Synthesis, structure and kinetics of Group 6 metal carbonyl complexes containing a new ' P_2N ' mixed donor multidentate ligand †

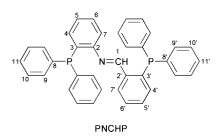
Eric W. Ainscough,* Andrew M. Brodie,* Paul D. Buckley, Anthony K. Burrell, Steven M. F. Kennedy and Joyce M. Waters

Chemistry - Institute of Fundamental Sciences, Massey University, Private Bag 11-222, Palmerston North, New Zealand. E-mail: A.Brodie@massey.ac.nz

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The 'P₂N' mixed donor ligand 2-(diphenylphosphino)-*N*-[2-(diphenylphosphino)benzylidene]aniline (PNCHP) reacted with [Cr(CO)₄(NBD)] and [M(CO)₄(C₅H₁₁N)₂] (M = Mo or W; NBD = norbornadiene; C₅H₁₁N = piperidine) to yield *cis*-[Cr(CO)₄(PNCHP- $\kappa^2 N, P$)] and *cis*-[M(CO)₄(PNCHP- $\kappa^2 P, P'$)] respectively. PNCHP behaves as a 'P,N' bidentate ligand when M is Cr and a 'P,P' bidentate ligand when M is Mo or W. Heating *cis*-[Cr(CO)₄(PNCHP- $\kappa^2 N, P$)] or *cis*-[M(CO)₄(PNCHP- $\kappa^2 P, P'$)] (M = Mo or W) in toluene affords the complexes *mer*-[M(CO)₃(PNCHP- $\kappa^3 P, N, P'$)] (M = Cr, Mo or W). Reaction of PNCHP with [Mo(CO)₃(CHT)] (CHT = 1,3,5-cycloheptatriene) in cold toluene yields *fac*-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)] which readily isomerises to *mer*-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)] in solution. PNCHP reacts with UV-irradiated tetrahydrofuran solutions of [M(CO)₆] (M = Cr or W) to yield phosphorus bound co-ordination isomers of [M(CO)₅(PNCHP- $\kappa^1 P$)]. The same reaction when carried out with one half equivalent of PNCHP yields the PNCHP bridged dinuclear species [{M(CO)₅}₂(PNCHP- $\kappa^2 P, P'$)]. The compounds have been characterised by spectroscopic means and the single crystal structures of *cis*-[Cr(CO)₄-(PNCHP- $\kappa^2 P, P'$)], *cis*-[W(CO)₄(PNCHP- $\kappa^2 P, P'$)], *mer*-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)] and [{W(CO)₅}₂(PNCHP- $\kappa^2 P, P'$)] has been studied to determine rate constants and thermodynamic activation parameters.

The mixed donor ligand, 2-(diphenylphosphino)-*N*-[2-(diphenylphosphino)benzylidene]aniline (PNCHP), can be visual-



ised as two triphenylphosphine moieties tethered together in the *ortho* positions by an imine linker. Since the imine bond has no plane of symmetry perpendicular to the C=N axis, the two phosphorus atoms are inequivalent which provides an excellent handle for characterising the metal complexes in solution by ³¹P NMR.

In this study we set out to establish the co-ordination preferences of the potentially tridentate 'P₂N' ligand in an octahedral environment. The Group 6 metal carbonyls were chosen because they offer an entry point into the realms of organometallic chemistry where such mixed donor ligand types have shown promise in catalysis.¹ One set of results obtained enabled the *fac*-to-*mer* isomerisation of the complex [Mo(CO)₃-(PNCHP)] to be characterised *via* a kinetic study. Recently, *fac*-*mer* isomerisation involving a tridentate 'P₂N' mixed donor ligand has been implicated in the C–C bond cleavage of terminal alkynes by water.² Kleverlaan *et al.* give a concise background to *fac*-*mer* isomerisation³ and in general the unassisted *fac-mer* isomerisation in six-co-ordinate octahedral complexes can occur in either of two ways. First, dissociation of a ligand can be followed by rearrangement of the resultant five-coordinate co-ordinatively unsaturated intermediate.⁴ Secondly, rearrangement may occur *via* a non-dissociative intramolecular mechanism.⁵ The majority of kinetic studies put forward to support either of the above isomerisation mechanisms have involved ligands of the mono- and/or bi-dentate variety.^{3,6,7} To our knowledge only one kinetic study has been reported involving a tridentate ligand.⁸

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Results and discussion

Synthesis and characterisation of PNCHP

The Schiff base condensation reaction of 2-diphenylphosphinobenzenaldehyde with 2-aminophenyldiphenylphosphine in refluxing benzene yields the ' P_2N ' ligand PNCHP. The choice of solvent is important since this reaction does not proceed in refluxing ethanol. PNCHP possesses two inequivalent phosphorus atoms and this is supported by the appearance of two signals at δ -14.2 and -14.8 [with no observable J(P-P) coupling] in the ³¹P-{¹H} NMR spectrum. The ¹³C-{¹H} and ¹H NMR spectral data, with assignments made using standard 2-D correlation and attached proton test (APT) techniques, are available as supplementary data. The proton spectrum displays an imine proton resonance at δ 8.92 which is split into a doublet by the nearest phosphorus atom with a coupling constant of 5.5 Hz [${}^{4}J(PH)$]. Four of the aromatic protons (H⁴, H⁵, H⁷, H^{5'} and $H^{7'}$) which are attached to the two central rings can clearly be identified at δ 7.03, 6.74, 7.96, 6.81 and 6.46 since they fall outside the main aromatic multiplet δ (7.16–7.36). In the ¹³C- ${^{1}H}$ spectrum the imine carbon (C¹) signal is the highest frequency resonance, appearing as a doublet at δ 157.9 [³*J*(PC) 25.0 Hz].

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[†] Electronic supplementary information (ESI) available: NMR data for the ligand, analytical and mass spectral data for the complexes. See http://www.rsc.org/suppdata/dt/b0/b002427i/

	³¹ P-{ ¹ H} NMR			¹ H NMR	
Complex	δ	² J(PMP)/Hz	¹ J(MP)/Hz	δ (N=CH)	⁴ J(PH)/Hz
1 mer-[Cr(CO) ₂ (PNCHP)]	79.6, 74.7	с		8.28 (s)	
$2 mer-[Mo(CO)_3(PNCHP)]$	59.0 (d), 53.7 (d)	93		8.56 (s)	
3 mer-[W(CO) ₃ (PNCHP)]	45.0 (d), 36.6 (d)	88	313, 315	8.50 (s)	
4 fac-[Mo(CO) ₃ (PNCHP)]	39.4 (d), 36.3 (d)	26		8.87 (s)	
5 cis-[Cr(CO) ₄ (PNCHP)]	53.2(s), -19.0(s)			c	
6 cis-[Mo(CO) ₄ (PNCHP)]	36.9 (d), 19.8 (d)	26		8.06 (s)	
7 cis-[W(CO) ₄ (PNCHP)]	22.2 (d), 3.0 (d)	27	238, 233	8.13 (s)	
$8[{Cr(CO)},{PNCHP}]$	55.2 (s), 54.2 (s)			8.65 (s)	
9 [{W(CO) ₅ },(PNCHP)]	19.2 (s), 18.1 (s)		244, 243	8.73 (s)	
10a [Cr(CO) ₅ (PNCHP)]	54.1 (s), -14.6 (s)			9.13 (d)	6.2
10b [Cr(CO), (PCHNP)]	54.5(s), -15.6(s)			8.44 (s)	
11a [W(CO) ₅ (PNCHP)]	18.6(s), -15.6(s)		244	9.13 (d)	6.1
11b [W(CO) ₅ (PCHNP)]	18.1 (s), -14.8 (s)		242	8.52 (s)	

^{*a*} Recorded at 109 MHz, chemical shifts are in ppm relative to 85% H₃PO₄, solvent CDCl₃. ^{*b*} Recorded at 270 MHz, chemical shifts are in ppm relative to TMS, solvent CDCl₃. ^{*c*} Not resolved.

