Ligand-controlled Synthesis, Reactivity and Oxo-Transfer Kinetics of Oxomolybdenum-(vi) and -(iv) Complexes

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Trifunctional (ONS) dianionic Schiff-base ligands $L^{2-}[H_{2}L = S$ -methyl 3-(2-hydroxyphenyl)methylenedithiocarbazate or 5-R substituent derivatives (R = H, Me, Cl, Br or NO₂) or a naphthyl derivative], in sharp contrast to the S-benzyl analogues (H_2L') form only Mo-O-Mo bridged oligomers [(MoO₂L),] in EtOH or MeOH, irrespective of the substituent R. However, these substituents R control the position of the $v(Mo - 0 \rightarrow Mo)$ vibration, the $Mo^{v_1} - Mo^{v}$ redox couple, the ligand-to-metal charge-transfer transition, as well as the chemical shift of the azomethine proton signal and the asymmetric v(Mo=0) vibration in solution, for both the polymetric complexes and, where relevant, the donor molecule (D) co-ordinated monomers [MoO₂L(D)] [D = pyridine (py), dimethylformamide (dmf) or Me₂SO]. Reaction of [MoO₂L] with PPh₃ in CH₂Cl₂, MeOH or MeCN or in donor solvents D (dmf or py) produced oxomolybdenum(iv) derivatives, [MoO(L)] or [MoOL(D)], respectively. The kinetics of oxo transfer from MoO₂²⁺ to PPh₃ occurs in a second-order process. The rate constant of the oxo-transfer reaction from the polymer $[(MoO_2L)_n]$ (R = H) to the PPh₃ substrate is $\approx 10^2$ times higher than that of the corresponding monomer $[MoO_2L(D)]$. Both [MoO(L)] and [MoOL(dmf)] react with the Me₂SO substrate in CH₂Cl₂ or dmf in a two-stage process. The first involves the equilibrium formation of a Me₂SO adduct while during the second stage an intramolecular oxo transfer occurs from Me₂SO to the MoO core via the elimination of Me₂S. The rate constant of the reverse oxo transfer (k_{-1}) is almost identical for both the polymer [MoO(L)] and monomer [MoOL(dmf)] but the equilibrium constant, K, for the formation of the Me₂SO complexed species is slightly higher for [MoOL(dmf)] than for [MoO(L)].

Oxo-transfer chemistry of molybdenum, and, to a lesser extent that of tungsten, ruthenium, osmium and rhenium, is of topical interest.¹ Molybdenum has a special place in this type of chemistry owing to the biological relevance of the model reaction $(1)^{\dagger}$ involving oxo-transfer molybdoenzymes.^{2–5} The

 $[Mo^{VI}O_2(ligand)_n] + X \rightleftharpoons [Mo^{IV}O(ligand)_n] + XO \quad (1)$

oxo-transfer ability of the MoO_2^{2+} core depends on the functionalities present in the ligands. In a number of model studies,⁶⁻⁹ the indispensibility of at least one sulfur donor was realised.¹⁰ Recently, however, it has been shown that the NNN functionalities generated by a pyrazolylborate ligand¹¹ also activate the MoO₂²⁺ unit towards oxo transfer. Extended X-ray absorption fine structure (EXAFS) spectroscopic studies have also implicated the presence of a sulfur atom, besides oxygen and nitrogen, at the active sites of oxo-transfer molybdoenzymes.¹²⁻¹⁵ We have shown ¹⁶ for the ONS-donor dianions of the Schiff-base ligands, S-benzyl 3-(5-R-2hydroxyphenyl)methylenedithiocarbazate, H2L', that phenyl substituents R (R = H, Me, Cl or Br) play a vital role in controlling whether the isolated dioxomolybdenum(vi) complexes are discrete monomers $[MoO_2L'(D)]$ (D = weak donor solvent molecule, e.g. EtOH or MeOH) or polymeric species, $[(MoO_2L')_n]$, containing ... (O)Mo=O→Mo(O)=O... chains, observed previously^{17,18} in [MoO₂(tridentate)] species. The electron-releasing 5-Me substituent yielded a polymer, whilst electron-withdrawing substituents (5-Cl or 5-Br) favoured the formation of monomers.

The present work shows that if the electron-releasing methyl group is substituted for benzyl at the unco-ordinated S atom of H_2L' the new ligands H_2L , shown in thione and thiol form



as A and B, respectively, afford only polymers, $[(MoO_2L)_n]$, even if the 5-R substituent is a strong electron withdrawer such as NO₂. However, for both sets of ligands H_2L' and H_2L it is possible to correlate the Hammett σ_p values of the 5-R substituents with various physico-chemical and spectroscopic parameters of the isolated complexes $[MoO_2L]$ or $[MoO_2L']$. This correlation study has a practical significance: the remotecontrol effect of the S- and phenyl-substituents on the structures and chemical and spectroscopic properties of the oxomolybdenum complexes studied here may help to identify the presence of different fragments or substituents attached to the active sites of relevant molybdoenzymes. The present work also shows that the oxo-transfer rate constant from MoO₂ to the substrate is much higher when the oxygen atom is transferred from an oxygen-bridged oligomer, $[(MoO_2L)_n]$ than from a discrete monomer, $[MoO_2L(dmf)]$ (dmf = dimethylformamide). A number of reports concerning substituent effects 6a,b,19 and oxotransfer kinetics 6c,20 in other oxomolybdenum systems, have been published previously.

