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Dimethyldioxirane oxidation of phosphide $PMo[N(R)Ar]_3$ [R = C(CD₃)₂Me, Ar = C₆H₃Me₂-3,5] 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 oxo transfer to a P₂Ni₂W 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 oxidation^{16,17} 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).

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(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 molybdenumphosphorus 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 Å,²⁹ while d(PO) for $OP(C_6H_{11})_3$ 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 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 $[(R^1)(R^2O)P]Mo[N(R)Ar]_3$ [**4**, R¹ = Me, R² = 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 low-symmetry conformation. A doublet (²*J*_{PH} 14 Hz) at δ 1.535 is attributed to the PMe moiety.

The three-coordinate phosphorus in 4 was shown by X-ray planar diffraction§ to be essentially 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(C₆H₁₁)_{2]4³⁴}. The conforma-} tion 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

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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 PI, 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) Å³, 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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