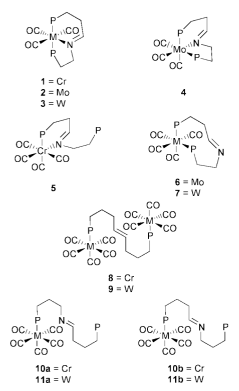


Fig. 1 The complexes showing the various co-ordination modes of PNCHP with the phenyl rings omitted for clarity.

Synthesis of PNCHP complexes

The complexes $[M(CO)_x(PNCHP)]$ (x = 3, 4 or 5, M = Cr, Mo or W) and $[{M(CO)_5}_2(PNCHP)] (M = Cr \text{ or } W)$ were prepared by conventional methods as described in the Experimental section. The air stable complexes all eventually convert in solution at room temperatures to $mer-[M(CO)_3(PNCHP-\kappa^3 P, N, P')]$, depicted by 1, 2 and 3 in Fig. 1, with the exception of the pentacarbonyl dimer species. This process takes hours for fac-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)] **4**, days for cis-[Mo(CO)₄-(PNCHP- $\kappa^2 P, P'$)] **6** and months for [Cr(CO)₅(PNCHP- $\kappa^1 P$)] 10a and 10b. The reverse reactions were not observed when dichloromethane solutions of the mer complexes were exposed to carbon monoxide gas at standard atmospheric pressure. The lability of similar systems, with respect to stability towards air and carbon monoxide substitution, appears to depend on the ease with which the nitrogen donor can be displaced. For example, in contrast to the tricarbonyl complexes in this paper, the 'P₂N' complex fac-[Mo(CO)₃{CH₃C(CH₂NHCH₃)(CH₂- $PPh_2_2 - \kappa^3 P, N, P'$] decomposes in air, even in the solid state,

and reacts spontaneously with carbon monoxide at standard atmospheric pressure to give the 'P,P' complex *cis*-[Mo-(CO)₄{CH₃C(CH₂NHCH₃)(CH₂PPh₂)₂- $\kappa^2 P$, P'}].⁹ The complexes are labelled and depicted in Fig. 1, NMR data are listed in Table 1, infrared and electronic spectral data in Table 2 and mass spectral (LSIMS) and microanalytical results, which are consistent with the formulations, have been deposited.

The key to obtaining the kinetic product fac-[Mo(CO)₃-(PNCHP- $\kappa^3 P, N, P'$)] **4** was its low solubility in toluene below 20 °C, enabling it to precipitate before its conversion into the thermodynamically favoured *mer* isomer. This facile conversion is most likely due to the ability of PNCHP easily to adopt the stable planar configuration when in the *mer* form. Unfortunately, the corresponding *fac* chromium and tungsten analogues could not be prepared, hence the *fac*-to-*mer* isomerisation kinetics was only carried out for the molybdenum complex. Kinetic and thermodynamic data for this isomerisation are shown in Table 3.

Single crystal X-ray diffraction studies

Crystal structure of *mer*-[Mo(CO)₃(PNCHP- $\kappa^{3}P, N, P'$)]. $2C_6H_5Me 2$. In complex 2 (Fig. 2 and Table 4), which has a *trans* arrangement of phosphorus atoms, the P(1)-Mo-P(2) angle is constrained from the ideal octahedral angle of 180° to be 159.29(3)° as a result of the strain imposed by the 5-membered chelate ring. This angle is typical of molybdenum carbonyl complexes containing five-membered chelate rings.^{11,12} As expected the carbonyl ligand which is trans to the N atom (Mo-C(3)) has a shorter Mo–C distance of 1.954(3) Å than the two carbonyl groups which are trans to one another (Mo-C(2), 2.019(3); Mo-C(4), 2.014(3) Å). The C(12)-N, N-C(1) and C(1)-C(42) bond lengths of 1.486(4), 1.258(3) and 1.445(4) Å respectively show that there is little, if any, electron delocalisation through the imine system. These lengths are similar to accepted values for C-N, C=N, and C-C bonds which are exo to an aromatic ring.¹³ (Similar results are found for 7, 5 and 9.) The Mo-N distance of 2.237(2) Å is similar to the lengths (2.294(3) and 2.278(4) Å) previously found in $cis-[Mo(CO)_4\{(CH_3)(C_6H_5)C=N(CH_2)\}_2-\kappa^2 N, N')]$.¹² The Mo–P distances of 2.425(2) and 2.431(2) Å do not differ significantly from one another despite the chemical inequivalence of the two P atoms. Similarly the six P-C bond lengths, which lie in the range 1.834(3)–1.840(3) Å, are in close agreement to each other.

Crystal structure of $cis-[W(CO)_4(PNCHP-\kappa^2 P, P')]$ **0.5CHCl₃ 7.** In complex 7 (Fig. 3 and Table 5), where the two P atoms occupy *cis* co-ordination sites, the largest deviation from the ideal co-ordination geometry is found, not unexpectedly, with the P(1)–W–P(2) angle of 102.27(4)° (*cf.* P(1)–Mo–P(2) of

	IR/cm ⁻¹		
Complex	v(C=O) ^a	v(C=N) ^b	UV-vis $\lambda/nm (\varepsilon_{max}/L mol^{-1}cm^{-1})^c$
1	1944w, 1847vs, 1828 (sh)	d	628 (1800)
2	1956w, 1857vs, 1836 (sh)	d	574 (1900) 551 (2200) ^e 586 (sh) (2500) ^f
3	1948w, 1846vs, 1824 (sh)	1571w	580
4	1932s, 1843s, 1825s ^b	1586w	536
5	2001m, 1911s, 1893s, 1857m	1582m	396 (2700)
6	2019m, 1919vs, 1900vs, 1865m	1634w	
7	2014m, 1911vs, 1892vs, 1863m	1637w	
8	2063m, 1983w, 1940s	1618w	
9	2071m, 1981w, 1938s	1613w	
10a,10b ^g	2063m, 1984w, 1940s	1614w	
11a,11b ^g	2071m, 1982w, 1939s	d	

^{*a*} In CHCl₃. ^{*b*} As Nujol mull. ^{*c*} M $\longrightarrow \pi^*_{imine}$ charge transfer band in CHCl₃. ^{*d*} Not assigned. ^{*e*} In acetone. ^{*f*} In toluene. ^{*g*} Isomer mixture.

Table 3 First-order rate constants and activation energies^{*a*} for geometrical isomerisation of *fac*-to-*mer*-[Mo(CO)₃(PNCHP- κP , N, P')]: $k_{obsd} = k_1 + k_{-1}, K_{eq} = k_1/k_{-1} = 7.3$

Solvent	<i>T</i> /°C	$10^{-5} k_{\rm obsd}/{\rm s}^{-1}$	$10^{-5} k_1/s^{-1}$	$10^{-5} k_{-1} / s^{-1}$
Acetone	19.5	1.39	1.22	0.170
	29.7	4.75	4.18	0.574
	40.0	19.6	17.2	2.08
	49.5	55.6	48.9	6.71
Acetone/Et ₃ N ^b	19.8	1.37	с	
CH ₂ Cl ₂	19.4	3.46	с	
MeČN	40.0	11.6	с	

^{*a*} $\Delta H_1^{\ddagger} = 95 \pm 3 \text{ kJ mol}^{-1}, \Delta H_{-1}^{\ddagger} = 94 \pm 2 \text{ kJ mol}^{-1}, \Delta S_1^{\ddagger} = -14.1 \pm 0.7$ J K⁻¹ mol⁻¹, $\Delta S_{-1}^{\ddagger} = -36 \pm 2 \text{ J K}^{-1} \text{ mol}^{-1}$. ^{*b*} 2 mol equivalents of triethylamine. ^{*c*} K_{eq} not recorded.

