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A Mössbauer Study of Solvation Effects of Five-Coordinated Iron(III) Complexes with Sulfur-Containing Ligands

By J. L. K. F. de Vries, J. M. Trooster, and E. de Boer*

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Mössbauer spectra of bis(N,N-diethylthiocarbamato)iron(III) chloride and thiocyanate and the tetrabutylammonium salts of bis(maleonitriledithiolato)iron(III) and bis(toluenedithiolato)iron(III) were measured. The observed large reduction of the quadrupole splitting on going from the solid state to the solution has been explained by a change from five- to six-coordination, brought about by the binding of a solvent molecule. Some four-line spectra observed for the solid state may be explained as superposition spectra of partly five- and partly six-coordinated iron.

Introduction

It is well known that iron(III) complexes with bidentate sulfur-containing ligands occur in the solid state as five-coordinated compounds. The structure of these compounds in solution, however, is unknown. In a recent review article it was suggested that six-coordination might occur in solution; the resulting complex will then contain one or two solvent molecules. Such a transformation should reveal itself in the Mössbauer spectra of these iron(III) compounds and we decided therefore to investigate the solvation effect by means of Mössbauer spectroscopy.

The spectra of bis(N,N-diethylthiocarbamato)iron(III) chloride and thiocyanate (Figure 1, I; dtc)

and the tetrabutylammonium salts of bis(maleonitriledithiolato)iron(III) (Figure 1, II; MNT) and bis(toluenedithiolato)iron(III) (Figure 1, III; TDT) show indeed a considerable decrease in the quadrupole splitting (QS) when these compounds are dissolved. We believe that the transformation to six-coordination is responsible for this.

Experimental Section

a. Preparation.—The complexes were prepared as described elsewhere, the purity being checked by C, H, and N analyses; deviations from the theoretical contents did not exceed 0.4%. The purity of the compounds under investigation was confirmed elsewhere, the purity being checked by C, H, and N analyses.

The spectra of bis(N,N-diethylthiocarbamato)iron(III) chloride and thiocyanate (Figure 1, I; dtc)

spectra. The ir and uv-visible spectra of Fe(dtc)Cl are also the same as published elsewhere.

b. Mössbauer Measurements.— Mössbauer spectra were recorded on a constant-acceleration apparatus, Co in palladium being used as a source. All spectra, listed in Table I, were recorded at liquid nitrogen temperature. The isomer shifts (IS) are given relative to sodium nitroprusside; the accuracy of the data is 0.04 mm/sec for the solid compounds and the dimethylformamide (DMF) and pyridine solutions but 0.08 mm/sec for the other solutions. In the latter case the accuracy is limited by the large line width and the small Mössbauer fraction of these dilute solutions. In nonpolar solvents like chloroform and benzene the absorption was insufficient to get reliable data; the best spectra were obtained in DMF, the solutions being about 0.03 M.

As Fe(dtc)Cl in solution can be hydrolyzed or converted to the tris-dtc complex, all solutions were prepared shortly before the measurements. They were injected in a polypropylene capsule filled with cotton, immediately sealed off, and cooled down to liquid nitrogen temperature. Spectra of samples not prepared under these conditions sometimes showed extra lines which were not reproducible and were therefore ascribed to decomposition products.

c. Other Physical Measurements.—Ir spectra were recorded on a Hitachi EPI-G3 in the 4000-700-cm⁻¹ region (KBr disks) and on a Hitachi EPI-L in the 700-200-cm⁻¹ region (CaI disks). Electronic spectra were taken with a Cary 14 spectrometer. In the 5000-12,000-cm⁻¹ region the compounds were also measured in the solid state; as samples we used KBr disks containing about 15 mg/cm² of complex. The C, H, and N analyses were performed on a Perkin-Elmer 240 instrument. For conductivity measurements use was made of a Wayne Kerr B 641 universal bridge.

Results and Discussion

The results of our measurements are collected in Tables I and II. The most striking result is the reduction of the QS on going from the solid state (about 2.8 mm/sec) to the solution (about 0.7 mm/sec).

