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carbamates, it is possible to obtain both a minimum and a maximum in the term-
far reported are in iron(III) complexes.^{2,4} In favorable cases, e.g. many dihydro-
been observed in a number of different thioxanthates. The best example of this so
High-spin \rightleftharpoons low-spin equilibrium equilibria (also called "cross-over") have

INTRODUCTION

chains are more stable than the straight-chain analogues.
central iron atom to form an iron(II) complex. Complexes with branched alkyl
when R is an alkyl group. Some long-chain ligands are capable of reducing the
of the ligand to reduce the central metal atom increases with increasing chain length
the case of the anion, the stability of the thioxanthates decreases and ability
Fe(S₂C-S-R)_n are at least as stable as the corresponding xanthates. Analogously to
kept fairly well if stored frozen in sealed tubes. The iron(III) thioxanthates
In general the complexes are rather unstable at room temperature, but some
adduct.

Pyridine forms green paramagnetic adducts in the solid with the nickel(II)
complexes which lose pyridine if left in the open, while in solution a concentration
dependent equilibrium is set up between the planar thioxanthates and its bis-pyridine
that.

that, though the low spin form predominates for the xanthates and thioxan-
thiates, high-spin \rightleftharpoons low-spin equilibrium shows that the ferric thioxanthates and
ture dependences of the magnetism show that the ferric thioxanthates exhibit a
xanthates and thioxanthates which are brownish red. The pressure and tempera-
In sulfidic dilute solutions, iron(III) thioxanthates are more red than
iron(III), nickel(II), and copper(II) are black and those of cobalt(III) are dark green.
Thioxanthate complexes closely resemble the xanthates and the thioocar-
bamates. Like the corresponding xanthates and thioocarbamates, thioxanthates of

Summary

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AND NICKEL(II)

MAGNETICALLY ANOMALOUS THIO COMPLEXES OF IRON(III)

perature dependence of the susceptibility because the relative proportions of high and low spin species follow a Boltzmann distribution,^{2,7} while the individual susceptibilities (χ) of the high and low spin forms each essentially follow a Curie-Weiss law. At low temperature ($kT \ll E$, where E is the difference in energy between high and low spin forms), the predominant effect on the susceptibility is the Curie-Weiss decrease of the low spin χ with temperature, which outweighs the increase in χ due to a small net conversion to high spin: at higher temperatures ($kT \approx E$) the proportion of high and low spin species is very temperature dependent, and the increase in χ is due to conversion of low to high spin species with increasing temperature outweighing the Curie-Weiss dependences for both the high and low spin forms. At still higher temperatures ($kT \gg E$) the complex is mainly in the high spin form and the Curie-Weiss dependence for the latter predominates, so that χ decreases again with temperature (Fig. 1).

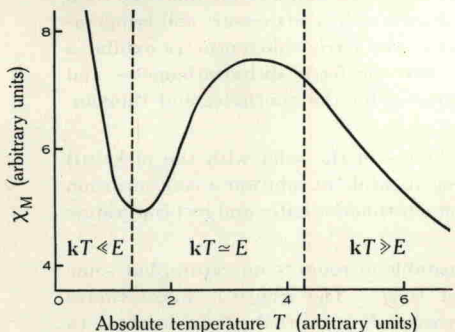


Fig. 1.—Temperature dependence (schematic) of magnetic susceptibility for a compound exhibiting a ${}^2T_1 - {}^6A_1$ equilibrium.

In simple paramagnetic substances, the susceptibility and the moment (μ_{eff}) are independent of pressure.² For complexes which exhibit an electronic equilibrium in solution without any change in structure, pressure shifts the equilibrium in favour of the low spin species, which have the smaller volume, so that χ and μ_{eff} decrease with increasing pressure.^{2,3}

The dependence of an electronic equilibrium in an octahedral d^6 configuration on temperature (T) and pressure (P) may be expressed quantitatively by the equation²

$$\mu_{\text{eff}}^2 = \frac{\frac{3}{4}g^2 + 105C \exp\{-(1+E/\zeta)x\} \exp(-P\Delta V/RT) + 8x^{-1}\{1-\exp(-\frac{3}{2}x)\}}{1 + 2 \exp(-\frac{3}{2}x) + 3C \exp\{-(1+E/\zeta)x\} \exp(-P\Delta V/RT)} \quad (1)$$

where x is ζ/kT , ζ the one-electron spin-orbital coupling constant, g the spectroscopic splitting factor applicable to the 2T_2 state, E the separation of the zero point energies of the 2T_2 state, C ($= Q_a/Q_t$) the ratio of the molecular vibrational partition functions in 6A_1 and 2T_2 , and ΔV the difference in molar volume of the complex in the two states. The factor g should be in the range 1.8–2.0, ζ between 300 and 400 cm⁻¹, while E can vary over any range, depending on the compound.

At atmospheric pressure $\exp(-P\Delta V/RT)$ is sufficiently close to unity to be neglected, but it becomes quite important at higher pressures. Thus, for the iron(III)

⁷ Van Vleck, J. H., "The Theory of Electronic and Magnetic Susceptibilities." p. 182. (Oxford University Press 1932, repr. 1965.)