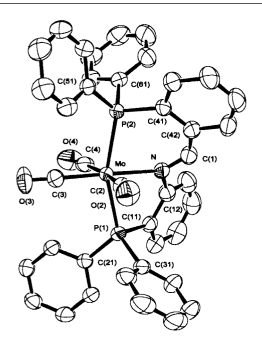


Fig. 2 An ORTEP¹⁰ diagram for the complex *mer*-[Mo(CO)₃-(PNCHP- $\kappa^3 P, N, P'$)] **2** showing the numbering system used. Thermal ellipsoids are at the 50% probability level. Hydrogen atoms have been omitted for reasons of clarity.

104.62(7)° found in *cis*-[Mo(CO)₄(PPh₃)₂]¹²). There is a small difference in the two W–P bond lengths (W–P(1), 2.5749(12); W–P(2), 2.5576(14) Å) but both are very similar to those observed in other tungsten–carbonyl containing derivatives where the triphenylphosphine is *trans* to a carbonyl group.¹⁴ No significantly different distances in the W–C bond lengths are noted for the two carbonyl ligands in *trans* co-ordination sites

Table 4 Selected bond lengths [Å] and angles [°] for *mer*-[Mo(CO)₃-(PNCHP- $\kappa^3 P, N, P'$)]·2C₆H₅Me

Mo–P(1)	2.431(2)	P(2)–C(41)	1.840(3)
Mo-P(2)	2.425(2)	P(2)–C(51)	1.838(3)
Mo–N	2.237(2)	P(2)–C(61)	1.836(3)
Mo-C(2)	2.019(3)	O(2)–C(2)	1.148(3)
Mo-C(3)	1.954(3)	O(3)–C(3)	1.162(3)
Mo-C(4)	2.014(3)	O(4)–C(4)	1.152(3)
P(1)-C(11)	1.834(3)	N–C(1)	1.258(3)
P(1)–C(21)	1.836(3)	N-C(12)	1.486(4)
P(1)–C(31)	1.840(3)	C(1)–C(42)	1.445(4)
$D(2) = M_{-1} = D(1)$	150 20(2)	$C(51)$ $D(2)$ M_{-}	110 74(0)
P(2)-Mo-P(1)	159.29(3)	C(51)-P(2)-Mo	119.74(9)
C(2)-Mo-P(1)	91.92(8)	C(61)-P(2)-Mo	120.49(9)
C(3)–Mo– $P(1)$	101.33(8)	O(2)–C(2)–Mo	175.4(2)
C(4)– Mo – $P(1)$	89.11(8)	O(3)–C(3)–Mo	177.8(2)
C(2)–Mo–P(2)	86.79(8)	O(4)–C(4)–Mo	173.3(3)
C(3)–Mo–P(2)	99.27(8)	C(1)–N–Mo	129.8(2)
C(4)–Mo–P(2)	95.36(8)	C(12)–N–Mo	114.3(2)
N-Mo-P(1)	76.71(6)	C(11)-P(1)-C(21)	105.79(11)
N-Mo-P(2)	82.82(6)	C(11)-P(1)-C(31)	103.60(12)
C(2)-Mo-C(3)	87.66(11)	C(21)-P(1)-C(31)	100.76(12)
C(2)-Mo-C(4)	171.07(10)	C(41)-P(2)-C(51)	102.90(11)
C(3)-Mo-C(4)	83.45(12)	C(41)–P(2)–C(61)	102.19(11)
C(2)–Mo–N	95.22(10)	C(51)-P(2)-C(61)	102.16(12)
C(3)–Mo–N	176.55(10)	C(1) - N - C(12)	115.5(2)
C(4)–Mo–N	93.64(11)	N-C(1)-C(42)	127.5(3)
C(11)–P(1)–Mo	100.74(9)	C(16)-C(11)-P(1)	125.6(2)
C(21)–P(1)–Mo	123.05(8)	C(12)-C(11)-P(1)	114.8(2)
C(31)–P(1)–Mo	120.59(9)	C(13)-C(12)-N	121.4(3)
C(41)–P(2)–Mo	106.77(8)	C(11)–C(12)–N	118.3(2)

to the P atoms. However these two W–C bonds (W–C(4) and W–C(5)) at 1.977(6) and 1.970(6) Å are both shorter than the two related *trans* distances of 2.035(6) and 2.016(6) Å for W–C(2) and W–C(3) respectively, a result not unexpected when the π acidity of the CO group is taken into account. The long distance between W and N (3.530(4) Å) indicates the absence of a bond between these two atoms. Support for the distinction made between the C and N atoms of the imine bridge on the basis of their thermal parameters comes from steric arguments. If these two atoms were interchanged then the H associated with the C would approach too closely to one of the carbonyl ligands (see Fig. 3).

Crystal structure of *cis*-[**Cr**(**CO**)₄(**PNCHP**- $\kappa^2 P$, *N*)]-0.5**CHCl**₃ **5.** In complex **5** (Fig. 4 and Table 6) there are two independent molecules in the asymmetric unit showing no significant differences from one another. One of the independent molecules (molecule A) has arbitrarily been selected for the purpose of this discussion. Complex **5** shows a much smaller deviation from the ideal octahedral co-ordination geometry than do **2** and **7** discussed previously. The N and one of the P atoms

Table 5 Selected bond lengths [Å] and angles [°] for cis-[W(CO)₄- (PNCHP- $\kappa^2 P, P'$)]-0.5CHCl₃

W–P(1)	2.5749(12)	P(2)–C(51)	1.833(5)
W-P(2)	2.5576(14)	P(2)–C(61)	1.845(5)
W-C(2)	2.035(6)	O(2)–C(2)	1.139(5)
W-C(3)	2.016(6)	O(3)–C(3)	1.150(5)
W-C(4)	1.977(6)	O(4) - C(4)	1.160(5)
W-C(5)	1.970(6)	O(5)–C(5)	1.156(6)
P(1)-C(11)	1.835(5)	N–C(1)	1.257(6)
P(1)-C(21)	1.841(5)	N-C(12)	1.405(6)
P(1)-C(31)	1.825(5)	C(1)–C(42)	1.451(6)
P(2)–C(41)	1.850(5)		
		a/a	
P(1)-W-P(2)	102.27(4)	C(5)–W–N	136.8(2)
P(1)-W-C(2)	82.84(13)	C(11) - P(1) - W	123.5(2)
P(1)-W-C(3)	94.93(13)	C(21)-P(1)-W	109.9(2)
P(1)-W-C(4)	168.3(2)	C(31)-P(1)-W	115.7(2)
P(1)-W-C(5)	86.7(2)	C(11)-P(1)-C(21)	102.5(2)
P(2)-W-C(2)	87.21(14)	C(11)-P(1)-C(31)	100.9(2)
P(2)-W-C(3)	90.93(14)	C(21)-P(1)-C(31)	101.5(2)
P(2)-W-C(4)	86.5(2)	C(41)-P(2)-C(51)	107.1(2)
P(2)-W-C(5)	171.0(2)	C(41)-P(2)-C(61)	101.8(2)
P(1)-W-N	55.25(7)	C(51)-P(2)-C(61)	96.5(2)
P(2)-W-N	51.08(7)	C(41)–P(2)–W	110.3(2)
C(2)-W-C(3)	176.7(2)	C(51)–P(2)–W	122.5(2)
C(2)-W-C(4)	90.0(2)	C(61)–P(2)–W	116.0(2)
C(2)-W-C(5)	93.3(2)	O(2)–C(2)–W	177.8(4)
C(3) - W - C(4)	92.6(2)	O(3)–C(3)–W	178.3(4)
C(3) - W - C(5)	89.0(2)	O(4)–C(4)–W	175.5(5)
C(4) - W - C(5)	84.5(2)	O(5)–C(5)–W	174.9(5)
C(2)-W-N	65.1(2)	C(1)-N-W	122.0(3)
C(3)-W-N	111.7(2)	C(12)–N–W	110.2(3)
C(4)-W-N	129.3(2)	C(1) - N - C(12)	125.6(4)