[†] Model studies mainly concern tertiary phosphines as the substrate X.

Experimental

Physical Measurements.—Infrared spectra were recorded as KBr discs on a Perkin-Elmer 597 spectrophotometer, and in dmf solution on a Perkin-Elmer 1710 FT-IR spectrophotometer using a demountable Irtran 2 (ZnS) cell, electronic spectra on a Hitachi U-3400 UV/VIS/NIR spectrophotometer, ¹H NMR spectra using a Varian EM 390 (90 MHz) spectrometer and SiMe4 as standard. Thermoanalyses were made on a Shimadzu thermoanalyser DT 30. Voltammetric measurements were made with a PAR model 370-4 electrochemistry system: model 174A polarographic analyser, model 175 universal programmer, model RE 0074 XY recorder, model 173 potentiostat, and model 377A cell system. All experiments were performed at 298 K under a dinitrogen atmosphere in a three-electrode configuration using a planar Beckman model 39273 platinum-inlay working electrode, a platinum-wire auxiliary electrode and a saturated calomel reference electrode (SCE). Coulometry was carried out using a 273 potentiostat/galvanostat with a 377 cell system equipped with a platinum-wire-gauze working electrode. The potentials reported are uncorrected for the junction contribution. Magnetic susceptibilities were obtained by the Gouy method using Hg[Co(NCS)₄] as the calibrant. Solution conductances were measured with a Systronics (India) model 304 digital conductivity meter. A Knauer (Berlin) vapour-pressure osmometer was used for the molecular weight determination, calibrated with benzyl solution in an appropriate solvent. Elemental analyses were performed using a Perkin-Elmer 240C elemental analyser and molybdenum was estimated by a gravimetric method.²¹ The kinetic measurements were performed under a dinitrogen atmosphere employing 2 \times 10⁻⁴ mol dm⁻³ dmf-CH₂Cl₂ solutions of the dioxomolybdenum(vi) and oxomolybdenum(IV) complexes. Pseudo-first-order conditions were maintained throughout by use of an excess of PPh₃ over the molybdenum concentration. The time dependence of the absorbance at 464 nm (for complexes of H_2L^1 and H_2L^2) and 462 nm (for complexes of H_2L^3 and H_2L^4) were recorded on a Philips Analytical SP8-150 UV/VIS spectrophotometer with a thermostatted (HAAKE F3) cell compartment. The observed rate constants were obtained from plots of $\ln(A_{\infty} - A_{t})$ or $\ln(A_t - A_{\infty})$ vs. time, where A_t and A_{∞} are the absorbances at time t and infinity, respectively. These were all linear up to four half-lives of the reaction. Data were extracted using a leastsquares computer-fitting program.

Materials.—Ammonium heptamolybdate (J. T. Baker, Phillipsburg, NJ), p-chlorophenol (AR BDH, UK) and p-cresol (S. D. Chemicals, India) were obtained commercially and used as such. The complexes $[MoO_2(acac)_2]$ (acac = acetylacetonate),¹⁷ [MoOL'] (L' = L^{1'}-L^{4'}),* S-methyldithiocarbazate²² and 5-nitrosalicylaldehyde²³ were prepared by the literature methods. 5-Bromosalicylaldehyde was obtained from Aldrich (USA) and the corresponding 5-methyl, 5-chloro and 5,6-C₄H₄ derivatives were prepared in good yields by employing the Duff reaction.²⁴ Triphenylphosphine (Sisco Research Laboratories, India) was recrystallized twice from ethanol-water. Dimethylformamide used for the electrochemical and kinetics studies was dried by distillation *in vacuo* over P₂O₅ and then stored over Linde AW-500 molecular sieves. Dimethyl sulfoxide was distilled from CaH₂, degassed and stored under dinitrogen.

Preparation of the Ligands $H_2L^1-H_2L^6$.—The ligands were obtained via a Schiff-base condensation reaction between the appropriate aldehyde (0.02 mol) and S-methyldithiocarbazate (0.02 mol) in ethanol. Yields: 85–90% [Found: C, 47.9; H, 4.4; N, 12.4. C₉H₁₀N₂OS₂(H₂L¹) requires C, 47.7; H, 4.4; N, 12.4.

Found: C, 50.2; H, 5.0; N, 11.7. $C_{10}H_{12}N_2OS_2(H_2L^2)$ requires C, 50.0; H, 5.0; N, 11.6. Found: C, 41.4; H, 3.5; N, 10.8. $C_9H_9CIN_2OS_2(H_2L^3)$ requires C, 41.4; H, 3.5; N, 10.7. Found: C, 35.4; H, 3.3; N, 9.3. $C_9H_9BrN_2OS_2(H_2L^4)$ requires C, 35.4; H, 3.0; N, 9.2. Found: C, 40.0; H, 3.3; N, 15.6. $C_9H_9N_3O_3S_3(H_2L^5)$ requires C, 39.8; H, 3.3; N, 15.5. Found: C, 56.7; H, 4.4; N, 10.3. $C_{13}H_{12}N_2OS_2(H_2L^6)$ requires C, 56.5; H, 4.4; N, 10.1%].

