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Dimethyldioxirane oxidation of phosphide PMo[N(R)Ar]₃ $[R = C(CD_3)_{2}Me, Ar = C_6H_3Me_2-3.5]$ efficiently provides **(OP)Mo[N(R)Ar]3, a complex shown by X-ray crystallography to contain a terminal phosphorus monoxide ligand, and which reacts with dimethylzirconocene to give addition of a Zr–Me bond across the phosphoryl moiety.**

Known phosphorus monoxide complexes^{$1-3$} are those in which PO^{4-10} serves as a triply bridging ligand. These have been prepared by αx transfer to a $P_2N_i^2W$ cluster,¹ and by hydrolysis of a μ_3 -P(NPrⁱ₂) functional group.³ Recent reports of terminal phosphide (P^{3-}) complexes^{11–15} hint at the possibility of preparing terminal PO complexes by oxygen-atom transfer to the terminal P ligand. Accordingly, dimethyldioxirane oxidation16,17 has been found to effect the desired conversion of PMo[N(R)Ar]3 **1** to (OP)Mo[N(R)Ar]3 **2**. Synthetic, structural, and initial reactivity data pertaining to **2** are detailed herein.

Treatment of a -78 °C gold-colored solution of 1 (1.0009 g, 1.4853 mmol) in dichloromethane (50 ml) with dimethyldioxirane (20 ml of a 0.075 m solution in acetone18) elicited a rapid color change to purple, the color of **2**.‡ Workup consisted of cold-filtration, precipitation of **2** by addition of acetonitrile (75 ml, pre-chilled to just above its melting point), and collection of solid **2** (73.8% yield) by filtration. Once thus obtained in pure form, solid **2** can be manipulated and maintained at 25 °C for at least *ca.* 5 days without noticeable decomposition. When dissolved in various solvents (*e.g.* benzene, diethyl ether, heptane and dichloromethane) **2** is more sensitive, undergoing gradual reversion to **1** according to 1H and 31P NMR spectroscopic monitoring.

Diamagnetic 2 exhibits a single ³¹P NMR signal at δ 269.8, shifted dramatically upfield relative to **1** but in a range typical of

Fig. 1 ORTEP drawing of the molecular structure of **2** with 35% probability ellipsoids. Selected bond lengths (Å) and angles (°): Mo–P 2.079(5), Mo– N(1) 1.95(2), Mo–N(2) 1.94(2), Mo–N(3) 1.994(13), P–O 1.49(2); Mo– P–O 177.6(8), P–Mo–N(1) 101.7(4), P–Mo–N(2) 101.4(5), P–Mo–N(3) 101.6(5), N(1)–Mo–N(2) 113.1(6), N(1)–Mo–N(3) 117.2(6), N(2)–Mo– N(3) 118.0(6). **Scheme 1**

two-coordinate phosphorus compounds.19 Proton NMR data for **2** are indicative of a single environment for the N(R)Ar ligands, consistent with the threefold (propellor) symmetry typical of complexes containing three $N(\overline{R})$ Ar ligands.^{20–25}

Single crystals of **2** suitable for an X-ray diffraction study§ were obtained by slow cooling to -35 °C of a diisopropyl ether solution. Molecular **2** evinces pseudo-threefold symmetry about the near-linear $[177.6(8)°]$ Mo–P–O moiety, which is flanked by the three *tert*-butyl substituents (Fig. 1). An intriguing feature of the structure is the molybdenum–phosphorus distance $d(MoP) = 2.079(5)$ Å, which is shorter by 0.04 Å than $d(MoP)$ for terminal phosphide **1**.11 Contraction of the molybdenum– phosphorus distance in this manner is explicable in terms of rehybridization at P upon addition of a terminal substituent.26 The phosphorus–oxygen distance [1.49(2) Å] in **2** is comparable to that measured by IR diode laser spectroscopy for gaseous PO [1.476370(15) Å)],^{27,28} and to $d(PO)$ values for μ_3 -PO complexes: $1.509(3)$,³ $1.462(9)$ and $1.480(10)$.¹ Phosphoryl halide species such as $OPCl₃$ contrastingly exhibit longer phosphorus–oxygen distances, on the order of 1.54–1.58 \AA ²⁹ while $d(PO)$ for $OP(C₆H₁₁)₃$ is 1.490(2) Å.³⁰ Other metrical parameters pertaining to the structure of **2** are unexceptional.

Previously reported $(SP)Mo[N(R)Ar]_3$ 3¹¹ was shown recently by X-ray crystallography to resemble **2** in terms of its monomeric nature and gross structural characteristics.31 The rough similarity of **2** and **3** was used to advantage to obtain assignment of the v_{PO} (1232 cm⁻¹) and v_{PS} (782 cm⁻¹) IR stretching frequencies for the two complexes, by digital subtraction of solution FTIR spectra. The v_{PO} value for 2 is similar to that reported for gaseous PO $(1220.25 \text{ cm}^{-1})$, and also is comparable to values reported for μ_3 -PO complexes $(1260, 1169$ cm⁻¹).