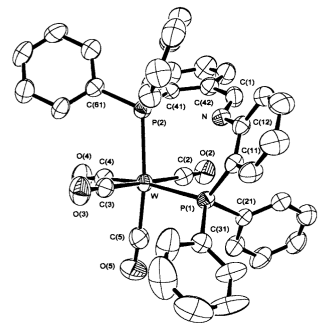


Fig. 3 An ORTEP diagram for the complex cis-[W(CO)₄(PNCHP- $\kappa^2 P, P'$)] 7. Details as in Fig. 2.

occupy *cis* co-ordination sites forming a six-membered ring with a P(2A)–Cr(1)–N(1A) angle of $82.06(13)^\circ$.‡ As expected the carbonyl *trans* to the N atom (Cr(1)–C(4A)) has the shorter Cr–C distance of 1.794(7) Å than the carbonyl group *trans* to the P atom (Cr(1)–C(5A), 1.854(7) Å). The bond lengths and angles

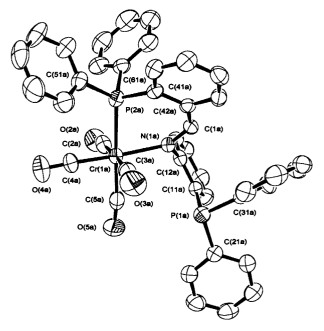


Fig. 4 An ORTEP diagram for the complex cis-[Cr(CO)₄(PNCHP- $\kappa^2 P$,N)] **5**. Details as in Fig. 2.

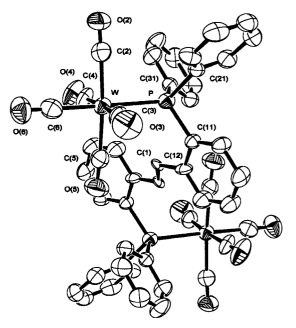


Fig. 5 An ORTEP diagram for the complex $[{W(CO)_5}_2(PNCHP-\kappa^2 P, P')]$ 9. Details as in Fig. 2.

around the Cr are virtually identical to those of the related complex *cis*-[{Cr(CO)₄}₂(P₂N₂- κ^4 P,N,N',P')], where P₂N₂ is N,N'-bis[o-(diphenylphosphino)benzylidene]ethylenediamine.¹⁶

Crystal structure of [{W(CO)₅}₂(PNCHP-\kappa^2 P, P')]·2C₆H₅Me 9. The dinuclear complex 9 (Fig. 5 and Table 7), where two W atoms are linked by the PNCHP bridge, also shows a much smaller deviation from the ideal octahedral co-ordination geometry than do 2 and 7. The W–P bond length of 2.554(2) Å is in accord with its counterparts in complex 7 and [W(CO)₅-(PCHO-\kappa^1 P)] (PCHO = 2-diphenylphosphinobenzaldehyde) (2.561(1) Å¹⁷) and it is in a *trans* **position to the shortest of the 5 W–C bonds. The C and N of the imine bridge were not distinguished due to the presence of the centre of symmetry at the midpoint of the C–N bond.**

Electronic and infrared spectroscopy

The colour of the complex was a reliable visual aid for

[‡] Previous studies (ref. 15) have indicated that PNCHP more readily binds a sulfur atom to the phosphorus atom three bonds removed from the imine nitrogen, implying that this maybe the more basic site. Incorporation of this P atom into the chelate ring in *cis*-[Cr(CO)₄(PNCHP)] would result in a five-membered ring. The single crystal structure shows that a six-membered ring is formed with the P atom four bonds removed from the co-ordinated imine N.

Cr(1)–P(2A)	2.3570(18)	P(2A)-C(51A)	1.825(6)
Cr(1)-N(1A)	2.141(5)	P(2A)-C(61A)	1.818(6)
Cr(1)-C(2A)	1.895(8)	O(2A) - C(2A)	1.141(7)
Cr(1)-C(3A)	1.879(8)	O(3A) - C(3A)	1.157(7)
Cr(1)-C(4A)	1.794(7)	O(4A) - C(4A)	1.162(6)
Cr(1)-C(5A)	1.854(7)	O(5A) - C(5A)	1.150(7)
P(1A) - C(11A)	1.830(6)	N(1A) - C(1A)	1.279(6)
P(1A) - C(21A)	1.828(6)	N(1A)-C(12A)	1.449(7)
P(1A) - C(31A)	1.825(6)	C(1A) - C(42A)	1.464(8)
P(2A) - C(41A)	1.817(6)		
N(1A)-Cr(1)-P(2A)	82.06(13)	C(21A) - P(1A) - C(31A)	105.6(3)
P(2A)-Cr(1)-C(2A)	95.0(2)	C(41A) - P(2A) - C(51A)	102.8(3)
P(2A)-Cr(1)-C(3A)	89.1(2)	C(41A)–P(2A)–C(61A)	102.1(3)
P(2A)-Cr(1)-C(4A)	94.3(2)	C(51A)–P(2A)–C(61A)	102.7(3)
P(2A)-Cr(1)-C(5A)	177.2(2)	C(41A)-P(2A)-Cr(1)	108.89(19)
C(2A)-Cr(1)-C(3A)	175.5(3)	C(51A)-P(2A)-Cr(1)	120.1(2)
C(2A)-Cr(1)-C(4A)	89.2(3)	C(61A) - P(2A) - Cr(1)	117.8(2)
C(2A)-Cr(1)-C(5A)	87.5(3)	O(2A)-C(2A)-Cr(1)	174.8(6)
C(3A)-Cr(1)-C(4A)	88.8(3)	O(3A)-C(3A)-Cr(1)	175.6(6)
C(3A)-Cr(1)-C(5A)	88.4(3)	O(4A)-C(4A)-Cr(1)	177.3(6)
C(4A)-Cr(1)-C(5A)	86.9(3)	O(5A) - C(5A) - Cr(1)	173.1(6)
C(2A)-Cr(1)-N(1A)	91.4(2)	C(1A)-N(1A)-Cr(1)	130.5(4)
C(3A)-Cr(1)-N(1A)	90.9(2)	C(12A)-N(1A)-Cr(1)	115.9(3)
C(4A)-Cr(1)-N(1A)	176.4(2)	C(1A)-N(1A)-C(12A)	113.6(5)
C(5A)-Cr(1)-N(1A)	96.7(2)	N(1A)-C(1A)-C(42A)	129.4(5)
C(11A) - P(1A) - C(21A)	100.3(3)	C(11A)-C(12A)-N(1A)	121.4(5)
C(11A)–P(1A)–C(31A)	100.2(3)	C(13A)-C(12A)-N(1A)	117.7(5)

Table 7 Selected bond lengths [Å] and angles [°] for $[{W(CO)_5}_2 - (PNCHP-\kappa^2P, P')] \cdot 2C_6H_5Me$

