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THE BEHAVIOUR OF COPPER AND THALLIUM HEXAFLUOROMOLYBDATES(V) IN ACETONITRILE

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Copper (II), copper (II), and thallium (III) hexafluoromolybdates (V), prepared by the oxidation of the metals in acetonitrile with molybdenum hexafluoride (A. Prescott, D.W.A.Sharp, and J.M. Winfield, J. Chem. Soc., Dalton Trans., 1975, 963) have been investigated by cyclic voltametry. Half wave potentials, E. V vs. Ag*/Ag were obtained using a evacuable cell equipped with an external Ag*/Ag electrode, enabling strict anerobic conditions to be maintained. A number of reversible or quasi-reversible electron transfer processes have been observed, enabling comparison with synthetic work to be made. Results for Cu¹ and Cu¹I hexafluoromolybdates (V) are in accord with preparative experience. MoF₆₄ MoV¹/MoVE_{k2} = +1.600V, oxidises Cu metal to Cu¹I in MeCN, and Cu¹I is reduced by Cu° to Cu¹, Cu¹I/Cu¹E_{k2} = +0.750 or +0.710V for Cu¹ and Cu¹I solutes respectively, Cu¹/Cu°E_{k2} = -0.720V not reversible. A wave at E_{k2} = -0.350V is assigned to MoV/MoIV by analogy with Ag¹ hexafluoromolybdate (D.W.A. Sharp, unpublished work). E_{k2} data for I₂ in MeCN, I₂/I₃ = 0.280, I₃-/I = -0.116V, suggest that reduction of MoF₆ by I¹ is not likely, in contrast to the situation in So₂ (A.J. Edwards and R.D. Peacock, Chem. Ind., 1960, 1441). Reduction of MoF₆ by Cu° in MeCN should be feasible, but appears to be very slow. Pure Tu¹II hexafluoromolybdate (V) is obtained from Tu° and MoF₆ only when the mole ratio MoF₆:Tu>5:1. Smaller ratios produce yellow solids in which Mo:Tû is ca. 2:1. Tu²III is a stronger oxidising agent than Cu¹I in MeCN, as oxidation of Cu¹I by Tu²III is rapid and quantitative. However a reversible electron transfer wave assignable to Tu²III/Tu² is not observed in the expected range +1.600 to +0.710V possibly because of solute-electrode interactions.

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REDOX BEHAVIOUR IN HEXAFLUOROMETALI ATE SYSTEMS

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The electrochemical processes $[MF_6]^{n-}$ \longrightarrow $[MF_6]^{-n+1}$ have been studied by cyclic voltammetry and polarography in acetonitrile for a series of transition metals. The observed potentials show correspondence with the number of d electrons and emphasise quantitatively the expected greater oxidising power of second-row elements over third-row elements. There are breaks in the plots of E against the number of d electrons at $d^3 + d^4$ corresponding to the spin pairing in the d^4 state.

Metal	VI/V	V/IV	IV/III	Metal	VI/V	V/IV	IV/III
Nb	-	-1.55	<-3.0	Ta	-	-2.55	<-3.0
Mo	+1.70	-0.35	-2.25	W	+0.70	-1.30	<-3.0
Tc	-	-	-	Re	+2.00	-0.20	<-3.0
_ Ru	>3.0	+0.85	-1.25	e0	+2.90	+0.25	-2.3

(volts vs. Ag⁺/Ag)