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SLOW PARAMAGNETIC RELAXATION OF SOME PSEUDO-TETRAHEDRAL FERROUS COMPOUNDS IN ZERO FIELD

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Abstract. Slow (zero field) paramagnetic relaxation is reported for undiluted tetrahedral ferrous monomers for the first time. The Mössbauer spectra of Fe(2,9-di-CH3-phenanthroline)(NCS)2 and Fe(2,9-di-CH3-4,7-di-phenanthroline)(NCS)2 exhibit magnetic hyperfine splitting in the range 20 K to 1.6 K while the corresponding chlorides are rapidly relaxing paramagnets at all temperatures in zero field. Low temperature susceptibility and Mössbauer results indicate that D changes sign from positive to negative from the Cl− to NCS− respectively. The spectra Mössbauer spectra to 1.6 K confirm rapid paramagnetic relaxation for other pseudo-tetrahedral analogues (e.g. Fe(biquinoline)X2, X=Cl−, NCS−). The slow relaxation is then peculiar to the 2,9-di-CH3-phenanthroline system.

Résumé.- La relaxation paramagnétique lente en l'absence de champ magnétique extérieur a été observée dans des composés tétraédriques de fer (II) pour la première fois. Les spectres Mössbauer de Fe(2,9 di-CH3-phenanthroline)(NCS)2 et Fe(2,9 di-CH3-4,7 di-phenanthroline)(NCS)2 montrent une structure hyperfine magnétique dans le domaine de 20 K à 1,6 K. Au contraire, les complexes correspondants de chlorure montrent un taux de relaxation rapide à toutes températures en l'absence de champ magnétique. Des résultats de mesures de susceptibilité magnétique à basse température et de spectroscopie Mössbauer indiquent que D change de signe (de positif en négatif) lorsqu'on passe de Cl− à NCS− respectivement. Les spectres Mössbauer jusqu'à 1,6 K confirment la relaxation paramagnétique rapide pour les autres composés analogues pseudo-tétraédriques (par exemple, Fe(biquinoline)X2, X=Cl−, NCS−). La relaxation lente est particulière au système 2,9-diCH3-phenanthroline.

1. Introduction and Experimental.- While still relatively rare for high-spin ferrous, slow paramagnetic relaxation has recently been demonstrated for six coordinate iron (II) compounds /1,2/. To our knowledge this phenomenon has not been previously documented for simple tetrahedral systems and in this work we report what is apparently the first observation of slow zero field paramagnetic relaxation and resolved magnetic hyperfine splitting in the Mössbauer spectra of distorted, monomeric, pseudo-tetrahedral coordination compounds of iron(II) with simple organo ligands. The compounds involved belong to the family [Fe(2,9 di-CH3-phenanthroline)X2], X=Cl−, Br−, I− and NCS−. Hereafter, the ligand 2,9-di-CH3-phenanthroline is abbreviated as dmp. We have studied the slow relaxation in this system of compounds using Mössbauer spectroscopy and Faraday susceptibility apparatus previously described /3,4/ for undiluted powder samples. The compounds are pseudo-tetrahedral monomers as shown by near infrared visible and far infrared spectra /5/ and isomer shifts /6/ as well as solution conductivity measurements /7/ (soluble non-electrolyte behavior) in accord with stoichiometries indicated by chemical analyses. In addition, X-ray powder studies /8/ show that Fe(dmp)Cl2 is isomorphous to Zn(dmp)Cl2 for which a single crystal X-ray investigation /9/ of its structure clearly indicates monomeric pseudo-tetrahedral structure units.

2. Mössbauer Spectroscopy Results.- The Mössbauer spectrum of the Cl− system over the range 1.6 to 300 K gives no evidence of magnetic hyperfine splitting implying positive zero field splitting (D>0) while the corresponding NCS− system exhibits progressively resolved splitting in the decreasing temperature range 20 to 1.6 K, implying D<0. It is likewise found that D goes from positive to negative ongoing from Fe(2,9 di-CH3-4,7-di-phenyl-phenanthroline)Cl2 to corresponding NCS− where in the latter hyperfine splitting occurs in the range 4.2 to 1.6 K suggesting a smaller |D| for the phenylated system. Some typical Mössbauer spectra for the extremes of slow and rapid relaxation are shown in figure 1 for the case of Fe(dmp)(NCS)2 where the transitions of the quadrupole doublet of the rapidly relaxing phase have completely disappeared at low temperatures. A small applied field (~3kG) at 4.2 K results in a Mössbauer spectrum quite similar to that observed in zero field at 3 K, suggesting field induced slow relaxation at higher temperatures. A more complete study of the field dependence of the Mössbauer spectra of these compounds over the range 1.6 K to 300 K and 0 to 100 kG is the subject of future publication and is now in progress. To conclude this section we point out that there is no evidence of the foregoing
relaxation broadening in the Mössbauer spectra of other tetrahedral analogues: e.g. Fe(biquinoline)X₂ (for H₀ = O) (X=Cl⁻, NCS⁻) behave as rapidly relaxing paramagnets to as low as 1.6 K. Thus the observed slow relaxation appears to be peculiar to the anion ligation of the 2,9 di-CH₃-phenanthroline systems. These compounds rise to a maximum of 8.10μB at 3.01 K and then decreases to 6.94μB at 1.5 K. In the case of D>O and for comparable [D], one expects μₜ > μ∥ and |μ⊥-μ∥| i.e. the anisotropy to be smaller with the maximum value of μ smaller as well. A powder sample of the latter type would then be expected to show a less pronounced low temperature maximum in μ if at all.

