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## Adducts of Molybdenum(v) Trichloride Sulphide

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The compound MoCl<sub>3</sub>S reacts with methyl cyanide to give MoCl<sub>3</sub>S,MeCN and MoCl<sub>3</sub>S,2MeCN, which have been characterised as containing six-co-ordinate molybdenum with bridging Mo-S-Mo and terminal Mo-S bonds respectively. The complex MoCl<sub>3</sub>S,2MeCN reacts with other ligands; 1,4-dioxan and 1,2-bis(methoxy)ethane give adducts MoCl<sub>3</sub>S,2diox and MoCl<sub>3</sub>S,bme, while pyridine gives the salt [Hpy][MoCl<sub>4</sub>S,py] and 1,10phenanthroline reduces molybdenum to the quadrivalent state in forming MoCl<sub>2</sub>S, phen. Direct reaction of MoCl<sub>3</sub>S with py gives MoCl<sub>3</sub>S,3py (one py being in the lattice). The salts WČl<sub>3</sub>S and MoCl<sub>3</sub>S did not react with any of the ligands tried even after prolonged reaction periods.

THE chemistry of molybdenum(v) trichloride oxide and tribromide oxide has been studied in some detail, 1-5 but reported work with the corresponding trihalide sulphide and selenide has been limited to their preparation.6,7 Some adducts of WCl<sub>3</sub>S have been prepared 7 from WClaS by reduction with various ligands. We now report attempts to prepare complexes of MoCl<sub>2</sub>S, MoCl<sub>3</sub>Se, and WCl<sub>3</sub>S by direct reaction of the trihalide sulphides or selenides with ligands.

## EXPERIMENTAL

As the compounds are very moisture sensitive, the preparative and handling procedures were carried out by means of conventional vacuum-line or dry-box methods. All ligands and solvents were rigorously dried and distilled on the vacuum line before use. The compounds MoCl<sub>3</sub>S, MoCl<sub>3</sub>Se, and WCl<sub>3</sub>S were prepared by heating the appropriate pentahalide with either Sb<sub>2</sub>S<sub>3</sub> or Sb<sub>2</sub>Se<sub>3</sub>.7

Conductivity measurements were carried out in cells specially adapted to eliminate air and moisture; molecular weights were determined by the elevation of b.p. method; oxidation states were determined by titration with cerium-(IV) sulphate.

Preparations.—Analytical data are given in the Table. (i) Molybdenum(v) trichloride sulphide-methyl cyanide (1/1) and (1/2). The compound MoCl<sub>3</sub>S (2 g) and MeCN (50 cm<sup>3</sup>) were sealed in an evacuated ampoule, and the mixture heated gently under reflux for 48 h. A green solution formed together with a green precipitate and the latter

- † 1 B.M. =  $0.927 \times 10^{-23}$  A m<sup>-2</sup>.
- <sup>1</sup> R. A. Walton, *Progr. Inorg. Chem.*, 1972, **16**, 1. <sup>2</sup> D. L. Kepert, 'The Early Transition Metals,' Academic Press, London, 1972.

  <sup>3</sup> K. Feenan and G. W. A. Fowles, Inorg. Chem., 1965, 4, 310.

