



## COMMUNICATION

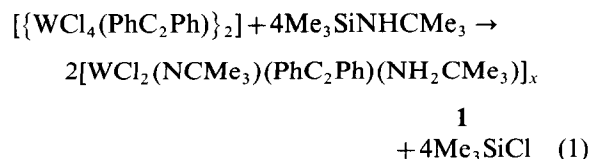
A  $d^0$  TO  $d^2$  TRANSFORMATION OF TUNGSTEN,  
PROMOTED BY FORMATION OF AN ORGANOIMIDO  
LIGAND AND INVOLVING DISRUPTION OF THE  
 $\pi$ -PERPENDICULAR BONDING COMPONENT OF  
CO-ORDINATED DIPHENYLACETYLENEALASTAIR J. NIELSON,\* PETER D. W. BOYD, GEORGE R. CLARK,  
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**Abstract**—Synthetic, spectroscopic and structural investigations as well as theoretical calculations show that reaction of the  $d^0$ , four-electron donor alkyne complex  $[\{WCl_4(PhC_2Ph)\}_2]$  with the silylamines  $Me_3SiNHR$  ( $R = CMe_3, CHMe_2, CH_2Me, 2,6-CHMe_2Ph$ ) leads to  $d^2$ , two-electron donor alkyne complexes containing an organoimido ligand.

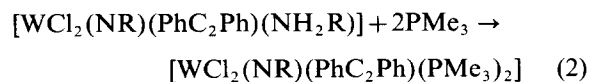
We have recently reported confirmatory evidence for the stabilisation of high oxidation states of tungsten by alkynes, in a manner comparable with an oxo or organoimido ligand.<sup>1</sup> For example the alkyne complex  $[\{WCl_4(PhC_2Ph)\}_2]$  shows similar chemistry and properties to  $[\{WCl_4(NPh)\}_2]$ <sup>2</sup> and may thus be regarded as a  $d^0$  complex. The alkyne–tungsten bonding in  $[\{WCl_4(PhC_2Ph)\}_2]$  incorporates the alkyne  $\pi_\perp$ -bonding components so that the diphenylacetylene ligand acts as a four-electron donor.<sup>3</sup> We report here the disruption of this alkyne–tungsten  $\pi_\perp$ -donor interaction, by an organoimido ligand which results in a  $d^0$  to  $d^2$  transformation of tungsten.

Reaction of  $[\{WCl_4(PhC_2Ph)\}_2]$  with *N*-trimethylsilyl-*t*-butylamine in benzene proceeds according to reaction (1)

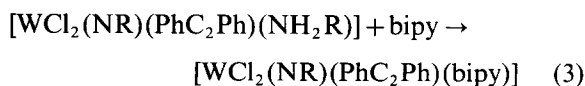


The IR spectrum of complex **1** contains a single W—Cl stretch characteristic of *trans*-chloro ligands while the *t*-butylimido and *t*-butylamine ligands are confirmed by specific quaternary carbon resonances in the  $^{13}C\{-^1H\}$  NMR spectrum.<sup>4</sup> The acetylenic carbons appear at  $\delta$  156 which is at a similar position to that found for the  $d^2$ , two-electron donor alkyne complex  $[WCl_2(NPh)(PhC_2Ph)(PMe_3)_2]$  ( $\delta$  156).<sup>5</sup> In the four-electron donor alkyne complex  $[WCl_5(PhC_2Ph)]NEt_4$ , a soluble derivative of  $[\{WCl_4(PhC_2Ph)\}_2]$ , the acetylenic carbon resonance occurs at  $\delta$  270. The  $^{13}C\{-^1H\}$  NMR spectra thus indicate the alkyne changes from a four-electron donor to a two-electron donor in reaction (1). Analogues of complex **1** with acetylenic carbon resonances in the vicinity of  $\delta$  156 can also be prepared from  $[\{WCl_4(PhC_2Ph)\}_2]$  and the silylamines  $Me_3SiNHR$  ( $R = CHMe_2$  and  $Et$ ).  $Me_3SiNH-2,6-CHMe_2Ph$  reacts similarly but requires heating to proceed.

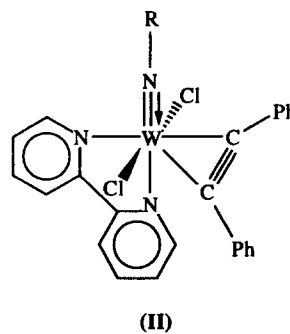
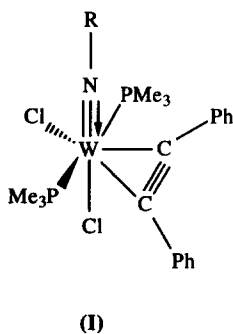
Derivatives of these complexes can be prepared by replacement of the amine ligand with other  $\sigma$ -donor ligands [reactions (2) and (3)].



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IR spectra of the phosphine adducts show two W—Cl stretches indicating *cis*-chloro ligands and the NMR spectra show *trans*-phosphines (structure I). Conversely, the bipy derivatives show only one W—Cl stretch consistent with *trans*-chloro ligands (structure II). These structures have been confirmed by X-ray crystallography and will be reported elsewhere.