Preparation of the Complexes $[MoO_2L]$ (L = L¹ 1a, L² 1b, L³ 1c, L⁴ 1d, L⁵ 1e or L⁶ 1f).—To a hot solution of the ligand (1 mmol) in ethanol or methanol (20 cm³), was added a solution of $[MoO_2(acac)_2]$ (0.32 g, 1 mmol) in the same solvent (10 cm³). The reaction mixture was gently refluxed for 30 min. It was then allowed to stand at 5 °C overnight whereupon the brown crystalline products formed were filtered off, washed with ethanol and dried *in vacuo*. Yield: 60–65%.

 $[MoO_2L^1(D)]$ $[D = pyridine (py) 1g, dmf 1h or Me_2SO 1i]$ and $[MoO_2L(py)]$ $(L = L^2 1j, L^3 1k or L^4 1l)$. These compounds were prepared by treating $[MoO_2L]$ with an excess of the monodentate donor ligands in ethanol. Yields: 60-65%.

[MoO(L¹)] **2a**. A solution of $[MoO_2L^1]$ (1 mmol) was dissolved in either methanol, dichloromethane or acetonitrile (50 cm³) by heating. Triphenylphosphine (0.4 g, 1.5 mol) was added as a solid to the solution with constant stirring. The reaction mixture was heated on a water-bath for 20 min producing a dark brown compound. The resulting precipitate was filtered off, and washed with ethanol. Yield: 85%.

[MoO(L)] ($\dot{L} = L^2$ 2b, L^3 2c or L^4 2d). Complexes, 2b–2d, were prepared directly from [MoO₂(acac)₂] by refluxing first with the respective ONS ligand and then with triphenyl-phosphine in methanol. Yields: 80%.

[MoOL(D)] (L = L¹, D = dmf 2e or py 2f; L = L⁴, D = dmf 2g or py 2h). These compounds were prepared in the same manner as described for 2a, using pyridine or dmf instead of methanol, acetonitrile or dichloromethane. Yields: $\approx 80\%$.

The analytical data for the isolated complexes are shown in Table 1.

Results and Discussion

(a) Oxomolybdenum(VI) Complexes.—Like H_2L' ,¹⁶ the free ligands $H_2L^1-H_2L^6$ (see structures **A** and **B**) used in this work exhibit identical vibrational²⁵ spectra characterising their functionalities. After metal co-ordination the vibrational bands are identically modified as for the H_2L' complexes¹⁶ indicating the attachment of H_2L ligands as L^{2-} obtained via the thiol form²⁶ **B**. The characteristic blue and red shifts¹⁶ of the specific ligand functions^{25,27} upon co-ordination, indicated that the MoO_2^{2+} group is bonded through the phenolic oxygen and azomethine nitrogen, besides the thiol sulfur. Moreover, comparison of the ¹H NMR spectra of $H_2L^1-H_2L^5$ with those of the complexes 1a-1e reveals that the signals of (*i*) the phenolic protons, (*ii*) the azomethine protons²⁸ (Table 2) and (*iii*) the position of the -SMe proton resonance in all the ligands, occurring at *ca*. δ 2.72, undergo virtually identical shifting as observed for the [MoO_2L'] complexes,¹⁶ supporting an ONS mode of co-ordination suggested from the IR data. Further endorsement comes from the appearance of new bands at 550– 600, 415–440 and 340–345 cm⁻¹ due to v(Mo–O),¹⁶ v(Mo–N) and v(Mo–S)¹⁶ stretches, respectively, in 1a-1f.

All the complexes 1a-1f show a typical band at 805 cm⁻¹ assignable to the Mo...O-Mo vibration,¹⁷ besides a terminal v(Mo=O) vibration at *ca*. 935 cm⁻¹, indicating the formation of polymeric complexes $[(MoO_2L)_n]$ containing ... (O)Mo=O-Mo(O)... chains.^{17,18} Hence, the structure of 1a-1f can be represented as C where the ONS functionalities span the three meridional positions in a pseudo-octahedral geometry. Curiously, in the polymeric $[(MoO_2L')_n]$ complexes¹⁶ there appear three other bands in the v(Mo=O) vibrational region, besides the band at 805 cm⁻¹ due to the bridging oxo group, whilst in the present complexes, 1a-1f, only one such v(Mo=O)

^{*} Ligands $L^{1'}\!-\!L^{4'}$ are the -SCH_2Ph analogues of the -SMe ligands $L^1\!-\!L^4.$

Table 1 Analytical,^a IR $(cm^{-1})^{b}$ and electronic spectral data^c for the complexes

