

Iodo(etiohemiporphycenato)iron(III). Unexpected Difference in Magnetic Behavior in Solution and Solid

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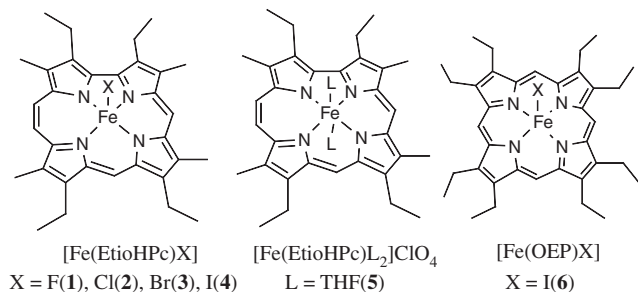
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Although iodo(etiohemiporphycenato)iron(III) showed an admixed intermediate-spin state ($S = 3/2$, $5/2$) with a major contribution of $S = 3/2$ in solution, the same complex exhibited the high-spin state ($S = 5/2$) in the solid phase. Importance of the crystal packing has been pointed out for the formation of the high-spin complex in the solid.

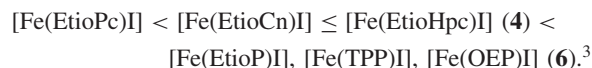
Porphyrin isomers such as porphycene (Pc), corphycene (Cn), and hemiporphycene (Hpc) have attracted much attention because of their characteristic molecular structures.¹ The central cavities surrounded by the four nitrogen atoms are quite different among Pc, Cn, and Hpc; they are rectangular, trapezoidal, and quadrilateral, respectively. Thus, each macrocycle serves a unique ligand field to the central metal ion, which could in turn give unique physicochemical properties to the metal complexes different from those of the corresponding porphyrin (P) complexes.^{2,3} Among the porphyrin isomers, Hpc is quite peculiar because the macrocycle possesses four nonequivalent nitrogen atoms. Thus, the d-orbitals of Hpc complexes could split differently to form the complexes with unique electronic structure. Although several groups have reported the synthesis and characterization of Hpc and its metal complexes,^{4–8} little is known on the physicochemical properties of the iron complexes. Here, we report the spin states of 5- and 6-coordinated (etiohemiporphycenato)iron(III) complexes, **1–5**, both in solution and in the solid.⁹ We also report the X-ray molecular structures of **4** and the analogous porphyrin complex(**6**).



(EtioHpc) H_2 was synthesized from 3,3'-diethyl-5'-formyl-4,4'-dimethyl-2,2'-dipyrrylmethane and 3,3'-diethyl-5,5'-diformyl-4,4'-dimethyl-2,2'-bipyrrrole.^{5,10} Insertion of iron followed by the HCl treatment afforded **2**, which was further converted to the corresponding μ -oxo-dimer. Complexes **1**, **3**, and **4** were prepared by the cleavage of the μ -oxo-dimer with HClO_4 in the presence of the corresponding potassium halides. **5** was pre-

pared by addition of AgClO_4 to a THF solution of **2** followed by the recrystallization from $\text{CH}_2\text{Cl}_2/\text{THF}$ solution.

Figure 1 shows the molecular structures of **4** and **6** determined by X-ray crystallographic analysis.^{11,12} Because **4** is a chiral complex, the crystals are racemic; both the enantiomers are placed as a pair around the center of symmetry. Table 1 lists the selected structural data of **4** and **6** together with those of some relevant complexes.^{3,12} In the previous paper, we have pointed out that the essentially pure intermediate-spin ($S = 3/2$) complexes are characterized by the narrower cavity and smaller iron displacement, ΔFe , than the high-spin ($S = 5/2$) complexes.³ The data in Table 1 indicate that the cavity size and ΔFe value are the smallest in the $S = 3/2$ complex, $\text{Fe}(\text{EtioPc})\text{I}$, and increase in the following order:



Since $[\text{Fe}(\text{EtioCn})\text{I}]$ has been characterized as a high-spin complex, it is reasonable to consider that **4** is also a high-spin complex from the X-ray crystallographic point of view. To further examine the spin states of **4** and **6** in the solid, we have measured the effective magnetic moments (μ_{eff}) by SQUID magnetometry over 2–300 K. Complexes **4** and **6** are high-spin be-

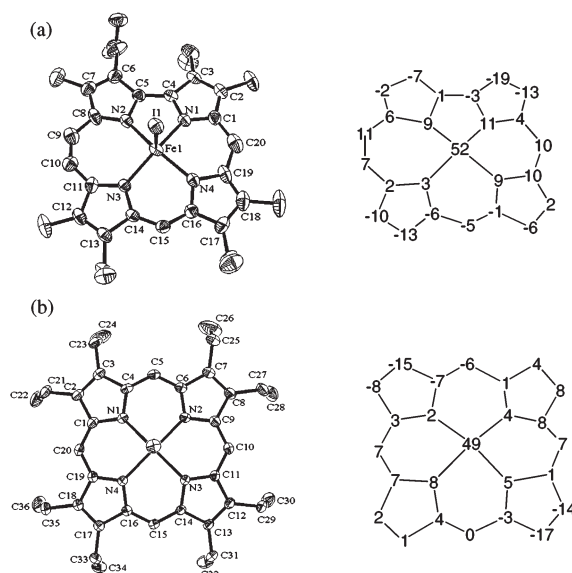


Figure 1. Molecular structures of (a) **4** and (b) **6**. Left: ORTEP drawings, right: perpendicular displacements of each atom from the least-squares plane of peripheral 24 atoms.

Table 1. Some structural data

Complexes	Fe–N _p /Å	Fe–X /Å	Cavity area /Å ²	ΔFe /Å	Spin state ^a	Ref
Fe(TPP)I	2.07	2.55	8.123	0.459	5/2(5/2)	12
Fe(EtioP)I	2.061(7)	2.617(1)	8.094	0.454(4)	5/2(5/2)	3
Fe(EtioCn)I	2.034(2)	2.615(1)	7.897	0.387(1)	5/2(5/2)	3
	2.056(2)					
Fe(EtioPc)I	1.956(3)	2.664(1)	7.355	0.343(2)	3/2(3/2)	3
Fe(EtioHpc)I(4)	2.042(4)	2.618(1)	7.882	0.438(2)	5/2(3/2)	b
	2.039(5)					
	2.051(4)					
	2.050(5)					
Fe(OEP)I(6)	2.068(2)	2.610(1)	8.151	0.466(1)	5/2(5/2)	b

^a Spin state in the solid. Spin state in solution is given in the parenthesis.

^b This work.

cause the μ_{eff} values are 5.5–5.9 μ_{B} above 30 K. The Mössbauer spectra have also suggested that these complexes are high spin at 77 K; the *IS* and *QS* values are 0.39 and 1.34 mm s^{−1} for **4** and 0.43 and 1.10 mm s^{−1} for **6**, respectively. It should be noted that the Mössbauer spectra of **4** contain ca. 20% of **2**; the *IS* and *QS* values of **2** are 0.36 and 0.58 mm s^{−1}, respectively, at 77 K.

The spin states of **1–6** have then been examined in solution. The ¹H NMR spectra of **1–4** were fairly complicated because these complexes have four nonequivalent methyl and ethyl groups. The μ_{eff} values determined by the Evans method in CH₂Cl₂ solution at 298 K were 5.4, 4.5, 4.1, and 5.3 μ_{B} for **2**, **4**, **5**, and **6**, respectively. The results suggest that, while **2** and **6** are in the *S* = 5/2 state with minor contribution of *S* = 3/2, **4** and **5** are in the *S* = 3/2 state with minor contribution of *S* = 5/2. The slightly larger μ_{eff} value of **4**, 4.5 μ_{B} , could be the result of contamination of high-spin **2** as is revealed from the Mössbauer result. We have then examined the EPR spectra in frozen CH₂Cl₂–toluene solution at 4.2 K. The EPR spectra of **1**, **3**, and **4** exhibited the hyperfine coupling with the axial ligands, indicating explicitly that F[−], Br[−], and I[−] actually bind to the iron. The *g* values are as follows; **1**: *g*_x = *g*_y = 5.86, *g*_z = 2.03; **2**: *g*_x = *g*_y = 5.80, *g*_z = 2.00; **3**: *g*_x = 6.09, *g*_y = 5.80, *g*_z = 2.00; **4**: *g*_x = *g*_y = 4.24, *g*_z = 2.00; **5**: *g*_x = 4.92, *g*_y = 3.84, *g*_z = 1.97; **6**: *g*_x = *g*_y = 5.80, *g*_z = 1.96. Consistent with the Mössbauer result, the EPR spectrum of **4** showed contamination of **2**. Nevertheless, the EPR *g* values clearly indicate that **4** and **5** adopt mainly the *S* = 3/2 state while **1**, **2**, **3**, and **6** exhibit *S* = 5/2.