Initial reactivity studies indicate that the phosphoryl phosphorus in **2** is susceptible to nucleophilic attack in a manner reminiscent of the Fischer carbene synthesis.32 Accordingly, treatment of **2** (282.0 mg, 0.4088 mmol) with dimethylzirconocene³³ (103.3 mg, 0.4108 mmol) in benzene (7.5 ml) led to formation of $[(\overline{R_1})(R_2O)P]Mo[N(R)Ar]_3$ [4, $R_1 = Me$, $R^2 = Zr(Me)(\eta - C_5H_5)_2$, which was isolated by crystallization

Fig. 2 ORTEP drawing of the molecular structure of **4** with 35% probability ellipsoids. Selected bond lengths (Å) and angles (°): Mo–P 2.169(2), Mo– N(1) 1.995(5), Mo–N(2) 2.011(5), Mo–N(3) 1.993(5), P–O 1.613(5), P–C(1) 1.832(8), Zr–O 2.000(4), Zr–C(2) 2.253(11); Mo–P–O 127.7(2), Mo–P–C(1) 133.9(3), O–P–C(1) 97.6(3), Zr–O–P 163.6(3), O–Zr–C(2) 97.1(4), P–Mo–N(1) 101.6(2), P–Mo–N(2) 103.2(2), P–Mo–N(3) 101.7(2), N(1)–Mo–N(2) 109.1(2), N(1)–Mo–N(3) 112.6(2), N(2)–Mo–N(3) 125.0(2).

from a 1:1 diethyl ether–heptane mixture as orange–brown plates in 75.0% yield (Scheme 1). Diamagnetic **4** exhibits a single ³¹P NMR signal at δ 284.4 {*cf.* δ 312 for $Mo[P(C₆H₁₁)₂]₄³⁴$. A reasonable oxidation state assignment for the Mo centre in **4** is +4. Proton NMR data for **4** are indicative of a single N(R)Ar ligand environment on the NMR timescale, although the solid-state structure of **4** reveals a lowsymmetry conformation. A doublet (${}^{2}J_{\text{PH}}$ 14 Hz) at δ 1.535 is attributed to the PMe moiety.

The three-coordinate phosphorus in **4** was shown by X-ray diffraction§ to be essentially planar [sum of angles = 359.2(3) $^{\circ}$], consistent with Mo–P π bonding in the complex (Fig. 2). The observed *d*(MoP) of 2.169(2) Å is also consistent with a molybdenum–phosphorus multiple bond {*cf.* $d(MoP)$ of 2.265(2) Å for $Mo[P(C_6H_{11})_2]_4^{34}$. The conformation of 4 is noteworthy in that one of the $N(R)$ Ar ligands is oriented with its *tert*-butyl group distal from phosphorus, presenting an Ar π cloud to the methyl substituent on phosphorus. The Zr–O–P(Me)–Mo core in **4** is reminiscent of $Zr-O-C(R')-M$ cores in zirconoxy carbenes prepared by addition of $Zr-R'$ ($R' = H$, alkyl) bonds across the CO unit in metal carbonyl complexes.35

We thank Monsanto Company for support of this work, and Michael K. Stern for helpful discussions. M. J. A. J. thanks NSERC for a predoctoral fellowship.

Footnotes and References

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- ‡ All manipulations were carried out under an atmosphere of dry nitrogen using solvents purified by standard methods. The new compounds **2** and **4** gave satisfactory elemental analyses (C, H and N).

§ *Crystallographic data*: For 2: triclinic, space group $P\overline{1}$, $a = 10.7615(8)$, $b = 11.4942(9), c = 15.1899(11)$ Å, $\alpha = 89.865(2), \beta = 80.838(2),$ $\gamma = 89.4900(10)^\circ$, $U = 1854.9(2)$ \AA^3 , $Z = 2$, 2988 unique reflections, data/

parameter ratio = 7.9, $R_1 = 0.1207$, $wR_2 = 0.2683$, GOF 1.050, residuals based on $I > 2\sigma(I)$.

For **4**: orthorhombic, space group *Pbca, a* = 11.3996(2), *b* = 23.2075(4), $c = 39.2443(7)$ Å, $U = 10382.3(3)$ Å³, $Z = 8$, 7456 unique reflections, data/parameter ratio 15, $R_1 = 0.0669$, $wR_2 = 0.2023$, GOF = 1.269, residuals based on $I > 2\sigma(I)$. CCDC 182/527.

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Received in Bloomington, IN, USA, 7th May 1997; 7/03105J