Reaction of *P*-Halogeno-phospha-alkenes with Alkene Complexes of Nickel and Platinum: η^2 -Co-ordination and Unusual Oxidative Addition Behaviour

D. Gudat,^a M. F. Meidine,^b J. F. Nixon,^{b*} and E. Niecke^{a*}

 Institut f
ür Anorganische Chemie, Universit
ät Bonn Gerhard-Domagk-Str. 1, D-5300 Bonn 1, Federal Republic of Germany

^b School of Chemistry and Molecular Sciences, University of Sussex, Falmer, Brighton BN1 9QJ, U.K.

Phospha-alkenes X–P=CTMS₂ (X = F, Cl, I; TMS = SiMe₃) display different reaction behaviour towards metal complexes (Bu₃P)₂Ni(cod) (cod = cyclo-octa-1,5-diene) or (Ph₃P)₂Pt(C₂H₄), respectively, undergoing either co-ordination to give η^2 -phospha-alkene complexes or alternatively oxidative addition of the P–X bond, forming phospha-alkenyl-metal(u) complexes, [(R₃P)₂M(X)(σ -P=CTMS₂)].

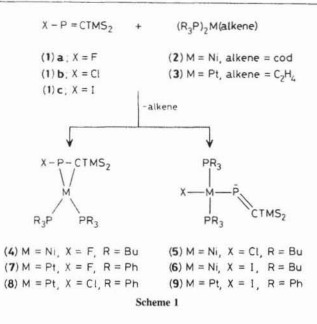
Phospha-alkenes react with a wide variety of transition metal compounds *via* ligand exchange to give complexes in which the phospha-alkenes may act as 2e- or 4e-donors and bonding of the ligand to the metal centre can occur through the phosphorus lone pair (η^1 -co-ordination), the π -electrons of

the double bond (η^2 -co-ordination), or a combination of both.¹ A different type of interaction, however, was observed during the reaction of Me₅C₅-P=C(SiMe₃)₂ with complexes of type [L₃M(CO)₃] (M = Mo, W; L = MeCN)² or [(R₃P)Ni-(alkene)₂],³ which proceeded *via* oxidative addition of the

Table 1. ³¹ P(¹ H) and ¹⁹ F n.m.r. data of complexes $[(\eta^2 - X - P_m = CTMS_2)M(P_{a,b}R_3)_2]$ and $[(P_aR_3)_2M(P_a)_2M$	$(X) (\sigma - P_x = CTMS_2)] (4-9).$
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	$\delta^{31}P^a$	${}^{1}J_{\mathrm{Pt,P}}^{}\mathfrak{b}$	${}^{2}J_{\mathbf{p},\mathbf{p}}\mathbf{b}$	$\delta^{19}F^a$	"J _{P,F} b
(4)	227.9 Pm		$21.8 J_{m,a}$	-176.8	872 J _{F,Pm}
	8.5 Pa		$24.0 J_{m,b}$		$2.2 J_{\rm F,Pa}$
	5.6 Pb		$21.3 J_{a,b}$		$11.6 J_{F,Pb}$
(5)	663.4 P _x		12.6		7,10
	$-0.6 P_{a}$				
(6)	645.4 P _x		6.5		
	0.2 Pa				
(7)	184.3 Pm	446	$8.0 J_{m,a}$	-166.3c	907 J _{F,Pm}
	23.2 Pa	3879	39 J _{m,b}		$14J_{F,Pb}$
	25.3 Pb	3145	20 J _{a,b}		1.10
(8)	128.6 Pm	432	6 J _{m.a}		
	19.5 Pa	3772	40 J _{m,b}		
	23.0 Pb	3311	15 $J_{a,b}$		
(9)	585.0 P _x	98	3.6		
	16.2 P _a	3404			

^a In p.p.m.^b In Hz. ^c ²J_{Pt.F} 104 Hz.



metal at the P–C(C₅Me₅) single bond. The η^5 -Me₅C₅-metal complexes thus formed contain a phosphorus-metal σ -bond and a phospha-alkenyl fragment acting as either a 1e-³ or 3e-donor.² By analogy, generation of transition metal complexes containing iminophosphanyl³ or diphosphenyl ligands⁴ has been reported, starting from Me₅C₅-substituted iminophosphane and diphosphene precursors.

We explored whether this type of oxidative addition was limited to Me_5C_5 -substituted phosphorus p- π -systems, or whether it might be extended to other systems containing reactive P-X single bonds. Since P-halogen bonds in phosphaalkenes are readily cleaved by organic⁵ and organometallic nucleophiles^{6,7} with substitution at phosphorus and retention of the double bond, we investigated the reactivity of derivatives X-P=C(SiMe_3)₂ [X = F (1a),⁸ Cl (1b),⁹ I (1c)]⁸ with d¹⁰-metal-alkene complexes, (Bu₃P)₂Ni(cod), (2), and (Ph₃P)₂Pt(C₂H₄), (3).