- (MoCl<sub>3</sub>S,MeCN) was isolated by filtration in vacuo. Evaporation of the solution gave a dark green powder of composition MoCl<sub>3</sub>S,2MeCN. The magnetic moment (in B.M.) of MoCl<sub>3</sub>S,MeCN was determined over a temperature range:  $\mu = 1.07$  (298); 1.04 (273); 1.03 (253); 1.03 (233); 1.00 (193); 0.99 (173); 0.98 (153); 0.97 (133); and 0.94113 K).†
- (ii) Molybdenum(v) sulphide-1,4-dioxantrichloride(1/2) and -1, 2-bis(methoxy)ethane (1/1). The adduct MoCl<sub>2</sub>S,2MeCN (2 g) was heated under reflux in a sealed ampoule with neat ligand (50 cm<sup>3</sup>; diox or bme) for 16 h. The green solids obtained on evaporation of the excess of ligand still contained some McCN (i.r. spectral evidence) so they were heated for a further period with two further successive quantities of fresh ligand.
- (iii) Pyridinium tetrachlorothiomolybdate(v)-pyridine (1/1). The adduct MoCl<sub>3</sub>S,2MeCN (2 g) and pyridine (py) (1:4 mol ratio) were heated gently in a sealed ampoule with benzene (50 cm<sup>3</sup>). The green solution formed initially gradually turned brown and deposited a green solid, [Hpy]-[MoCl<sub>4</sub>S,py], which was isolated by filtration, washed with benzene, and pumped free from solvent. An intractable tar remained when the brown solution was evaporated.
- (iv) Molybdenum(v) trichloride sulphide-pyridine (1/3). The compound MoCl<sub>3</sub>S (2 g) was heated gently under reflux for 24 h with py (50 cm3) in a sealed ampoule, and the green solution evaporated to leave MoCl<sub>3</sub>S,3py.
- (v) Molybdenum(IV) dichloride sulphide-1,10-phenanthro-
- 4 P. C. Crouch, G. W. A. Fowles, P. R. Marshall, and R. A.
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line (1/1). The adduct MoCl<sub>3</sub>S,2MeCN (2 g) was allowed to react with 1,10-phenanthroline (phen) (1:2 mol ratio) in dichloromethane solution at room temperature. The green solution which formed initially gradually turned red, and red crystals of MoCl<sub>2</sub>S, phen were formed as the solvent was removed by the pump.

(vi) Other attempted preparations. (a) Only nonstoicheiometric materials were obtained from the reaction of MoCl<sub>3</sub>S,2MeCN with PPh<sub>3</sub>, tetrahydrofuran (thf), Et<sub>2</sub>O, and MeSCH<sub>2</sub>CH<sub>2</sub>SMe. (b) Non-stoicheiometric materials also resulted when MoCl<sub>3</sub>S,MeCN was heated under reflux with pv, diox, thf, and bme. (c) The compounds MoCl<sub>3</sub>Se

MoCl<sub>2</sub>S Complexes were prepared by direct reaction with MeCN and py, and the methyl cyanide complex was used as a starting material for other complexes. Thus methyl cyanide can be replaced by diox, 1,2-bis-(methoxy)ethane, py, and phen to give well defined complexes, although with triphenylphosphine, diethyl ether, thf, and 1,2-bis(methylthio)ethane the products were non-stoicheiometric. The Table lists the products, their analytical data, and their i.r. spectra in the 200— 400 cm<sup>-1</sup> region, together with the peak at ca. 480 cm<sup>-1</sup> tentatively assigned to a v(Mo=S) stretching mode.4

Analytical and i.r. data (400—200 cm<sup>-1</sup>) of the complexes prepared from MoCl<sub>3</sub>S