	Analysis (%)				Colocted		1 /am 1	
Complex	С	н	N	Мо	$v(Mo=O)^d$	D vibrations	v(C–O)	$(\log \varepsilon/dm^3 \text{ mol}^{-1} \text{ cm}^{-1})$
$1a [MoO_2L^1]$	30.7 (30.6)	2.2 (2.3)	8.0 (7.9)	27.2 (27.1)	930, 810°	_	1555m	23 696 (3.04)
1 b [M ₀ O ₂ L ²]	33.0 (32.8)	2.8 (2.7)	7.6 (7.6)	25.9 (26.2)	935, 810°		1560m	23 256 (3.10)
	28.2 (27.9)	1.9 (1.8)	7.0 (7.2)	24.4 (24.8)	938, 800 °	_	1556m	23 866 (3.07)
ld [MoO₂L ⁴]	25.2 (25.1)	1.7 (1.6)	6.2 (6.5)	22.0 (22.2)	935, 800°		1555m	23 866 (3.07)
$1e [MoO_2L^5]$	27.3 (27.2)	1.9 (1.8)	10.4 (10.6)	23.8 (24.1)	920, 800 <i>°</i>		1557m	24 760 (3.18)
1f [MoO2L ⁶]	39.0 (38.8)	2.5 (2.5)	7.0 (6.9)	24.0 (23.8)	920, 820 <i>°</i>		1558s	21 740 (3.10)
$lg [MoO_2L^1(py)]^f$	38.8 (39.0)	3.1 (3.0)	9.8 (9.7)	22.4 (22.2)	920, 898	1455s, 1035m	1556s	23 256 (3.08)
1h $[MoO_2L^1(dmf)]^f$	33.7 (33.9)	3.6 (3.5)	10.0 (9.9)	22.7 (22.5)	927, 900	1655s, ^g 685m ^h	1558m	23 474 (3.04)
1i $[MoO_2L^1(Me_2SO)]^f$	30.9 (30.7)	3.2 (3.3)	6.7 (6.5)	22.5 (22.3)	920, 895	1015s ⁱ	1556m	23 640 (3.10)
$1j [MoO_2L^2(py)]$	40.8 (40.4)	3.4 (3.4)	9.5 (9.4)	20.9 (21.5)	928, 902	1450m, 1040m	1557s	23 410 (3.02)
$lk [MoO_2L^3(py)]$	36.4 (36.1)	2.6 (2.6)	8.8 (9.0)	20.2 (20.6)	930, 902	1450m, 1045m	1556m	23 310 (3.08)
11 $[MoO_2L^4(py)]$	33.1 (32.9)	2.5 (2.4)	8.0 (8.2)	18.9 (18.8)	925, 900	1450m, 1050m	1555 (sh)	23 310 (3.09)
2a $[MoO(L^1)]$	32.2 (32.1)	2.3 (2.4)	8.4 (8.3)	28.4 (28.5)	965		1554s	21 550 (3.43), 14 390 (3.00)
$2b \left[MoO(L^2)\right]$	34.7 (34.3)	2.9 (2.9)	7.8 (7.8)	27.1 (27.4)	970		1558m	21 550 (3.45), 14 285 (2.08)
2c $[MoO(L^3)]$	29.5 (29.2)	1.9 (1.9)	7.7 (7.6)	25.6 (25.9)	972	_	1558m	21 645 (3.44), 14 470 (2.05)
$2d [MoO(L^4)]$	26.3 (26.1)	1.7 (1.7)	6.8 (6.8)	23.0 (23.2)	970		1557m	21 645 (3.44), 14 325 (2.03)
$2e [MoOL^1(dmf)]^j$	35.3 (35.2)	3.6 (3.7)	10.4 (10.2)	23.1 (23.4)	970	1655s,# 685m*	1556m	k
$2f [MoOL^2(py)]$	40.6 (40.5)	3.0 (3.1)	10.3 (10.1)	23.4 (23.1)	975	1455s, 1040m	1556m	21 275 (3.73), 14 535 (1.97)
$2g [MoOL^3(dmf)]$	29.7 (29.5)	3.0 (2.9)	8.7 (8.6)	19.2 (19.0)	962	1650s,ª 680m h	1554m	k
2ĥ [MoOL⁴(py)]	34.2 (34.0)	2.4 (2.4)	8.6 (8.5)	19.1 (19.4)	960	1450m, 1010m	1556m	21 460 (3.73), 14 535 (1.97)

^{*a*} Calculated values given in parentheses. ^{*b*} As KBr disc. ^{*c*} In dmf; intraligand transitions are not listed. ^{*d*} All the absorptions are very strong. ^{*e*} Strong and broad band. ^{*f*} Thermal analysis data (temperature range; see text). ^{*g*} Amide I band. ^{*h*} δ (NCO). ^{*i*} ν (SO). ^{*j*} Molecular weight in MeCN-dmf: 412 (409). * Same as 2a and 2c, respectively.

Table 2 Selected cyclic voltammetric,^a proton NMR^b and FT-IR^c data for oxomolybdenum(vi) complexes

Complex	$E_{\rm pc}/{\rm V}(n^d)$	δ(CH=N) ^e	$\Delta \delta^{f}$	v(Mo=O)/ cm ⁻¹
1a	$-0.86(0.97)^{g}$	8.66	0.32	928.8, 904.2
1b	- 0.90	8.61	0.30	933.3, 902.9
lc	-0.84	8.64	0.34	930.0, 905.4
1d	-0.83	8.66	0.36	929.0, 905.4
le	-0.76	8.74	0.18	928.9, 909.0

^a Solvent dmf, solute concentration $\approx 10^{-3}$ mol dm⁻³, reference electrode SCE. ^b Proton NMR in CDCl₃-(CD₃)₂SO (10:1) at room temperature. ^c In dmf solution. ^d n = Q/Q' where Q' is the calculated coulomb count for a one-electron transfer and Q is the observed value after exhaustive electrolysis of 0.001 mol of solute. ^e Signal for the free ligand: H₂L¹, 8.34; H₂L², 8.31; H₂L³, 8.30; H₂L⁴, 8.30; H₂L⁵, 8.56. ^f $\Delta \delta = (\delta_{complex} - \delta_{ligand})$. ^g Coulometry was carried out at -1.06 V.