W–P	2.554(2)		P–C(31)	1.836(8)
W-C(2)	2.025(10)		O(2) - C(2)	1.152(9)
W-C(3)	1.998(10)		O(3) - C(3)	1.162(9)
W-C(4)	1.986(10)		O(4) - C(4)	1.162(9)
W-C(5)	2.036(10)		O(5) - C(5)	1.135(9)
W-C(6)	1.970(11)		O(6)–C(6)	1.167(10)
P-C(11)	1.838(7)		C(1)–C(1)#1	1.281(12)
P-C(21)	1.840(8)		C(1)–C(12)	1.427(10)
C(2)–W–P	89.7(2)		O(5)–C(5)–W	176.5(8)
C(3)-W-P	91.1(2)		O(6) - C(6) - W	179.0(8)
C(4)-W-P	92.1(2)		C(11) - P - W	115.5(2)
C(5)-W-P	93.4(2)		C(21)–P–W	116.3(3)
C(6)-W-P	178.2(2)		C(31) - P - W	115.0(3)
C(2) - W - C(3)	93.1(3)		C(11) - P - C(21)	101.1(3)
C(2) - W - C(4)	89.6(3)		C(11) - P - C(31)	106.5(3)
C(2) - W - C(5)	175.6(4)		C(21) - P - C(31)	100.6(3)
C(2) - W - C(6)	88.7(3)		C(12)–C(11)–P	120.9(6)
C(3) - W - C(4)	175.9(4)		C(16)–C(11)–P	119.9(6)
C(3) - W - C(5)	89.9(4)		C(22)–C(21)–P	122.0(6)
C(3) - W - C(6)	88.2(4)		C(26)–C(21)–P	118.6(6)
C(4) - W - C(5)	87.3(4)		C(32)–C(31)–P	124.1(6)
C(4) - W - C(6)	88.7(3)		C(36)–C(31)–P	117.5(6)
C(5) - W - C(6)	88.2(3)		C(1)#1-C(1)-C(12)	118.3(8)
O(2) - C(2) - W	177.1(8)		C(13)-C(12)-C(1)	121.7(7)
O(3) - C(3) - W	176.5(8)		C(11)-C(12)-C(1)	120.0(6)
O(4)–C(4)–W	177.1(7)			
Symmetry tran	nsformation	used	to generate equival	ent atoms:
#1 - x, -v, -z.			C 1	

determining the environment of the imine nitrogen. In species where the nitrogen atom was co-ordinated an electronic absorption in the range 396 to 628 nm with ε_{max} values 1800 to 2700 L mol⁻¹ cm⁻¹ was observed as listed in Table 2. Metal-to-ligand charge transfer bands (MLCT) are expected when the ligand has low lying empty orbitals. Unsaturated ligands with empty π^* orbitals are prime candidates.¹⁸ Hence the absorption responsible for giving the complexes their characteristic 'rich' and intense colours as found for the blood red *cis*-[Cr(CO)₄-(PNCHP- $\kappa^2 P$,N)] **5**, the crimson *fac*-[Mo(CO)₃(PNCHP- $\kappa^3 P$,N,P')] **1**, **2**, **3** complexes is most likely a M $\longrightarrow \pi^*_{imine}$ charge transfer. In further support of the assignment, the absorption exhibited ε_{max} values and solvatochromism (see Table 2) typical of a MLCT band.¹⁸ Conversely, when the nitrogen atom was not co-ordinated, light reddish orange and yellow colours were observed, as found for the tetracarbonyl complexes *cis*-[M(CO)₄(PNCHP- $\kappa^2 P, P'$)] (M = Mo 6 or W 7) and for all the pentacarbonyl complexes.

The contrasting colours of the light reddish orange molybdenum and tungsten tetracarbonyl complexes in comparison to the intense dark red of the chromium tetracarbonyl complex can now be explained in terms of the absence of the $M \longrightarrow \pi^*_{imine}$ charge transfer band in the former cases. For molybdenum or tungsten complexes, both phosphorus atoms of PNCHP are co-ordinated with the nitrogen atom non-coordinated, as confirmed by ³¹P NMR spectroscopy and X-ray crystallography. In contrast, the chromium tetracarbonyl complex has only one phosphorus atom bound, with the imino nitrogen atom binding in preference to the second phosphorus atom. This phenomenon can be explained by evoking the Hard Soft Acid Base (HSAB) principle. This co-ordination set is also observed in the 'P₂N₂' ligand N,N'-bis[o-(diphenylphosphino)benzylidene]ethylenediamine which binds to the 'Cr(CO)₄' moiety via one of its two phosphorus donor atoms and one imine nitrogen atom.16

The wavenumbers and number of IR bands in the carbonyl stretching region (Table 2) are typical for each type of carbonyl species.^{9,16,19-21} When the imine nitrogen was non-co-ordinated, as found for the pentacarbonyl and molybdenum and tungsten tetracarbonyl complexes, the ν (C=N) band was at similar or slightly higher wavenumbers when compared with the free PNCHP ligand (1620 cm⁻¹). However, as expected, when the imine nitrogen was co-ordinated, the ν (C=N) band was lowered by 35 to 50 cm⁻¹ in energy as shown in Table 2.

NMR spectroscopy

Proton NMR spectra (Table 1) showed the expected integration and peak multiplicities. In all complexes where the phosphorus nearest to the imine carbon of PNCHP was involved in coordination, the imine proton signal shifts to lower frequencies relative to 'free' PNCHP [δ 8.92, ⁴*J*(PH) 5.5 Hz] and the P–H coupling is no longer observed. In contrast, for the complexes where this phosphorus was non-co-ordinated a higher frequency shift was observed, as seen in the mixtures of the $[M(CO)_5(PNCHP-\kappa^1 P)]$ (M = Cr 10a, 10b or W 11a, 11b) complexes.

When one equivalent of PNCHP is treated with the ' $M(CO)_{5}$ ' (M = Cr or W) moiety both phosphorus co-ordination isomers are obtained. However, the ratio of the two isomers is not a statistical 1:1 mixture but 2:1 in favour of 10a and 11a. This was concluded by considering the NMR signal multiplicity of the imine proton. In the "free" PNCHP ligand it is split by the phosphorus nucleus nearest to the imine carbon $({}^{4}J(PH) = 5.5)$ Hz). However, when the same phosphorus is bound to sulfur, as in PNCHP=S, the imine proton is a singlet suggesting that coordination of the phosphorus four bonds removed from the nitrogen means that P-H coupling is no longer observed. When sulfur is bound to the other phosphorus atom, as in S=PNCHP, a doublet imine proton is observed.¹⁵ The complexes $[Cr(CO)_{5}(PNCHP-\kappa^{1}P)]$ 10a and $[W(CO)_{5}(PNCHP-\kappa^{1}P)]$ 11a both display doublet imine proton signals while [Cr(CO)5-(PNCHP- $\kappa^{1}P$)] 10b and [W(CO)₅(PNCHP- $\kappa^{1}P$)] 11b have singlet signals (Table 1). The comparable ligand N-(2-diphenylphosphinobenzylidene)-2-(2-pyridyl)ethylamine²² shows no imine proton-phosphorus coupling in the "free" ligand, further supporting the imine proton coupling assignment in PNCHP. Hence we have assigned the major isomer of the above reaction to $[M(CO)_5(PNCHP-\kappa^1 P)]$ 10a, 11a and the minor isomer to $[M(CO)_{5}(PNCHP-\kappa^{1}P)]$ **10b**, **11b** based on the chemical shifts and coupling constants listed in Table 1 and supported by the reaction of sulfur with PNCHP. These assignments have allowed us to assign the weakest resonances in the ³¹P NMR spectra to the minor isomer 10b and 11b. ${}^{1}J(WP)$ coupling values of 244 and 242 Hz respectively are typical.23