	Analyses/(%)																		Diffuse-
	Found						Calc.						/ :(NIC) /			Oxida	reflectance		
Complex	Mo	S	Cl	С	Н	N	Mo	S	CI	Н	Н	N	Colour	$_{ m B.M.}^{\mu/}$	/ (Mo=S) ( cm <sup>-1</sup>	M a	tion state	I.r. spectra b	spectra $(10^{-3} \bar{\nu}/\text{cm}^{-1})$
MoCl <sub>3</sub> S,MeCN	35.0	12.3	38.2	9.0	1.0	5.1	34.9	11-6	38-7	8-7	1.1	5.1	Green	1.07			5.2	340s, 285m, 270m	37·2(sh), 30·0, 21·8, 10·5, 8·0(sh)
MoCl <sub>3</sub> S,2McCN	29-9	9.7	33.8	13.2	2.0	6.7	30.3	10.1	33.7	15.2	1.9	8.8	Green	1.85	480	330	4-9	320s, br, 285(sh), 245w	37.0, 24.0, 13.0
MoCl <sub>3</sub> S,2diox	23.7	7.4	26.4	$22 \cdot 1$	3.7		23.4	7.8	26.0	23.4	3.9		Green	1.71	475	400	5.0	<i>320</i> s, br, 285s	37.0, 34.2, 29.0, 22.2, 15.6
MoCl <sub>3</sub> S,bme	29.0	8.9	31.9	13.0	2.8		29.6	9-9	32.8	14.8	3.1		Green	1.68	470		5.2	315,s br	37·5, 34·2, 30·0, 18·0
MoCl <sub>3</sub> S,3py	19-9	6.5	22.0	37.4	3.2	8.6	20.4	6.8	22.6	38.2	$3 \cdot 2$	8.9	Green	2.01	475	280 c	4.8	320s, 300s, 280(sh)	37.6, 31.8, 26.2, 22.0, 16.0
[Hpy][MoCl <sub>4</sub> S,py]	21.8	7-0	32-8	29.()	$2 \cdot 6$	7.3	22.4	7.5	33.1	28.0	2.4	6.5	Green	1.74	490	d	5.1	340s, 322s, 255	36·0, 28·6, 36·4, 21·0, 7·0
MoCl <sub>2</sub> S,phen	25.3	6.6	18.4	39.6	$2 \cdot 7$	8.3	25.4	8-5	18.7	38.0	2.1	7.4	Red	1.99	460		3.8	<i>335</i> s, 330s, <i>295</i> s, 235w	44.6, 37.0, 25.6, 19.0, 14.5

a Measured by elevation of b.p. in benzene solution. b Italicised bands suggest probable Mo-Cl modes. c Dissociation to MoCl<sub>3</sub>S,py and py. d 1:1 Electrolyte in McCN.

and WCl<sub>2</sub>S were heated under reflux with all the ligands referred to under reactions of MoCl<sub>3</sub>S,2MeCN, but despite prolonged reflux periods no reaction took place; analysis showed the products to be unchanged MoCl<sub>3</sub>Se and WCl<sub>3</sub>S.

## RESULTS AND DISCUSSION

The trichloride and tribromide oxides of molybdenum and tungsten have the NbCl<sub>3</sub>O structure with six-coordinate metal atoms bridged by both chlorine and oxygen atoms, 8,9 although there is a modification of

MoCl<sub>3</sub>O which contains terminal oxygen atoms. The sulphur and selenium analogues under discussion also have the NbCl<sub>3</sub>O structure, (I).<sup>10</sup> Hence it is interesting to note that while the trihalide oxides react directly with a number of ligands, often under relatively mild conditions,4 the trihalide sulphide and selenide are appreciably less reactive. The compounds MoCl<sub>3</sub>Se and WCl<sub>3</sub>S, for instance, did not react with any of the ligands tried, even after prolonged periods under reflux, and only MeCN and py reacted with MoCl<sub>3</sub>S. This reduced activity suggests that the M-S-M bridging is more resistant to attack by donor ligands than the comparable M-O-M bridges.

<sup>8</sup> P. C. Crouch, G. W. A. Fowles, J. L. Frost, P. R. Marshall, and R. A. Walton, *J. Chem. Soc.* (4), 1968, 1061.