1. Bis(N,N-diethylthiocarbamato)iron(III) Chloride.

a. The Mössbauer Spectra.—The structure of Fe(dtc)Cl in the solid state has been depicted in Figure 2A. It exists in monomeric form and is five-coordinated, the Fe-Cl distance being almost the same as the Fe-S distances. According to calculations of Ake and Harris Loew the unusually large electric field gradient

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obtained. Using these values the QS was calculated in good agreement with method the occupation numbers of the iron 3d and 4p orbitals are recently carried out extended Hückel calculations for Fe(dtc)2.

of this Fe(III) complex (see Table I) arises solely from the lattice. These authors assumed, however, that covalency effects could be entirely neglected. However, strong covalency is known to occur in dithiocarbamates of related metals,14,15 and a contribution to the electric field gradient from the valence electrons cannot be excluded. Recent calculations of the electric field gradient in some dithiolene complexes by Golding et al.,16 support this. These authors explained the total field gradient by the valence electrons, taking into account covalency.16a

It is tempting to interpret the remarkable reduction of the QS on going from the solid state to the solution as a change in the coordination number from 5 to 6 caused by the binding of a solvent molecule, thus yielding a complex of higher symmetry. This change of geometry of the complex, resulting in a different order of the molecular orbitals involved is then responsible for the observed change in QS. This explanation is supported by the experimental data obtained for Fe(dtc)3, in which compound the iron atom is six-coordinated and located at the center of a distorted octahedron.17 This six-coordinated complex is like Fe(dtc)2Cl11 present in the monomeric form, both in the solid state and in solution.17 In chloroform solution about the same magnetic susceptibility has been measured for both compounds.11,17

Figure 2.—Mössbauer spectra of Fe(dtc)3 (a) in solid state and (b) in DMF solution and of Fe(dtc)2Cl (c) in DMF solution, (d) in solid state, and (e) after complete removal of the DMF from sample (c). All spectra were recorded at liquid nitrogen temperature.

Figure 3 shows that the Mössbauer spectrum of Fe(dtc)3 in solution is almost identical with the spectrum of the six-coordinated Fe(dtc)3 in solution. The similarity in magnetic susceptibility and in the IS and QS parameters (cf. Table I) suggests a geometrical correspondence between both complexes in solution, which can be attained by the binding of a solvent molecule to the sixth coordination site of the Fe(dtc)2Cl monomer. In principle, the solvent molecule may enter the complex either cis or trans with respect to the chlorine atom. Therefore on the basis of the Mössbauer spectra alone we cannot decide which configuration is formed. However, chemical evidence favors the cis model.

It is known that Fe(dtc)2Cl can be rapidly and re-

Note Added in Proof.—In cooperation with Dr. C. P. Keijzers we recently carried out extended Hückel calculations for Fe(dtc)2Cl, by which method the occupation numbers of the iron 3d and 4p orbitals are obtained. Using these values the QS was calculated in good agreement with the experimental values. This supports the statement that the lattice contribution to the electric field gradient in Fe(dtc)2Cl is negligible.

Table I

<table>
<thead>
<tr>
<th>Compound</th>
<th>IS Solid state</th>
<th>QS Solid state</th>
<th>IS Solution</th>
<th>QS Solution</th>
</tr>
</thead>
<tbody>
<tr>
<td>Fe(dtc)Cl</td>
<td>0.70</td>
<td>2.76</td>
<td>0.70</td>
<td>0.70</td>
</tr>
<tr>
<td>Fe(dtc)SCN</td>
<td>0.68</td>
<td>2.65</td>
<td>0.68</td>
<td>0.79</td>
</tr>
<tr>
<td>Fe(dtc)2</td>
<td>0.70</td>
<td>0.52</td>
<td>0.69</td>
<td>0.66</td>
</tr>
<tr>
<td>Fe(MNT)2-</td>
<td>0.62</td>
<td>2.73</td>
<td>0.69</td>
<td>0.84</td>
</tr>
<tr>
<td>Fe(TDT)2-</td>
<td>0.59</td>
<td>3.09</td>
<td>0.68</td>
<td>0.85</td>
</tr>
</tbody>
</table>

Table II

<table>
<thead>
<tr>
<th>Solvent</th>
<th>IS (mm/sec)</th>
<th>QS (mm/sec)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Dimethylformamide</td>
<td>0.70</td>
<td>0.70</td>
</tr>
<tr>
<td>Acetonitrile</td>
<td>0.72</td>
<td>0.77</td>
</tr>
<tr>
<td>Dimethyl sulfoxide</td>
<td>0.71</td>
<td>0.84</td>
</tr>
<tr>
<td>1,2-Dimethoxyethane</td>
<td>0.73</td>
<td>0.89</td>
</tr>
<tr>
<td>Pyridine</td>
<td>0.70</td>
<td>0.71</td>
</tr>
<tr>
<td>Thiocyanate in DMF</td>
<td>0.69</td>
<td>1.40</td>
</tr>
</tbody>
</table>

Figure 2.—Molecular structure diagrams of (A) bis(N,N-diethylthiocarbamato)iron(III) iron and (B) bis(maleonitriledithiolato)iron(III) iron.14,15
versibly converted to Fe(dtc)$_2$Cl. This can easily be understood for the cis configuration, since the iron atom has two neighboring sites available for binding the bidentate ligand. Therefore we favor a cis configuration for the Fe(dtc)$_2$(Cl)(solv) complex as depicted in Figure 4.