The ³¹P NMR data, shown in Table 1, of the molybdenum and tungsten tetracarbonyl species showed all phosphorus nuclei resonances at higher frequency than the 85% H₃PO₄ reference, with an AB splitting pattern and coupling constants of 26 and 27 Hz respectively. This is indicative of PNCHP coordinated through both phosphorus atoms in *cisoid* fashion.²⁴ In contrast, the chromium tetracarbonyl species showed only one phosphorus nuclei resonance at higher frequency than H₃PO₄, the other resonance being at a lower frequency with no phosphorus-phosphorus coupling. This is consistent with one co-ordinated and one non-co-ordinated P^{III} respectively. A small amount (<5%) of another chromium species with δ 55.4 and δ 38.4, ²J(PP) 35 Hz, is observed in the ³¹P NMR spectrum of the crude product. This is most likely the chromium analogue of the 'P,P'-bound molybdenum and tungsten complexes. The expected tungsten satellites are observed in the ³¹P NMR spectrum of $cis-[W(CO)_4(PNCHP-\kappa^2 P, P')]$ 7, the $^{1}J(WP)$ values of 238 and 233 Hz being typical.^{23,25}

The most significant differences found in the ³¹P NMR spectra of the tricarbonyl complexes (compared to the tetracarbonyl species) are the much larger ²*J*(PP) coupling constants of *ca.* 90 Hz. These are indicative of a *transoid* arrangement of the phosphorus atoms around the metal,²⁴ supporting a meridional arrangement of the PNCHP ligand. In general the tricarbonyl species exhibit shifts of some 20 ppm higher in frequency than their tetracarbonyl counterparts which is possibly due to the increased ring strain, particularly for the five membered ring component,²⁵ when PNCHP behaves as a tridentate ligand. The isomer *fac*-[Mo(CO)₃(PNCHP-κ³*P*,*N*,*P'*)] **4** displays a ²*J*(PP) coupling constant of 26 Hz typical of *cisoid* phosphorus atoms, hence supporting the *fac* isomer assignment.²⁶

Kinetic and equilibrium studies

Preliminary studies of the geometrical isomerisation of fac-[Mo(CO)₃(PNCHP- $\kappa^{3}P, N, P'$)] **4** to mer-[Mo(CO)₃(PNCHP- $\kappa^{3}P, N, P'$)] **2** were made in acetone, acetonitrile, chloroform and dichloromethane by following the increase in the absorbance at 700 nm as the purple **2** product formed. The *fac*-to-*mer* isomerisation proceeds with considerable complex decom-

position in chloroform and to a lesser, but still significant, extent in dichloromethane while solubility problems were encountered with acetonitrile as solvent at the lower temperatures. In acetone the solubility was sufficient and the isomerisation took place without decomposition over the temperature range 19.5–49.5 °C. Hence, comprehensive kinetic and thermodynamic parameters could only be obtained for the isomerisation in this solvent. As depicted in eqn. (1), k_1 and k_{-1} denote

$$fac \xrightarrow[k_{1}]{k_{1}} mer$$
 (1)

the forward and reverse rate constants, respectively. There was no spectroscopic evidence for detectable concentrations of any intermediates. In each kinetic run, a plot of $\ln(A - A_{\infty})$ versus time gave a straight line for at least three half-lives (A and A_{∞} denote the absorbance at 700 nm at time t and at infinite time respectively, where the *fac-mer* mixture was at thermodynamic equilibrium). The slope gave the first-order rate constant k_{obsd} which was determined at four different temperatures (19.5, 29.7, 40.0 and 49.5 °C). The rate constants for isomerisation were calculated from the equilibrium constant K_{eq} , obtained from the integrated ¹H NMR signals of the imine proton in equilibrated solutions (eqn. 2), and the observed rate constant

$$K_{eq} = k_1 / k_{-1} = \int \delta C H = N_{mer} / \int \delta C H = N_{fac}$$
(2)

(eqn. 3). The K_{eq} value was found to be 7.3 and independent of

$$k_{\rm obsd} = k_1 + k_{-1} \tag{3}$$

temperature in acetone. The k_{obsd} values along with the forward (k_1) and reverse (k_{-1}) rate constants are collected in Table 3. The activation enthalpies for the forward (ΔH_1^{\ddagger}) and reverse $(\Delta H_{-1}^{\ddagger})$ reactions were 95 ± 3 and 94 ± 2 kJ mol⁻¹, respectively. The activation entropies for the forward (ΔS_1^{\ddagger}) and reverse $(\Delta S_{-1}^{\ddagger})$ reactions were -14.1 ± 0.7 and -36 ± 2 J K⁻¹ mol⁻¹, respectively.

Driving forces for *fac-mer* isomerisation can be attributed to electronic advantages and/or steric pressure exerted by the ligands around the metal.^{4,27} In this case the PNCHP ligand prefers to be coplanar, as it is in the *mer* isomer. This is probably due to the structural rigidity and inherent inflexibility of the PNCHP ligand imposed on the ligand by the imine double bond with the phosphino moieties locked *trans* to the double bond.

In general, a large positive activation enthalpy (>100 kJ mol⁻¹), that is of bond breaking magnitude, coupled with a positive activation entropy is supportive of a dissociative mechanism whilst a small activation enthalpy, less than bond breaking magnitude, coupled with a large negative entropy $(\approx -200 \text{ J K}^{-1} \text{ mol}^{-1})$ is supportive of a twist mechanism.²⁸ For our system, ΔH_1^{\ddagger} and ΔH_{-1}^{\ddagger} values of 95 and 94 kJ mol⁻¹ coupled respectively with ΔS_1^{\ddagger} and ΔS_{-1}^{\ddagger} values of -14.1 and -36 J K⁻¹ mol⁻¹ are inconclusive with respect to which mechanism the isomerisation follows. Note that if solvent restriction occurs ΔS^{\ddagger} could be negative for a dissociative mechanism. Rate constants did decrease slightly with increasing co-ordinating powers of the solvent. For example, for the co-ordinating solvent acetonitrile, the rate constant halves compared with acetone whilst moving to the poor coordinating solvent dichloromethane the rate constant doubles. One might equate this observation with a dissociative mechanism, however the addition of Et₃N to acetone did not affect the rate. In addition, whilst following the isomerisation by ³¹P NMR in the presence of a tenfold excess of triphenyl phosphite no intermediates on the NMR timescale were detected. Triphenyl phosphite may have been expected to bind to the metal if a dissociative mechanism was in effect. Without thermodynamic activation parameters for the isomerisation in acetonitrile and dichloromethane, a more substantial conclusion with respect to the solvent effect cannot be drawn.

Conclusion

Related compounds of the type $[{M(CO)_x}_{\nu}]_{\nu}(PNCHP)]$ where M is Cr, Mo or W and x = 3, 4 or 5, and y = 1 or 2 have been synthesized. The 'P2N' mixed donor multidentate ligand PNCHP adopts $\kappa^2 P, N$ cis co-ordination in cis-[Cr(CO)₄-(PNCHP- $\kappa^2 P, N$)] but a $\kappa^2 P, P'$ cis co-ordination for the complexes of Mo and W. For the compounds [M(CO)₃(PNCHP- $\kappa^{3}P, N, P'$], PNCHP co-ordinates in a *mer* fashion. The *fac* isomer is also isolatable for Mo but readily converts into the *mer* isomer. The first-order forward rate constants (k_1) for the fac-to-mer isomerism in acetone are 1.22×10^{-5} , 4.18×10^{-5} , 1.72×10^{-4} and 4.89×10^{-4} s⁻¹ at 19.5, 29.7, 40.0 and 49.5 °C respectively, with thermodynamic activation parameters ΔH_1^{\ddagger} and ΔS_1^{\ddagger} of 95 kJ mol⁻¹ and -14.1 J K⁻¹ mol⁻¹ respectively. For $[M(CO)_5(PNCHP-\kappa^1 P)]$ (M = Cr or W) the favoured binding mode is via the phosphorus atom nearest to the imine nitrogen atom. PNCHP will also bridge (via the two phosphorus atoms) two pentacarbonyl moieties.