(i) MoCl<sub>3</sub>S,2MeCN and MoCl<sub>3</sub>S,MeCN.—The 1:2 adduct is monomeric in benzene solution and behaves as a non-electrolyte in methyl cyanide. The roomtemperature magnetic moment (1.85 B.M.) rules out any interaction through neighbouring molybdenum atoms, so the structure would appear to be a simple six-coordinate monomer. This requires a terminal Mo=S bond and the i.r. spectrum shows a peak at 480 cm<sup>-1</sup> that can be tentatively assigned to  $\nu(Mo=S)$ . There is no such peak in the spectrum of the 1:1 adduct, on the other hand, and this implies a Mo-S-Mo bridging system rather than a terminal Mo=S bond. Insolubility of the adduct in the usual solvents ruled out measurements of molecular weight and conductance but is consistent with a polymeric structure, (II). The low room-temperature magnetic moment (1.07 B.M.) and the other magnetic data (see Experimental section) indicate some Mo-Mo interaction and this could possibly be by way of the bridging sulphur atom.

In the reaction of WCl<sub>2</sub>O with pv <sup>4</sup> two analogous complexes (WCl<sub>3</sub>O,2py and WCl<sub>3</sub>O,py) were also obtained, the 1:2 adduct being a six-co-ordinate monomer and the 1:1 adduct an oxygen-bridged polymer that shows magnetic interaction.4

(ii) MoCl<sub>3</sub>S,bme and MoCl<sub>3</sub>S,2diox.—The i.r. spectra of both complexes contain bands associated with  $\nu(\text{Mo=S})$  and that of the diox complex shows both modified and unmodified COC stretching frequencies,

<sup>&</sup>lt;sup>9</sup> M. G. B. Drew and I. B. Tomkins, Acta Cryst., 1970, **B26**,

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showing the ligand to be unidentate.4,11 The diox complex is monomeric in benzene solution and is clearly six-co-ordinate. The bme complex is also likely to be a monomer with a chelating ligand molecule. Thus the room-temperature magnetic moment of 1.68 B.M. rules out Mo-Mo interactions, and the i.r. spectrum shows the ligand to be in the gauche chelating form.<sup>12</sup>

(iii) MoCl<sub>3</sub>S,3py and [Hpy][MoCl<sub>4</sub>S,py].—The direct reaction of MoCl<sub>3</sub>S with py gives MoCl<sub>3</sub>S,3py as product, whereas MoCl<sub>3</sub>S,2MeCN reacts with py to give a product with analysis corresponding to the overall composition MoCl<sub>4</sub>S,2py. The complex MoCl<sub>3</sub>S,3py appears to be six-co-ordinate MoCl<sub>3</sub>S,2py with the third molecule of py in the lattice; an analogous situation has been reported for TiCl<sub>3</sub>,4py.<sup>13</sup> The i.r. spectrum shows the presence of both co-ordinated and unco-ordinated py molecules, <sup>14</sup> but no bands attributable to Hpy<sup>+</sup>. When the complex was dissolved in benzene, molecular-weight measurements indicated dissociation, but conductivity measurements on methyl cyanide solutions showed the species to be non-conducting. MoCl<sub>4</sub>S,2py, on the other hand, appears to be a pyridinium salt, [Hpy]-

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1972, 8, 975.

[MoCl<sub>4</sub>S,py], since it behaves as a 1:1 electrolyte in methyl cyanide solution 15 and its i.r. spectrum shows both the pyridinium cation and co-ordinated py to be present. The Hpy+ proton presumably comes from one of the excess of pyridine molecules.16

(iv) MoCl<sub>2</sub>S,phen.—Oxidation-state titrations show molybdenum to be present in the quadrivalent state in this complex; the magnetic moment (1.99 B.M.) is well below the spin-only value but very similar to that of  $[MoBr_4(pdma)_2]$  $\lceil pdma = o-phenylenebis(dimethyl$ arsine)].<sup>17</sup> The complex behaves as a non-electrolyte in methyl cyanide solution, and we suggest that it may contain six-co-ordinate molybdenum through bridging chlorine atoms. Thus the i.r. spectrum shows a Mo=S bond to be present, which rules out Mo-S-Mo bridging, and the complex low-frequency spectrum is consistent with both terminal and bridging Mo-Cl bonds being present; the usual bands of co-ordinated phen are found but there is no evidence for a protonated cation.

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