Tables I and II show that the IS does not vary markedly upon a change of coordination number or ligand. This has been observed previously: the IS's of Fe(MNT)$_3^{3-}$ and Fe(MNT)$_2^{2-}$ differ by only 0.06 mm/sec, whereas Fe(MNT)$_2^{2-}$ and Fe(MNT)$_3$(py)$_2^{3-}$ have the same IS. Also no change of IS is observed in the series Fe(dtc)$_2$X, X being Cl, Br, I, NCS, or C$_6$F$_5$CO$_2$Cl.

As can be seen in Table II, a second site was observed for solutions of Fe(dtc)$_2$Cl in pyridine and in DMF to which potassium thiocyanate was added. The large IS suggests that the extra lines belong to a high-spin iron (II) complex of which no further identification has been attempted.

b. The Electronic Spectra.—The proposed change of the coordination number must also reveal itself in a change of the d–d transitions in the electronic spectrum. For that purpose we measured the electronic spectrum of Fe(dtc)$_2$Cl, both in solid form and in solution, in the region from 5000 to 12,000 cm$^{-1}$. Figure 5 shows the similar spectrum was obtained by Martin and White by dissolving Fe(dtc)$_2$Cl in the very weakly coordinating solvent CCl$_4$. In other solvents with better coordinating power no spectrum was observed. One may conclude from this that CCl$_4$ is not capable of coordinating with Fe(dtc)$_2$Cl. Unfortunately we could not verify this statement from the Mössbauer spectrum of Fe(dtc)$_2$Cl in CCl$_4$, due to the low solubility and the small Mössbauer fraction. In solution no absorption bands are observed in the 8000–12,000 cm$^{-1}$ region, even for saturated solutions and very long cells. The absorption spectrum outside this region (visible and uv) is the same as published spectra in the literature. It should also be noted that the same visible–uv spectrum is produced by the material having the small QS of 0.70 mm/sec.

The absence of absorption bands for solutions of Fe(dtc)$_2$Cl in the optical region, where d–d transitions for five-coordinated iron(III) complexes occur, strongly supports our conclusion that a change in coordination number takes place on dissolving Fe(dtc)$_2$Cl.

c. The IR Spectra.—The presence of bound solvent molecules must also reveal itself in the ir spectra of the solutions. Unfortunately the solubility of Fe(dtc)$_2$Cl in various solvents was not high enough to measure ir spectra. However, evidence for solvent binding was found by measuring ir spectra of the solid material, obtained by evaporating a Fe(dtc)$_2$Cl solution in DMF, which is a strongly coordinating solvent. In Figure 6 some relevant ir spectra are shown. The ir spectrum of the material, obtained from the solution, clearly shows peaks due to DMF. They are shifted with respect to corresponding positions in pure DMF. For example, in the complex peaks are found at 655, 1110, 1242, and 1405 cm$^{-1}$ (indicated by arrows in Figure 6), while in pure DMF these peaks are at 660, 1097, 1254, and 1388 cm$^{-1}$. The Mössbauer spectra of these samples all contained the small QS of 0.70 mm/sec thought to be indicative of six-coordination.

Removal of a weakly coordinating solvent like acetonitrile or chloroform resulted in unchanged Fe(dtc)$_2$Cl, as could be concluded from its ir and Mössbauer spectra. Hence no decomposition takes place by dissolving the compound.
d. Reactions in DMF.—Some interesting phenomena occurring in DMF but not observed in other solvents merit special attention.

After evaporating to dryness a Fe(dtc)2Cl solution in DMF, the remaining bound solvent molecules could be completely removed by slightly heating the residue on the vacuum line. The ir spectrum of the residue resembled very much that of Fe(dtc)2Cl and no peaks due to DMF were present any more. The Mössbauer spectrum, however, surprisingly shows a four-line spectrum (see Figure 3e). It appeared to be a superposition of the spectrum of Fe(dtc)2Cl in the solid state and in solution. Thorough C, H, and N analyses of these samples proved that we are dealing with a compound having the same composition as the pure starting material. Anal. Calcd: C, 30.97; H, 5.20; N, 7.22. Found: C, 30.31, 30.47, 30.31, 30.29; H, 5.08, 5.23, 5.14, 5.15; N, 7.17, 7.30, 7.34, 7.30. Recrystallization of these samples from chloroform yielded the original complex, as indicated by its Mössbauer and ir spectra.

