ether. Yield 0.91 g, 83%.

 $cis-[PtCl_2(dpppO-P)_2]$ (1g). dpppO (0.45 g, 1.05 mmol) was added to a suspension of $[PtCl_2(COD)]$ (0.187 g, 0.5 mmol) in benzene (12 cm³). The mixture was heated under reflux for 1 h, after which the resultant yellow solution was decanted from the sticky residue. The solution was evaporated to dryness and the residue triturated with diethyl ether and then dissolved in dichloromethane. n-Hexane was added, at intervals of 2 h to this solution, at 4°C, to give the required product as a white powder. Yield 0.32 g, 57%.

 $cis-[PtMe_2(dppeO-P)_2]$ (1f). A solution of dppmO (0.87 g, 2.1 mmol) in warm benzene (20 cm³) was added to a solution of [PtMe₂(COD)] (0.353 g, 1.0 mmol) in benzene (20 cm³). The mixture was stirred for 16 h, then evaporated to dryness and the residue triturated with pentane, giving the required product, which was obtained as a white powder from benzene/hexane. Yield 0.88 g, 84%.

cis-[PtMe₂(dpppO-P)₂] (1h). A mixture of dpppO (0.87 g, 2.05 mmol) and [PtMe₂(COD)] (0.37 g, 1.0 mmol) in benzene (25 cm³) solution was warmed (40 °C) for 30 min. The mixture was evaporated to dryness, giving a glass, which was triturated with diethyl ether/n-hexane and then stored under diethyl ether at -20 °C. This produced the required product as a white powder. Yield 0.54 g, 48%. Attempts at recrystallization were unsuccessful.

cis-[PtMe₂(dppbO-P)] (1i). This was prepared similarly and formed white microcrystals from dichloromethane/cyclohexane. Yield 57%.

 $[Pd{Ph_2PCH_2P(=O)Ph_2}_2][PF_6]_2$ (3a). A solution of silver nitrate (0.10 g, 0.6 mmol) in ethanol (5 cm³)/water (0.5 cm³), was added to a solution of 2d (0.25 g, 0.25 mmol) in dichloromethane (2 cm³). The resultant mixture was stirred for 1 h, in the dark, and then the silver salts were filtered off and the filtrate evaporated to ca. 3 cm³. A solution of ammonium hexafluorophosphate (0.098 g, 0.6 mmol) in ethanol (18 cm³) was then added to give a white precipitate. This was collected etc. and recrystallized from acetone/diethyl ether, to give the required product as white microcrystals. Yield 0.13 g, 42%.

 $[Pt{Ph_2PCH_2P(=O)Ph_2-PO}_2][NO_3]_2$ (3b). A suspension of 1a (0.27 g, 0.25 mmol) in acetone (20 cm³) was treated with a solution of silver nitrate (0.10 g, 0.6 mmol) in methanol (9 cm³)/water (1 cm³). The resultant mixture was heated to ca. 50 °C for 12 h and then evaporated to dryness. The required product was isolated with dichloromethane and formed white microcrystals from dichloromethane/diethyl ether. Yield. 0.20 g, 72%.

 $[Pt{Ph_2PCH_2P(=O)Ph_2-PO}_2][PF_6]$ (3c). A solution of TIPF₆ (0.128 g, 0.52 mmol) in acetone (10 cm³) was added, with stirring to a solution of **1a** (0.25 g, 0.23 mmol) in dichloromethane (10 cm³). After 2 h the solution was filtered from thallous salts and evaporated to dryness. The required product was extracted into acetone from which it was precipitated with diethyl ether. It formed white microcrystals from dichloromethane/chlorobenzene. Yield 0.21 g, 74%.

 $[Pt{Ph_2P(CH_2CH_2)P(=O)Ph_2-PO}_2][PF_6]_2$ (3d). A solution of silver nitrate (0.20 g, 1.2 mmol) in ethanol (20 cm³)/water (0.5 cm³) was added to a solution of 1e (0.55 g, 0.5 mmol) in dichloromethane (4 cm³), in the dark. The mixture was then heated under reflux for 1 h, filtered and the filtrate evaporated to dryness. The residue was extracted with ethanol (15 cm³) and the extract treated with a solution of NH₄PF₆ (0.19 g, 1.15 mmol) in ethanol (9 cm³)/water (1 cm³). The resultant

white precipitate was recrystallized from acetone/diethyl ether to give the required product. Yield 0.49 g, 75%.

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