R = H, Me, CI, Br, NO₂ or 5,6-C₄H₄

vibration appears. This indicates that MoO_2 complexes of L'^{2-1} and L²⁻ ligands are not strictly isostructural. Also, when the complexes [(MoO₂L)_n] 1a-1f react with donor solvents D, such as pyridine (py), dmf or dimethyl sulfoxide (Me₂SO), to form monomeric complexes [MoO₂L(D)] 1g-11 the v(Mo····O \rightarrow Mo) vibration (ca. 805 cm⁻¹) disappears (Table 1), indicating a discrete monomeric structure D for the complexes. As expected,¹⁶ each of the complexes **1a-11** shows only one

ligand-to-metal charge-transfer (l.m.c.t.) transition (see Table 1) in its electronic spectrum.

(b) Oxomolybdenum(IV) Complexes.—The oxomolybdenum-(IV) species, [MoO(L)] (L = L¹-L⁴, 2a-2d) or [MoOL(D)](L = L¹, D = dmf 2e or py 2f; L = L⁴, D = dmf 2g or py 2h) are formed when the appropriate [MoO₂L] complex reacts with PPh₃ in CH₂Cl₂, MeOH or MeCN or in the respective donor solvents (py or dmf), whereby the MoO_2^{2+} unit transfers one oxygen atom to PPh₃ converting it to OPPh₃. As such, 2a-2d are polymeric^{16,29} but show only one v(Mo=O) vibration (Table 1) eliminating polymerisation via oxo bridging. Polymerisation involving phenoxide oxygen as bridging ligand is also not supported ¹⁶ by IR data (Table 1). Complexes 2a-2h are diamagnetic and non-electrolytes in dmf. The donor-coordinated monomers, 2e-2h, also show only a single v(Mo=O) vibration (Table 1). Molecular weight data on 2f (Table 1), further suggests that the species are monomeric and hence five-co-ordinate. This idea is nicely supported by thermoanalytical data. Mass-loss values calculated from the thermogravimetric analysis (TGA) curves of complexes 2e and 2f (Fig. 1) correspond to the loss of one dmf (2e) or py (2f). Interestingly, the differential thermal analysis (DTA) curves (Fig. 1) show that after the endothermic loss of the donor ligand, another exotherm appears at 180 (2e) or 200 °C (2f), which is not accompanied by any mass loss in the TGA curve. This can be attributed to phase transitions arising from a structural change in 2e and 2f, and by inference, also of 2g and 2h, from discrete five-co-ordinate species to polymeric entities. Assuming meridional disposition of the ONS ligand, as in the dioxomolybdenum(vi) complexes, a square-pyramidal geometry with a vacant co-ordination site trans to the Mo=O bond is a reasonable likelihood for the structures of 2e-2h.

The electronic spectra of the [MoO(L)] complexes 2a-2d in degassed dmf solution show two bands and are essentially the same as those of the dmf adducts (2e and 2g) (Table 1). The bands appear at ≈ 462 and ≈ 690 nm (Table 1), very much as found for the H₂L' complexes.¹⁶

(c) Electrochemistry.---The cyclic voltammetry of the isolated dioxomolybdenum(vi) and oxomolybdenum(iv) complexes has







Fig. 2 Cyclic voltammograms for $\approx 10^{-3}$ mol dm⁻³ solutions of the [MoO₂L] complexes in dmf (scan rate 50 mV s⁻¹): R = H (----), Me (----), Cl (····), Br (--··-) or NO₂ (--·--)

been studied using a platinum working electrode and NEt₄ClO₄ as supporting electrolyte in dmf solvent. Complexes **1a-1e** (Fig. 2) exhibit irreversible single-electron (exhaustive coulometry) reduction waves in their cyclic voltammograms at the potentials given in Table 2. No detectable oxidation wave was seen during the reverse anodic scans. This type of behaviour was also noted earlier for other dioxomolybdenum(vI) complexes.^{6,16} The reduction potentials (E_{pc}) are sensitive to the nature of the 5-R substituents on the salicyl phenyl ring. As the substituents become more electron withdrawing (Me < H < Cl < Br < NO₂) the shift of E_{pc} of the Mo^{VI} \rightarrow Mo^V couple becomes more anodic indicating easier reducibility of the respective molybdenum(vI) centres.