Experimental

Materials and instrumentation

NMR measurements were performed on a JEOL GX270W spectrometer. IR spectra were recorded on a BIO-RAD FTS-40 instrument, UV-vis spectra using a Shimadzu UV-310PC UV-VIS-NIR scanning spectrometer. Mass spectra were obtained using a Varian VG70-250S double-focussing magnetic sector spectrometer by the method of liquid ion secondary mass spectroscopy (LSIMS) using *m*-nitrobenzyl alcohol as the matrix. Isotope abundance calculations were performed to identify the parent ion. Microanalyses were performed by the Campbell Microanalytical Laboratory, University of Otago. 2-Diphenyl-(2-PCHO),²⁹ phosphinobenzaldehyde 2-aminophenyldiphenylphosphine (2-PNH₂),³⁰ [Cr(CO)₄(NBD)],³¹ [M(CO)₄- $(pip)_2$] (pip = piperidine) $(M = Mo \text{ or } W)^{32}$ and $[Mo(CO)_3-(CHT)]^{33}$ (CHT = cycloheptatriene) were prepared according to literature. Solvents were dried by standard procedures. All reactions were carried out under an inert atmosphere.

Syntheses

2-(Diphenylphosphino)-*N*-**[(2-diphenylphosphino)benzylidene]**aniline (PNCHP). To a three-necked flask fitted with a Dean-Stark trap and condenser was added 2-PCHO (1.160 g, 4 mmol), 2-PNH₂ (1.108 g, 4 mmol), toluene-*p*-sulfonic acid (20 mg, 0.10 mmol) and benzene (120 cm³). The solution was refluxed for 7 h and then the solvent removed by rotary evaporation to produce a yellow oil. This was dissolved in the minimum amount of dichloromethane, then methanol was added until crystallisation began. After standing for half an hour PNCHP was filtered off as pale yellow crystals, washed with methanol and dried *in vacuo.* A second crop was obtained by concentration of the mother liquor by rotary evaporation and crystallisation from dichloromethanemethanol. Yield: 1.825 g (80%). mp 176–178°C. IR: 1620 cm⁻¹ [ν (C=N)].

mer-[Cr(CO)₃(PNCHP- $\kappa^3 P, N, P'$)]·2C₆H₅Me 1. A mixture of [Cr(CO)₄(NBD)] (0.143 g, 0.558 mmol) and PNCHP (0.306 g, 0.557 mmol) in 25 mL of toluene was heated to reflux for 1.5 h giving a dark green solution. The cooled solution was filtered through Kieselguhr and the filtrate allowed to stand overnight. The resulting solid was isolated, washed with n-pentane and dried *in vacuo*, yielding 0.130 g (34%) of *mer*-[Cr(CO)₃-(PNCHP- $\kappa^3 P, N, P'$)]·2C₆H₅Me as dark green crystals, mp 207–210 °C.

mer-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)]·CH₂Cl₂ 2. A mixture of [Mo(CO)₃(CHT)] (0.140 g, 0.529 mmol) and PNCHP (0.291 g, 0.530 mmol) in 25 mL of toluene was heated to reflux for 25 min giving a dark blue solution. The cooled solution was filtered through Kieselguhr and the filtrate allowed to stand overnight. The resulting solid was isolated, washed with n-pentane and dried *in vacuo*, to give *mer*-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)]·CH₂Cl₂ as dark blue crystals, mp 114–115 °C.

fac-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)] **4.** An ice-cold solution of [Mo(CO)₃(CHT)] (0.099 g, 0.365 mmol) dissolved in toluene was added to an ice-cold solution of PNCHP (0.200 g, 0.364 mmol) dissolved in toluene. The mixture was stirred for *ca*. 1 h, keeping the temperature below 0 °C, followed by 23 h at room temperature. The resulting precipitate was isolated, washed with n-pentane, and dried *in vacuo*, yielding 0.215 g (73%) of *fac*-[Mo(CO)₃(PNCHP- $\kappa^3 P, N, P'$)] as a crimson powder, mp 154–155 °C.

mer-[W(CO)₃(PNCHP- $\kappa^3 P$, *N*, *P'*)]·CH₂Cl₂ 3. A mixture of *cis*-[W(CO)₄(pip)₂] (0.206 g, 0.440 mmol) and PNCHP (0.243 g, 0.440 mmol) in 10 mL of toluene was heated to reflux for 2.5 h giving a dark blue solution. The solvent volume was reduced *in vacuo* to approximately 5 mL and the solution allowed to stand overnight to give dark purple crystals. The crystals were isolated then recrystallised from dichloromethane–methanol at 5 °C yielding 0.129 g (36%) of *mer*-[W(CO)₃(PNCHP- $\kappa^3 P$, *N*, *P'*)]·CH₂Cl₂ as a dark purple crystalline solid, mp 118–123 °C.

cis-[Cr(CO)₄(PNCHP- $\kappa^2 P$,N)] 5. A mixture of [Cr(CO)₄-(NBD)] (0.039 g, 0.151 mmol) and PNCHP (0.083 g, 0.151 mmol) in 12 mL of dichloromethane–acetonitrile (1:1) was stirred at room temperature for 48 h. The resulting dark brownish red precipitate of *cis*-[Cr(CO)₄(PNCHP- $\kappa^2 P$,N)] was isolated by filtration, washed with n-pentane and dried *in vacuo*. The filtrate was allowed to stand for two days, under argon, giving a second crop of dark brownish-red microcrystals. The combined yield was 0.077 g (71%), mp 196–200 °C (decomp.).

cis-[Mo(CO)₄(PNCHP-κ²*P*,*P'*)] 6. A mixture of *cis*-[Mo(CO)₄(pip)₂] (0.201 g, 0.531 mmol) and PNCHP (0.294 g, 0.534 mmol) in 10 mL of dichloromethane was heated to reflux for 15 min. The dark red solution was filtered through Kieselguhr and the solvent volume reduced *in vacuo* to approximately 3 mL. Co-crystallisation of reddish orange and maroon crystals occurred on the addition of *ca*. 1–2 mL of methanol after 3 h at 5 °C. Both crystal types were isolated, washed with cold methanol and dried *in vacuo*, to give a combined yield of 0.364 g (90%). Separation of the two products was achieved by washing away the very soluble maroon product with chloroform, leaving behind the less soluble *cis*-[Mo(CO)₄-(PNCHP-κ²P,P')] as reddish orange crystals, mp 168–170 °C (decomp.).

cis-[W(CO)₄(PNCHP-κ²P,P')] 7. A mixture of *cis*-[W(CO)₄-(pip)₂] (0.201 g, 0.431 mmol) and PNCHP (0.238 g, 0.435 mmol) in 10 mL of toluene was heated to *ca*. 60 °C until all of the *cis*-[W(CO)₄(C₅H₁₁N)₂] had been taken up into solution (*ca*. 30 min). The resulting clear green solution was allowed to cool and then filtered through Kieselguhr. Crystallisation was initiated by solvent volume reduction *in vacuo*. The mixture was then kept at 5 °C for 3–4 h. The reddish orange solid that formed was isolated by filtration and washed with toluene then recrystallised from dichloromethane to yield 0.091 g (25%) of *cis*-[W(CO)₄(PNCHP-κ²P,P')] as a red crystalline solid, mp 187–190 °C (decomp.).

The reaction of $Cr(CO)_6$ with PNCHP to give complexes 10a and 10b. $[Cr(CO)_6]$ (0.223 g, 1.01 mmol) in 30 mL of tetra-

hydrofuran was irradiated using a UV lamp for 1 h, resulting in an orange solution. PNCHP (0.538 g, 0.979 mmol) in 15 mL of tetrahydrofuran was added and the red solution stirred for 30 min. The solution was filtered through Kieselguhr. The solvent was removed and unchanged $[Cr(CO)_6]$ sublimed, *in vacuo*. Preparative scale, thin layer chromatography, eluting with dichloromethane–n-hexane (1:1), was used to isolate a $[Cr(CO)_5(PNCHP-\kappa^1 P)]-[Cr(CO)_5(PCHNP-\kappa^1 P)]$ **10a–10b** mixture (2:1), however, these two isomers could not be separated. In addition, the mixture could not be isolated as an analytically pure solid since conversion into tetracarbonyl species occurred.