The  $^{13}\text{C}$ - $\{^1\text{H}\}$  NMR spectra of these derivatives again show acetylenic carbon resonances in the vicinity of  $\delta$  156. X-ray photoelectron spectroscopy (XPS) shows the complexes have  $\text{W}(4f_{7/2})$  binding energies in the range considered to be tungsten(IV). For example the  $\text{W}(4f_{7/2})$  binding energy found for  $[\text{WCl}_2(\text{NCMe}_3)(\text{PhC}_2\text{Ph})(\text{bipy})]$  is 33.8 eV which is below that observed for  $[\{\text{WCl}_4(\text{PhC}_2\text{Ph})\}_2]$  where the  $\text{W}(4f_{7/2})$  binding energy (35.1 eV)<sup>1</sup> falls in the range considered to be tungsten(VI).<sup>6</sup>

Hartree Fock (HF)<sup>7</sup> as well as scattered wave  $\text{X}\alpha$  calculations ( $\text{MSX}\alpha$ ),<sup>8</sup> suggest that the model complex  $[\text{WCl}_2(\text{NCH}_3)(\text{HC}_2\text{H})(\text{PH}_3)_2]$  (3) is best described as a  $d^2$  tungsten system. The  $5d$  populations,  $n_{5d}$ , are presented in Fig. 1 along with data obtained from  $\text{SWX}\alpha$  calculations made on  $\text{WCl}_6$

and the  $d^0$ ,  $d^1$  and  $d^2$ , four-electron donor alkyne model systems  $[\text{WCl}_5(\text{HC}_2\text{H})]^-$ ,  $[\text{WCl}_3(\text{HC}_2\text{H})(\text{PH}_3)_2]$  and  $[\text{WCl}_2(\text{HC}_2\text{H})(\text{PH}_3)_3]$ .<sup>1</sup> The  $n_{5d}$  value for the imido complex is clearly greater than that obtained for  $[\text{WCl}_5(\text{HC}_2\text{H})]^-$ . The calculated charge for the imido complex ( $\text{SWX}\alpha$  calculation) is significantly smaller than that found for  $[\text{WCl}_5(\text{HC}_2\text{H})]^-$  and this difference correlates well with the lower  $\text{W}(4f_{7/2})$  binding energy found for the analogous complex  $[\text{WCl}_2(\text{NCMe}_3)(\text{PhC}_2\text{Ph})(\text{bipy})]$ .

The presence of the imido ligand, another  $\pi$ -donor, significantly affects the characteristics of the acetylene ligand bond. The imido ligand Frontier Molecular Orbitals (FMOs) are able to interact extensively with the same  $d$  Atomic Orbitals (dAOs) as the acetylene ligand. Population analysis shows a distinct difference in the ligand donor properties in  $[\text{WCl}_5(\text{HC}_2\text{H})]^-$  and  $[\text{WCl}_2(\text{NMe})(\text{HC}_2\text{H})(\text{PH}_3)_2]$ . In both complexes  $\text{W}(5d_\pi)$  to  $\text{C}_2\text{H}_2(\pi^*)$  back donation ( $b_2$  interaction in the local  $\text{C}_{2v}$  point group) and  $\text{C}_2\text{H}_2(\pi_\parallel)$  to  $\text{W}(5d_\pi)$  donation ( $a_1$  interaction) operate, but in the imido complex  $\text{C}_2\text{H}_2(\pi_\perp)$  to  $\text{W}(5d_\pi)$  donation ( $b_1$  interaction) is minimised as the nitrogen  $p_z$  ( $\text{N}_{p_z}$ ) orbital overlap with the  $d_{yz}$  AO dominates (Fig. 2).  $\text{W}(5d_\pi)$  to  $\text{C}_2\text{H}_2(\pi^*)$  back-bonding is activated by overlap of the  $\text{N}_{p_x}$  donor

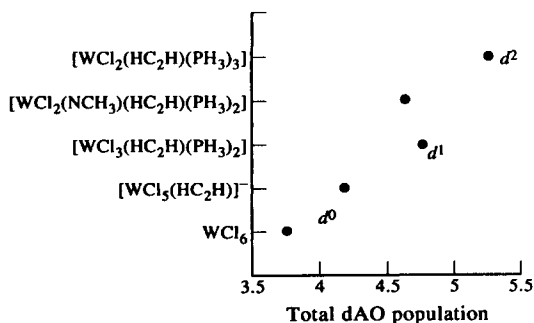


Fig. 1.

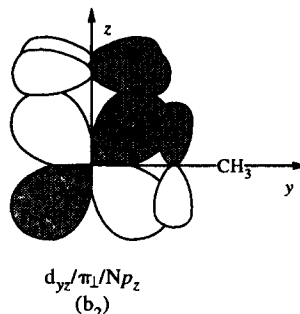


Fig. 2.

orbital with the  $d_{xy}$  AO which also overlaps with the  $\pi_{\perp}^*$  FMO of acetylene.

The combined results of these studies show that the four-electron donor alkyne ligand in  $[\{\text{WCl}_4(\text{PhC}_2\text{Ph})\}_2]$  may be formally considered a net two-electron donor alkyne after an organoimido ligand is added to the complex. The properties associated with  $[\{\text{WCl}_4(\text{PhC}_2\text{Ph})\}_2]$  all point to a  $d^0$  system, formally tungsten(VI).<sup>1</sup> Addition of the organoimido ligand disrupts the alkyne-metal  $\pi_{\perp}$  bonding component leading to a  $d^2$  complex formally considered as tungsten(IV). Alternatively, the alkyne ligand in  $[\{\text{WCl}_4(\text{PhC}_2\text{Ph})\}_2]$  may be regarded as being in a reduced state compared with the alkyne ligand in the organoimido complexes. Overall, reaction (1) represents a change at the metal centre, effectively of two electrons, and thus the alkyne ligand in  $[\{\text{WCl}_4(\text{PhC}_2\text{Ph})\}_2]$  may be regarded as an internal redox ligand.

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