The cyclic voltammograms of the [MoOL¹(dmf)] complexes in dmf–NEt₄ClO₄ (0.1 mol dm⁻³) exhibit an irreversible oxidative response at +0.15 V (vs. SCE) showing only a feeble reductive response in the reverse sweep at +0.05 V. On the negative side of the SCE, a cathodic scan shows a reduction wave at -0.97 V (vs. SCE) and the reverse sweep does not show any anodic counterpart, analogous to the H₂L' complexes.¹⁶ The corresponding H₂L complexes with L¹–L⁶ are unstable under the cyclic voltammetric experimental conditions. However, in the Mo^{IV}–H₂L' complexes, the values of E_{pa} move in the cathodic direction as indicated by the order R = Me < H < Cl \approx Br (Table 3), parallelling the electronic effect transmitted by the substituents. Similarly, values of E_{pc} also shift in the anodic direction following the substituent order as indicated above.^{6b} J. CHEM. SOC. DALTON TRANS. 1993

Complex	E_{pc}/V	$E_{\mathrm{pa}}/\mathrm{V}$
$[Mo(L^1)]$	-0.97	+0.15
$[MoO(L^{1'})]^{c,d}$	- 0.94	+0.42
$[MoO(L^{2'})]^d$	-0.98	+0.40
$[MoO(L^{3'})]^d$	-0.82	+0.45
$[MoO(L^{4'})]^d$	-0.83	+ 0.46

^{*a*} Solvent dmf. ^{*b*} Ligands $L^{1'}-L^{4'}$ correspond to $L^{1}-L^{4}$ but with $X = CH_2Ph$ instead of Me. ^{*c*} Data from ref. 16. ^{*d*} For **1a**-1d see Experimental section.



Fig. 3 Spectral changes in the reaction of $[MoO_2L^1]$ (2 × 10⁻⁴ mol dm⁻³) and PPh₃ (6 × 10⁻⁴ mol dm⁻³) in CH₂Cl₂ solution at 25 °C



Fig. 4 Spectral changes in the reaction of $[MoO_2L^1]$ (2 × 10⁻⁴ mol dm⁻³)

(d) Kinetics.—(i) Oxo transfer from Mo^{VI} to substrate. All the molybdenum(VI) complexes undergo oxygen-atom transfer reactions to the substrate, PPh₃, as shown in equations (2) and (3) forming $[Mo^{IV}O(L)]$ or $[Mo^{IV}OL(D)]$ as the products.

$$[MoO_2L^1] + PPh_3 \xrightarrow{CH_2Cl_2} [MoO(L^1)] + OPPh_3 \quad (2)$$

 $[MoO_2L(dmf)] (L = L^1 - L^4) + PPh_3 \xrightarrow{dmf}_{k_3}$ $[MoOL(dmf)] + OPPh_3 \quad (3)$

These reactions (PPh₃ in excess) have been examined spectrophotometrically, as shown in Figs. 3 and 4. Plots of $\ln(A_{\infty} - A_t)$ against time (t) are linear (see Experimental



Scheme 1 (a) In CH₂Cl₂. (b) In dmf

 Table 4
 Kinetic data for oxo-transfer reactions of molybdenum-(vi) and -(iv) complexes

Complex	Substrate	Solvent	<i>T/</i> °C	$10^3 \times \text{Rate constant}^a$	
1a	PPh ₃	CH ₂ Cl ₂	25	$(5.60 \pm 0.36) \times 10^3 (k_2)$	
	PPh ₃	dmf	30	$14.87 \pm 0.24 (k_3)$	
1b	PPh ₃	dmf	30	$12.62 \pm 0.20 (k_3)$	
lc	PPh ₃	dmf	30	$18.08 \pm 0.32 (k_3)$	
1d	PPh ₃	dmf	30	$18.59 \pm 0.87 (k_3)$	
2a	Me ₂ SO	CH ₂ Cl ₂	20	$2.58 \pm 0.11^{b} (k_{-1})$	
2e	Me ₂ SO	dmf	20	$6.15 \pm 0.14^{\circ} (k_{-1})$	
$k_2/dm^3 mol^{-1} s^{-1}, k_3/dm^3 mol^{-1} s^{-1}, k_{-1}/s^{-1}.$ $K = 16.5.$ $K = 79.0.$					

section) showing that the reaction is first-order in molybdenum complex. The dependence of k_{obs} on [PPh₃] is linear with a nearly zero intercept. As for other oxygen atom-transfer reactions from molybdenum(v1) complexes to PPh₃, the reactions follow the rate law, $-d[MoO_2L^1]/dt = k_2[MoO_2L^1][PPh_3]$ in CH₂Cl₂ or $-d[MoO_2L(dmf)]/dt = k_3[MoO_2L(dmf)][PPh_3]$ (L = L¹-L⁴) in dmf. The oxo transfer is more facile for 1a, studied in CH₂Cl₂ solvent in which the polymeric structure is retained, in comparison to that of 1h (Table 4). This is due to the fact that compared to the terminal MoO₂ oxygens in the discrete mononuclear complexes, the Mo...O-Mo oxygen present in the oligomers is more labile and hence reacts faster with the substrate, leading to a higher rate constant value. This phenomenon in oxo-transfer reactions is reported here for the first time.

(ii) Oxygen-atom transfer from substrate. In order to assess the oxo-transfer behaviour of [Mo^{IV}O(ONS)] in CH₂Cl₂ and in dmf, kinetic studies were carried out under pseudo-first-order conditions using a concentration variation of Me₂SO (50–1000 molar excess). The oxygen-atom transfer from substrate to the Mo^{IV}O core had previously been investigated in dmf for an NS₂ ligand.^{20a} The present study has focussed attention on the modification of the reactivity of the complex molecule with a variation of solvents (donor vs. non-donor). Complexes [MoO(ONS)] and [MoO(ONS)(dmf)] react with Me₂SO in two steps (Scheme 1) to yield [MoO₂(ONS)] and Me₂S.