The observations can be rationalized, if dimer formation is assumed according to

$$2\text{Fe(dtc)2(Cl)(solv)} \rightleftharpoons [\text{Fe(dtc)2Cl}]^+ + 2\text{DMF}$$

(1)

Apparently the DMF coordinates so strongly that in the paste, formed by evaporation of DMF, the complex is able to retain six-coordination by forming dimers. The six-coordinated iron atom in the dimer will give a Mössbauer spectrum similar to the solvated Fe(dtc)2Cl in solution.

Another interesting observation was made when a DMF solution of Fe(dtc)2Cl was very slowly evaporated on a vacuum line. At the beginning of the evaporation process black crystals were formed, the ir spectrum being given in Figure 7, together with the ir spectrum of pure Fe(dtc)3. On close inspection it can be noticed that this spectrum is a superposition of the ir spectra of pure Fe(dtc)3 and Fe(dtc)2Cl. This can be most clearly seen in those regions where the ir spectra of the two latter compounds differ markedly (in Figure 7 indicated by arrows). We conclude, therefore, that slow evaporation yields crystals which contain a fair amount of Fe(dtc)3. This conclusion is further supported by the Mössbauer spectrum of these crystals (a four-line spectrum due to Fe(dtc)2Cl and Fe(dtc)3) and the C, H, and N analyses.

e. Dissociation.—Slow addition of DMF to a solution of Fe(dtc)2Cl in acetonitrile causes a 10-fold increase in the conductivity. It was shown earlier that Fe(dtc)2Cl is a weak electrolyte and a dissociation in Fe(dtc)+ and Cl- ions was suggested. We prefer to formulate the dissociation process, in accord with our proposed model

$$\text{Fe(dtc)(Cl)(solv)} + \text{solv} \rightleftharpoons \text{Fe(dtc)(solv)H}^+ + \text{Cl}^-$$

in which a chlorine atom is replaced by a solvent molecule, thus producing a disolvated Fe complex. The Mössbauer spectra of these solutions were indistinguishable from the other solution spectra.

In conclusion, the various reactions occurring in DMF can be exemplified by the reaction scheme

$$\text{Fe(dtc)(solv)H}^+ + \text{Cl}^- \rightleftharpoons \text{Fe(dtc)(solv)}^2+ + \text{Cl}^-$$

2. Dithiolene Complexes.—The five-coordination in the dithiolene complexes occurs in the solid state by association to dimers (Figure 2B),2,3 in the solvents which we used, Fe(MNT)2- and Fe(MNT)2_ complexes are monomeric.19 In the solid state these compounds have a doublet spin state, whereas the monomers in solution have a quartet spin state.20 The reduction of the QS (see Table I and Figure 8) on dissolving these complexes is similar to the effect observed for the di-thiocarbamate complex, suggesting that also for these compounds six-coordination takes place in solution.

A comparison with the Mössbauer data of the complexes Fe(MNT)33- and Fe(MNT)32-, which in solid state have a QS of 1.69 and 1.57 mm/sec, respectively,10 cannot be made because these two complexes have one and two unpaired electrons, respectively.

\begin{figure}
\centering
\includegraphics[width=\textwidth]{figure8}
\caption{Mössbauer spectra of Fe(MNT)33- in solid state (a) and (b) in DMF solution, at liquid nitrogen temperature.}
\end{figure}


(20) L. Carlin, \textit{Transition Metal Chem.}, 1, 263 (1965).
On the analogy of the dithiocarbamate complexes we assume that a cis configuration is favored in the Fe-(MNT)$_2$(solv)$_2$~ and Fe(TDT)$_2$(solv)$_2$~ complexes. On heating in air, a Fe(MNT)$_2$~ solution in acetone readily produces$^{21}$ Fe(MNT)$_3^{2-}$. When a solution of Fe(MNT)$_2$~ or of Fe(TDT)$_2$~ in DMP or in other solvents is evaporated on the vacuum line, the original Mössbauer spectrum is found again.

It is noteworthy that Stukan, et al.,$^{22}$ in studying the related bis(naphthadithiolato)iron(III) complex (Figure 1, IV), found four-line Mössbauer spectra. The authors discussed the possibility of the simultaneous occurrence of a cis- and trans-solvated complex. According to our view such complexes will not have strongly different Mössbauer spectra; therefore we propose an explanation similar to the one presented for Fe(dtc)$_2$Cl: the spectrum consists of one site with five-coordination [IS = 0.47, QS = 3.0 mm/sec; cf. Fe(TDT)$_2$~ (solid state) IS = 0.59, QS = 3.09 mm/sec] and one site with six-coordinated iron [IS = 0.58, QS = 0.82 mm/sec; cf. Fe(TDT)$_2$~ (in solution) IS = 0.68, QS = 0.85 mm/sec].

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