The reaction of W(CO)₆ with PNCHP to give complexes 11a and 11b. [W(CO)₆] (0.130 g, 0.367 mmol) in 30 mL of tetrahydrofuran was irradiated for 1 h, resulting in a yellow solution. PNCHP (0.222 g, 0.409 mmol) in 20 mL of tetrahydrofuran was added and the orange solution stirred for 40 min. The solvent and any unchanged [W(CO)₆] were removed *in vacuo*. The resulting orange-red oil was dissolved in hot dichloromethane under a stream of argon gas, then cooled to 5 °C. After 48 h a mixture of [W(CO)₅(PNCHP- $\kappa^1 P$)] 11a and [W(CO)₅(PCHNP- $\kappa^1 P$)] 11b (2:1) was isolated as a pale yellow solid.

[{Cr(CO)₅]₂(PNCHP-κ²P,P')] 8. [Cr(CO)₆] (0.201 g, 0.913 mmol) in 25 mL of tetrahydrofuran was irradiated for 1 h, resulting in an orange solution. PNCHP (0.250 g, 0.455 mmol) in 15 mL of tetrahydrofuran was added and the now orange solution stirred for 30 min. The solvent and any unchanged [Cr(CO)₆] were removed *in vacuo*. The resulting residue was dissolved in toluene and filtered to remove insoluble solids. The filtrate was allowed to stand for 48 h at 5 °C under argon. [{Cr(CO)₅}₂(PNCHP-κ²P,P')] was isolated as yellow crystals, washed with n-pentane and dried *in vacuo* to yield 0.166 g (39%), mp 160 °C (decomp.).

[{W(CO)₅]₂(PNCHP- $\kappa^2 P, P'$)] 9. [W(CO)₆] (0.400 g, 1.14 mmol) in 25 mL of tetrahydrofuran was irradiated for 1 h, resulting in a yellow solution. PNCHP (0.312 g, 0.568 mmol) in 15 mL of tetrahydrofuran was added and the orange solution stirred for 45 min. The solvent and any unchanged [W(CO)₆] were removed *in vacuo*. The resulting residue was dissolved in toluene and filtered to isolate insoluble tan solids. These were dissolved in boiling toluene, under a stream of argon gas, and the resulting solution allowed to stand for 48 h at 5 °C. [{W(CO)₅]₂(PNCHP- $\kappa^2 P, P'$] was isolated as yellow diamond shaped crystals, washed with n-pentane and dried *in vacuo* to give 0.320 g (47%), mp 241–246 °C.

Crystallography

Crystal data. Complex 2, mer- $[Mo(CO)_3(PNCHP-\kappa^3 P, N, P')] \cdot 2C_6H_5Me$. C₅₄H₄₅MoNO₃P₂, M = 913.79, monoclinic, a = 12.490(10), b = 17.680(10), c = 20.630(10) Å, $\beta = 90.76(3)^\circ$, U = 4555(5) Å³, T = 292 K, space group $P2_1/c$, Z = 4, μ (Mo-K α) = 0.40 mm⁻¹, 8394 reflections measured, 8000 unique ($R_{int} = 0.0160$) which were used in all calculations. The final $Rw(F_0^2) = 0.0801$, $R(F_0) = 0.0416$.

Complex 5, cis-[Cr(CO)₄(PNCHP- $\kappa^2 P$,N)]. C₄₁H₂₉CrNO₄-P₂, M = 713.59, monoclinic, a = 14.429(3), b = 29.433(6), c = 17.389(3) Å, $\beta = 108.40(3)^{\circ}$, U = 7007(2) Å³, T = 295 K, space group P_{2_1}/c , Z = 8, μ (Mo-K α) = 0.461 mm⁻¹, 6871 reflections measured, 6533 unique ($R_{int} = 0.0359$) which were used in all calculations. The final $Rw(F_0^2) = 0.1030$, $R(F_0) = 0.0363$.

Complex 7, cis- $[W(CO)_4(PNCHP-\kappa^2 P, P')] \cdot 0.5 CHCl_3$. C_{41.5}-H_{29.5}Cl_{1.5}NO₄P₂W, M = 904.62, monoclinic, a = 23.476(5), b = 18.128(4), c = 19.254(4) Å, $\beta = 116.08(3)^\circ$, U = 7360(3) Å³, space group C2/c, Z = 8, μ (Mo-K α) = 3.38 mm⁻¹, 7068 reflections measured, 3438 unique ($R_{int} = 0.0198$) which were used in all calculations. The final $Rw(F_o^2) = 0.0450$, $R(F_o) = 0.0281$.

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Complex 9, $[\{W(CO)_5\}_2(PNCHP-\kappa^2 P, P')\} \cdot 2C_6H_5Me. C_{61}$ -H₄₄NO₁₀P₂W₂, M = 1378.61, triclinic, a = 10.387(2), b = 12.076(2), c = 12.229 Å, a = 98.79(3), $\beta = 100.59(3)$, $\gamma = 107.99(3)^\circ$, U = 1397.6(4) Å³, T = 290 K, space group $P\overline{1}$, Z = 1, μ (Mo-Ka) = 4.23 mm⁻¹, 2754 reflections measured, 2588 unique ($R_{int} = 0.0271$) which were used in all calculations. The final $Rw(F_0^2) = 0.0653$, $R(F_0) = 0.0384$.

X-Ray analysis. Single crystals of the complexes were grown as described under their respective syntheses and attached to the ends of glass fibres with cyanoacrylate glue. Data collection, solution and refinement were performed as previously described.³⁴⁻³⁶ Solvent molecules were located in 2, 7 and 9. In complex 2 two molecules of toluene were found, the first of which appeared to show little signs of disorder. However the second was disordered over two sites with occupancy factors of 0.579 and 0.421 and the atoms of each ring were refined as a rigid group. In 7 a half-weighted chloroform molecule was located. The half-weighted nature results from the close approach (C(81)–C(81') 0.87 Å) of a symmetry-related chloroform molecule in the unit cell. Isotropic thermal motion was assumed for C(81) of this group. In 9 the complex lies across a centre of symmetry with the centre at the midpoint of the C-N bond. No attempt was made to distinguish the C and N atoms by assigning the space group as P1 and they were included in the calculations as C. The molecule of toluene located in the asymmetric unit showed little signs of disorder. No solvent was located in 5.

CCDC reference number 186/2053.

See http://www.rsc.org/suppdata/dt/b0/b002427i/ for crystallographic files in .cif format.

Kinetic and equilibrium studies of complex 4

Acetone was freshly distilled over 4A molecular sieves, chlorinated solvents and acetonitrile were freshly distilled over CaH₂ and all were degassed with argon. Stock solutions of fac- $[Mo(CO)_3(PNCHP-\kappa^3 P, N, P')]$ 4 were prepared by dissolving an appropriate amount of the solid complex immediately before each experiment. Samples were transferred to a 1 cm path-length sealed quartz cuvette placed in the constant temperature (±3%) jacket of the spectrometer. Kinetic runs as a function of temperature (19.5-49.5 °C) were performed with a HP 8452A uv-vis spectrometer at 700 nm. Absorbance readings were taken until the reaction had progressed to equilibrium. Values of k_{obsd} were obtained from linear plots of $\ln(A - A_{\infty})$ against time, using a minimum of 6 absorbance/time pairs. The first-order rate constants were calculated using a linear regression method. All plots gave correlation coefficients greater than 0.9993. Activation energy was calculated from an Arrhenius plot and activation enthalpy and entropy from an Eyring plot.

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