Spectrophotometric examination of the above reactions shows a clean isosbestic point at 358 nm (Fig. 5) in CH₂Cl₂ and at 382 nm (Fig. 6) in dmf. The final spectra are identical to



Fig. 5 Spectral changes in the reaction of $[MoO(L^1)]$ (2 × 10⁻⁴ mol dm⁻³) and Me₂SO (1.4 × 10⁻² mol dm⁻³) in CH₂Cl₂ solution at 20 °C



Fig. 6 Spectral changes in the reaction of [MoOL¹(dmf)] (2×10^{-4} mol dm⁻³) and Me₂SO (2.8×10^{-2} mol dm⁻³) in dmf solution at 20 °C



Fig. 7 Dependence of the rate of reaction of [MoOL¹(dmf)] in dmf solution at 20 °C with concentration of Me₂SO

those of the isolated $[MoO_2L(Me_2SO)]$ complexes. The plots of $\ln(A_t - A_{\infty})$ against t are linear up to four half-lives of the reactions, indicating a first-order dependence of rate on the molybdenum(iv) complexes. Again, a plot of the observed rates against $[Me_2SO]$ shows that at sufficiently high concentrations the rates (k_{obs}) , become virtually independent of $[Me_2SO]$, *i.e.*, substrate saturation kinetics is obtained (Fig. 7).

As is apparent from Scheme 1, in dmf solution, an equilibrium exists between the dmf- and sulfoxide-ligated molybdenum(IV) complexes. On the other hand, in CH₂Cl₂ medium, an equilibrium exists between $[{MoO(L)}_n]$ and $n[MoOL(Me_2SO)]$. As for other oxygen-atom transfer reactions from Me₂SO to molybdenum(IV) complexes in dmf, the rate law ^{20a} is given by equation (4) (Y = dmf + Me₂SO).

$$\frac{d[MoOL(Y)]}{dt} = -k_{-1}[MoOL(Me_2SO)] = -k_{-1}\left\{\frac{K[Me_2SO]}{K[Me_2SO] + [dmf]}\right\}[MoOL(Y)] \quad (4)$$

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Fig. 8 Plot of (a) v_{asym} (Mo=O) vs. Hammett σ_p ; (b) l.m.c.t. vs. σ_p ; (c) l.m.c.t. vs. E_{pc} ; (d) E_{pc} vs. σ_p ; (e) k_3 vs. E_{pc} ; (f) k_3 vs. v_{asym} (Mo=O); (g) $\Delta\delta$ vs. σ_p [note that the abscissa is not extended up to σ_p (NO₂) (see text)]; for [MoO₂L(dmf)] [L = L¹ (R = H), L² (R = Me), L³ (R = Cl), L⁴ (R = Br) or L⁵ (R = NO₂)]

In CH_2Cl_2 the reaction follows a simpler rate law ³⁰ given by equation (5).

$$\frac{\mathrm{d}[\mathrm{MoOL}(\mathrm{Me}_{2}\mathrm{SO})]}{\mathrm{d}t} = -k_{-1}[\mathrm{MoO}(\mathrm{L})][\mathrm{Me}_{2}\mathrm{SO}] = \frac{-k_{-1}K[\mathrm{Me}_{2}\mathrm{SO}]}{1+K[\mathrm{Me}_{2}\mathrm{SO}]}$$
(5)

Parameter k_{-1} denotes the rate constant accompanying the intramolecular oxo transfer from co-ordinated substrate, Me₂SO, to the [MoO(L)] moiety as shown in Scheme 1, and values of K are the respective equilibrium constants of the reactions. The values of k_{-1} and K were evaluated from plots of $1/k_{obs}$ vs. 1/[Me₂SO], and the same values are obtained from the reciprocals of the intercepts and slopes,^{20a} respectively (Table 4) of the straight lines thus obtained. The equilibrium constants for **2a** and **2e**, are two orders of magnitude lower than that for the reaction of Me₂SO with an [MoO(NS₂)(dmf)] complex,^{20a} but the rates of the oxo transfer are almost the same. Thus, the activated species in our ONS ligated complex is thermodynamically less stable than that of the NS₂ ligated system.^{20a} Again, comparing the K values of the polymeric and monomeric species described here, it is found that the values are only slightly smaller for E than for F. This may be apparent from Scheme 1, which implies that E is derived by the breaking of polymers to monomers assisted by Me₂SO co-ordination, while F is generated *via* a nucleophilic substitution reaction.