Treatment of $[{}^{2}H_{6}]$ -benzene solutions of (2) or (3) at ambient temperature with 1 equiv. of (1a-c) immediately produced yellow to deep red coloured solutions. ¹H and ³¹P n.m.r. spectroscopic analyses confirmed the selective formation of complexes (4-9) (Figure 1), and ethylene or cycloocta-1,5-diene as the only spectroscopically detectable byproducts. The Pt complexes (7-9) were isolated as cream to orange coloured microcrystalline solids, following treatment of conc. solutions with hexane. After several hours, solutions of (4) displayed additional n.m.r. signals arising from decomposition, and no pure product could be isolated. Compounds (5) and (6) were found to be stable in solution for several hours in the presence of cyclo-octa-1,5-diene, enabling n.m.r. spectroscopic characterization. Attempts to remove the solvent and excess alkene *in vacuo*, however, led to decomposition and formation of $[NiX_2(PBu_3)_2]$ together with other, as yet unidentified products.

Structural formulation of compounds (4-9) was made on the basis of their ³¹P{¹H} n.m.r. data (Table 1). For (4), (7) and (8), ABM or ABMX type spectra were observed (A,B,M = ${}^{31}P$; X = ${}^{19}F$), which for the phospha-alkene resonance showed the considerable co-ordination shift1.10 and small magnitude of 1JPLP11 expected for bis(t-phosphine) (n2phospha-alkene)metal(0) complexes. The value of ${}^{1}J_{P,F}$ in (4, 7) shows a marked decrease with respect to free ligand $({}^{1}J_{P,F})$ 1087 Hz⁸) or the η^1 -complex [(Ph₃P)₂RhCl(η^1 -F-P=CR₂)] (1JP,F 1127 Hz12), suggesting a decrease in s-character for the P-F bond as a consequence of the η^2 -co-ordination. ³¹P{¹H} N.m.r. spectra of (5), (6) and (9) exhibit two multiplets of an AX2 system with the chemical shift of the X-part (8 584-663 p.p.m.) in the downfield region which is characteristic of transition metal substituted phosphorus p-n-systems.2-4.6.7 The observed equivalence of the phosphine ligands together with the comparatively small value of ${}^{2}J_{PAPX}$ suggests their formulation as *trans*-bis(t-phosphine) (σ -phospha-alkenyl)halogenometal(II) complexes. The phospha-alkenyl ligand in (9) exhibits an unusually low value (98 Hz) for ${}^{1}J_{Pt,P}$, which is however in accord with the small magnitude of ${}^{1}J_{M,P}$ in other systems containing phosphorus-metal single bonds.3.13 In the $^{13}C{^{1}H}$ n.m.r. spectra of (5), (6) and (9) the presence of a P=C double bond is indicated by a characteristic downfield multiplet [(5) δ 188.7 (${}^{1}J_{P,C}$ 105 Hz); (6) δ 183.9 (${}^{1}J_{P,C}$ 104 Hz); (9) δ 180.6 (${}^{1}J_{P,C}$ 106 Hz)].^{2,3,8} These findings are consistent with the presence of bent phospha-alkenyl ligands acting as 1e-donors,2.3 the observed equivalence of the Me₃Si resonances at ambient temperature† being a dynamic phenomenon caused by a rapid isomerization of the double bond.2

⁺ Selected ${}^{13}C{}^{1}H{}-n.m.r.$ data (20.0 MHz, ext. TMS): (5) (20 °C, [${}^{2}H_{6}$]-benzene) δ 3.1 (m, SiC₃); (-50 °C, [${}^{2}H_{6}$]-toluene) δ 3.4 (m, SiC₃) and 2.7 (d, SiC₃).

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Our findings indicate that the reactivity of phospha-alkenes (1a-c) towards transition metals is determined by a delicate balance of different factors.

The presence of a P-X bond of high bond energy (E) in (1a) $[E(P-F) 490 \text{ kJ mol}^{-114}]$ favours η^2 -co-ordination with retention of the P-X bond, whereas in the case of a weaker bond in (1c) $[E(P-I) 184 \text{ kJ mol}^{-114}]$ the oxidative addition is preferred. Compound (1b) represents an intermediate situation $[E(P-Cl) 319 \text{ kJ mol}^{-114}]$, and the product of the reaction depends on the nature of the transition metal fragment, since both complexation [with Ni(CO)₄¹³ and (3)] and insertion reactions [with (2)] are observed. Clearly, the strength of an individual P-X bond is of major importance for the reactivity of *P*-functionalized phospha-alkenes, and the behaviour of the phospha-alkenes involving other labile *P*-element single bonds towards low valent metal complexes is currently under study.

We gratefully acknowledge the loan of platinum metal salts from Messrs. Johnson Matthey, and financial support provided by the S.E.R.C. (M. F. M.) and the Deutsche Forschungsgemeinschaft and the Fonds der Chemischen Industrie (D. G.).

Received, 4th January 1989; Com. 9/00067D

J. CHEM. SOC., CHEM. COMMUN.

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