The question arises as to why the spin state of **4** in solution is different from that in the solid. One possible reason for this anomaly is an extraordinarily labile nature of the iodide ligand in **4**. In fact, we found that **4** was completely converted to **2** by addition of 1.0 equiv. of Bu₄N⁺Cl[−]. By contrast, only 40% of **6** was converted to [(OEP)FeCl] under the same condition. The results suggest that the coordination of iodide ion to the iron in **4** must be much weaker than that in **6**. As a result, the iron is dragged toward the N4 cavity and, consequently, the complex adopts the intermediate-spin state in solution.^{13–15} In the crystal lattice, however, the packing force could contract the labile Fe–I bond. Therefore, the iron is lifted toward the iodide ion, resulting in the formation of a typical high-spin 5-coordinated structure. The hypothesis mentioned above is supported by the fact that the Fe–I bond length in **4** is shorter by 0.046 Å while the ΔFe is longer by 0.095 Å than the corresponding values in [Fe(EtioPc)I]. On the basis of these results, we have concluded that the crystal packing is one of the important factors for the

formation of the high-spin complex in the solid.

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- Abbreviations: EtioPc, TPrPc, EtioHpc, OEP, TPP; dianions of 3,6,13,16-tetraethyl-2,7,12,17-tetramethylporphycene, 2,7,12,17-tetrapropylporphycene, 3,6,13,17-tetraethyl-2,7,12,18-tetramethylhemiporphycene, 2,3,7,8,12,13,17,18-octaethylporphyrin, and 5,10,15,20-tetraphenylporphyrin, respectively.
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- Crystallographic details*: Pure crystals of **4** and **6** were obtained from the CHCl₃ solutions. *Crystal data for 4* (FeC₃₃H₃₇Cl₃N₄I, *M* = 778.77) were collected on Rigaku RAXIS-RAPID Imaging plate diffractometer at 298 K with MoKα radiation (λ = 0.71069 Å). Crystal system, triclinic; Space group, *P*-1(#2); *a* = 9.430(1), *b* = 13.089(1), *c* = 14.392(1) Å, α = 115.73(1)°, β = 106.72(1)°, γ = 90.69(1)°, *V* = 1513.2(2) Å³, *Z* = 2. A total 13614 reflections were collected and of 6591 unique reflections 4679 (*I* > 2σ(*I*)) were used. Final *R* factors were *R*1 = 0.070 (observed data) and *wR*2 = 0.209 (all data); CCDC reference number 200304. *Crystal data for 6* (FeC₃₇H₄₅Cl₃N₄I, *M* = 834.87) were collected similarly as in the case of **4**. Crystal system, triclinic; Space group, *P*-1(#2); *a* = 10.342(2), *b* = 13.846(2), *c* = 14.815(3) Å, α = 71.12(1)°, β = 83.80(1)°, γ = 76.42(1)°, *V* = 1950.0(6) Å³, *Z* = 2. A total 15664 reflections are collected and of 8691 unique reflections 7061 (*I* > 2σ(*I*)) were used. Final *R* factors were *R*1 = 0.070 and *wR*2 = 0.209 (all data). CCDC reference number 200305.
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