3. Susceptibility Studies.- The foregoing Mössbauer results, i.e D<O for Fe(dmp)(NCS)₂ and D=0 for the chloride are further supported in the temperature dependence of the effective moment (Figure 3). The figure 3 shows μ_eff versus T for an applied field of 1.6 kG for the thiocyanate in powder form. The complex, however, apparently behaves as an oriented single crystal sample in our Faraday balance sample bucket owing to large magnetic anisotropy and significant sample polarization and alignment in the field. The temperature dependence of the molar susceptibility is unremarkable. However, the novel low temperature maximum and large value of μ are expected /10,11/ for a single crystalline sample with μₜ > μ∥ implying D<O and large (Δν = 2 to -5 cm⁻¹). In the present NCS⁻ compound μ has a value of 5.60μB at 104 K.

4. High Field Mössbauer Spectra.- Magnetically perturbed Mössbauer spectra have been determined for Fe(dmp)Cl₂ for fields up to 25 kG at 300 K at which temperature it is a rapidly relaxing paramagnet. The spectra indicate that γ₀ and that quadrupole coupling constant is positive. Since Q(⁷³Fe) is positive, ν_ZZ is also positive. These results are consistent with an elongation along the two-fold axis of the tetrahedral FeN₂Cl₂ chromophore. This stabilizes the |x²-y²> orbital, or a linear combination of |x²-y²> and |x²-y²> weighted heavily with the
former to give an orbital singlet \((\mathcal{S})\) ground term. The elongation of the \(\text{FeN}_2\text{Cl}_2\) chromophore in \(\text{Fe(dmp)}\text{Cl}_2\) is a consequence of a small "bite" angle between the imine nitrogens of the bidentate ligand.

From a single crystal study of the isomorphous \(\text{Zn(dmp)}\text{Cl}_2\), this "bite" angle (N-Zn-N) was measured to be 81°. For complex with an undistorted tetrahedral geometry, the "bite" angle is 109° 28'. It is expected that the other pseudotetrahedral complexes, \(\text{Fe(biquinoline)}\text{Cl}_2\), \(\text{Fe(biquinoline)(NCS)}_2\) and \(\text{Fe(dmp)(NCS)}_2\) should also have a similar structure with a "bite" angle (N-Fe-N) of the imine nitrogens of 2,2'-biquinoline and 2,9-dimethyl-1,10-phenanthroline of \(\approx 80^\circ\). The sign of \(\nu_{zz}\) for each of the other pseudotetrahedral complexes should also be positive using reasoning analogous to the foregoing and this is, indeed, observed to be the case for their perturbed spectra.

5. Discussion. - It may not be totally unexpected that slow paramagnetic relaxation is readily observed (i.e. in zero field above 1 K) in pseudotetrahedral iron(II) compounds and perhaps more so than for octahedral systems. For an undistorted six coordinate iron(II) complex (in \(T_d\) symmetry), the ground state is \(^5T_2\) which contains residual orbital angular momentum. The system can then, rapidly relax through interactions between the electronic state and phonons of the lattice vibrations (spin-lattice relaxation). Most "real" octahedral ferrous complexes are distorted such that the orbital ground state is a singlet \(^4S\) or \(^8S\) which is further split via spin-orbit coupling to give a non-magnetic ground singlet and two excited Kramers doublets thus precluding slow relaxation. On the other hand, a compound with undistorted tetrahedral geometry (\(T_d\) symmetry) has a \(^4E\) ground state. Any orbital angular momentum present in the ground state, is obtained by mixing the \(^5T_2\) excited state into the ground state via spin-orbit coupling. For the present systems this mixing effect should be small since the centroid of the excited \(^5T_2\) state is \(\approx 6000\) to \(7000\) cm\(^{-1}\) above the ground \(^4E\) state, and both of these states are split in complexes whose symmetry is at most \(C_2\)\(_y\). Our near-infrared spectra indicate an excited \(^5T_2\) state splitting of the order of 900 to 1000 cm\(^{-1}\) and a tetrahedral \(10\Delta Q\) of 6500 cm\(^{-1}\). The relatively temperature independent, quadrupole splitting \(\Delta(\Delta E_Q)\) \(\approx 0.3\) mm/s over the range 300 to 4.2 K) and large magnitude \(\approx 2.5\) to \(2.8\) mm/s) for the present systems correspond to a ground \(^4E\) state splitting at least \(\approx 500\) cm\(^{-1}\) to 1000 cm\(^{-1}\). Since the free ion term spin-orbit coupling for the ferrous ion is \(\approx 100\) cm\(^{-1}\), the preceding considerations indicate that spin-orbit coupling enhancement of the ground state orbital angular momentum will not be large. In the absence of residual orbital angular momentum such a tetrahedral complex can less readily interact with the lattice phonons and the spin-lattice relaxation times are expected to be longer as the temperature decreases.

The remaining electronic relaxation process is then, spin-spin relaxation whose rate will be small for a ground \(m_s = \pm 2\) corresponding to \(D\langle 0\rangle\), since highly forbidden \(\Delta m_s = \pm 4\) transitions are involved. The exact nature of the high temperature relaxation process, i.e. spin-spin vs. spin-lattice must wait dilution studies. Thus we have initiated studies of the temperature dependence of \(^57\text{Fe}\) diluted in the isomorphous diamagnetic \(\text{Zn(dmp)}\text{Cl}_2\) and \(\text{Zn(dmp)}\) (NCS)\(_2\) hosts. However, previous experience with ferrous systems suggests that spin-lattice relaxation is the dominant relaxation mechanism.

References