(e) Influence of Ligand Substituents on the Structural and Electronic Aspects of the Complexes.—The unco-ordinated S-Me group present in the H_2L ligands renders the molybdenum centre in the MoO₂ complexes too electron rich to exhibit discriminatory properties in forming both monomeric or polymeric complexes for the range of 5-R substituents [H (0.00),* Me(-0.17), Cl, Br(+0.22), NO₂(+0.77)] or 5,6-C₄H₄ studied here. As a result, only the polymers, [(MoO₂L)_n] 1a-1f,

* Figures in parentheses indicate the respective Hammett σ_p parameter.

have been obtained. This electron richness makes $[MoO_2L]$ species less easily reducible than the corresponding complexes $[MoO_2L']$ (compare Table 2 with Table 3 of ref. 16). Interestingly, this behaviour is also reflected in the oxomolybdenum(IV) complexes. The E_{pa} value for the Mo^{IV}-Mo^V couple for [MoOL¹(dmf)] (Table 3) is found to be more cathodic than that of the corresponding [MoOL'(dmf)] (R = H) species (Table 3). Similarly, the E_{pc} value for the Mo^{IV}-Mo^{III} couple in the oxomolybdenum(IV) also becomes more cathodic for X = Me(L) compared to $X = CH_2Ph(L')$ (Table 3).

However, despite the levelling effect (coarse tuning) imposed on the R substituents by the electron-donating methyl substituent towards the position of the monomer-polymer equilibrium, the nature of the R substituents still controls (by fine tuning) various spectroscopic, electrochemical and reactivity parameters of the respective polymers or $[MoO_2L(D)]$ complexes derived from these. These parameters include the position of the $v(Mo \rightarrow Mo)$ and v(Mo = O)vibrations, the band position of the $S_{p\pi} \rightarrow Mo_{d\pi}$ l.m.c.t. transition, the Mo^{v_1} -Mo^v reduction potential for the MoO_2 complexes, the oxidation (Mo^{lv} -Mo^v) and reduction (Mo^{lv} -Moⁱⁿ) potentials of the [MoO(L)] complexes, the rate constant of the oxo-transfer reaction represented by the forward reaction in equation (1) and the position of azomethine proton resonances in appropriate complexes. Regarding the position of the $v(Mo \rightarrow Mo)$ vibration, the gross substituent dependence is $NO_2 \approx Br \approx Cl < H \approx Me < 5,6-C_4H_4$ (Table 1) indicating that electron-withdrawing substituents destabilize the oxygen bridges. Interestingly, the v_{asym} (Mo=O) vibration (Table 2), and the energy (in cm⁻¹) of the l.m.c.t. electronic transition, of $[MoO_2L(dmf)]$ (L = L¹-L⁵) (Table 1) in dmf solution, show a linear correlation⁹ with the Hammett σ_p values of the R substituents (Fig. 8). Similarly for the $[MoO_2L^1(D)]$ (D = py 1g, dmf 1h or Me₂SO 1i) complexes, the v_{asym} (Mo=O) vibration in the solid state increases in the order D = Me₂SO < py < dmf. The temperature loss of donor molecules D (TGA data) for the above complexes in the solid state also decreases in the same order indicating that strong donor binding to molybdenum reduces $O_{p\pi} \rightarrow Mo_{d\pi}$ overlap in the MoO_2^{2+} moiety.

An investigation into the redox chemistry of the $[MoO_2L]$ complexes suggests that a linear dependence exists between values of E_{pc} (Table 2) and the Hammett σ_p parameter of the R substituents^{6b} of the respective ligands, and also between E_{pc} and positions of the l.m.c.t. transitions (Fig. 8). This demonstrates the sensitivity of the molybdenum electron density (and hence the optical electronegativity of the metal) on the Hammett σ_p parameter of the R substituent.

Also, the extent of the downfield shift ($\Delta\delta$) of the azomethine proton signal of H₂L upon complexation is less in the present complexes (X = Me) than the corresponding values observed in complexes with L' (X = CH_2Ph).¹⁶ For the present series of complexes $\Delta \delta$ varies linearly with the Hammett σ_p values of the R substituent (except for $R = NO_2$, see Fig. 8).

Finally, the observed oxo-transfer rate constant (k_3) from [MoO₂L(dmf)] to PPh₃ shows a linear relationship with the values of E_{pc} for the Mo^{VI}-Mo^V couples (Fig. 8). This indicates that the electronic effects that cause systematic changes in E_{pc} values also modify the electrophilicity of the oxide ligands which is manifested in their acceptor properties towards phosphorus donors

All these correlations indicate the prominence of ligand control (by variation of R) towards metal reactivity in these systems.

Conclusion

The nature of the neighbouring group SX attached to a >C-SH functionality imparts a profound electronic effect at the metal centre depending on the nature of X. When X is an ambivalent substituent, e.g. CH₂Ph, the mono- or oligonuclearity of the resulting ONS donor complex depends on the

nature of the R substituents attached to the salicyl phenyl groups. However when X is an electron-repelling group such as Me, the complex $[MoO_2(L)]$ is always an oligomer irrespective of the nature of the R substituents. For the oligomeric complexes [(MoO₂L)_n], or in the donor-co-ordinated monomers [MoO₂L(D)] obtained from the oligomers, the R substituent almost quantitatively controls various spectroscopic, electrochemical and reactivity parameters of the complexes. For the MoO₂ complexes the rate of oxo transfer to PPh₃ substrate is much higher when the oxygen atom involved occurs as a bridge between two molybdenum atoms, $Mo - O \rightarrow Mo, \{[(MoO_2L)_n]\}$ rather than as a terminal oxygen of the MoO_2^{2+} group {[MoO_2L(D)]}.

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