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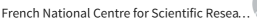
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Reaction of a Carbon Equivalent of Iodosylbenzene with Iron-Porphyrins: A Ready Access to N-Alkylporphyrin Complexes Involving Five-Membered Fe-O-C-C-N Metallocycles

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Abstract: Iron(II)-porphyrin complexes, Fe^{II}P (P = tetraarylporphyrin: TPP, TpClPP, and TTP), react with the iodonium ylide $C_6H_5I=X$ (X = $CC(O)CH_2C(CH_3)_2CH_2C(O)$) to give Fe^{II} -bis-N-alkylporphyrin complexes exhibiting a structure in which the metal and two trans pyrrole nitrogen atoms are bound respectively to the oxygen and the carbon atom of two X moieties. The bismetallocyclic structures of these Fe^{II} high-spin complexes result from the insertion of two -O-C-C- groups of the X moiety into two trans Fe-N bonds. These structures deduced from X-ray crystallography for one complex and proposed from the others agree with their elemental analysis and ¹H NMR, UV-vis, and mass spectra. In contrast, complexes Fe^{III}(P = tetraarylporphyrin = TPP, TpClPP, TTP, and TMP)(ClO₄) react with the iodonium ylide $C_6H_5I=X$ to give the corresponding high-spin monometallocyclic Fe^{III} -N-alkylporphyrin complexes whose structures have been deduced from (i) their elemental analysis, (ii) their mass and ¹H NMR spectra, and (iii) the complete transformation of three of them (TPP, TpClPP, and TTP) into the corresponding bismetallocyclic Fe^{II} -bis-N-alkylporphyrin complexes upon reaction of their one-electron reduction products with the iodonium ylide. The corresponding monometallocyclic Fe^{II} -N-alkylTMP complex fails to react with C_6H_5I =Xunder identical conditions probably because of the greater hindrance of the TMP derivative. The formation and physical properties of this new class of iron-N-alkylporphyrin complexes involving five-membered Fe-O-C-C-N metallocycles are discussed in relation to the complexes formed during the suicidal inactivation of cytochrome P-450 by terminal monosubstituted 1-alkenes.

High-valent Fe^{IV}- or Fe^V-oxo^{2,3} complexes have been prepared by the reaction of iron-porphyrins with potential oxygen atom donors. They are believed to be the active species involved in alkane hydroxylation and alkene epoxidation by iodosylbenzene catalyzed by cytochrome P-4504 or iron porphyrins.5 nitrogen analogues, the Fe^{IV}- or Fe^V-imido (or iron(II)- or iron(III)-nitrene) complexes, seem to be formed upon oxidation of 1,1-dialkylhydrazines by cytochrome P-450.6 Porphyrin-iron-nitrene complexes, $Fe^{IV} = N - NR_2 \leftrightarrow Fe^{II} \leftarrow N - NR_2$, prepared by oxidation of such hydrazines, have been completely characterized.⁷ Such iron-nitrene complexes might be involved

in the transfer of N-tosyl moiety of the nitrogen analogue of PhIO, PhI=N—tosyl, into aliphatic C-H bonds⁸ or double bond of alkenes,9 catalyzed by iron(III)-porphyrins.

Their carbon analogues, the iron-carbene complexes, seem to be formed upon oxidation of certain compounds by cytochrome P-450,¹⁰ and several porphyrin-iron-carbene complexes have now been prepared and characterized.¹¹ The reactions of the carbon analogues of PhIO, the iodonium ylides, PhI=CRR', with ironporphyrins are therefore of interest.

The results described therein show that, depending upon the reaction conditions, such a iodonium ylide, compound 1 (CRR' = $CC(O)CH_2C(CH_3)_2CH_2C(O)$ = X), reacts with ironporphyrins to give either N-alkylporphyrin complexes involving one five-membered Fe-O-C-C-N metallocycle or bis-N-alkylporphyrin-iron(II) complexes involving two Fe-O-C-C-N me-

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Table I. UV-vis, Mass Spectrometry and Electrochemical Characteristics of Complexes of Type 3 and 5

			UV-visa			r h	m	iss spectra ^c n	n/e
complex	λ (1	nM)	€	(mM ⁻¹ -cm ⁻¹)	$\frac{{E_{1/2}}^b}{\mathrm{Fe^{II}}/\mathrm{Fe^{II}}}(\mathrm{mV})$		rel intnsty %	
3a	454	524	580	655	736	-205		806	668
	(84)	(18)	(8.7)	(4)	(4.9)			(50)	(100)
3b	457	524	578	652	726	-103		942	804
	(104)	(22)	(11)	(5.6)	(6.4)			(100)	(62)
3c	458	526	38 Í	654	736	-174		864	724
	(83)	(18)	(9.6)	(5.4)	(4.9)			(82)	(100)
3d	448	518	574	638	696	-300	1009	975	836
	(71.4)	(12.5)	(7.5)	(3.6)	(sh)		(3)	(100)	(18)
5a	464	562	615	746		+410	944	806	668
	(114)	(10.9)	(16.4)	(4.4)			(100)	(30)	(25)
5b	467	562	620	750		+540	1080	942	804
	(110)	(12.3)	(20)	(5.1)			(98)	(84)	(100)
5c	466	562	622	754		+340	1000	864	724
	(116)	(9.9)	(17.7)	(5.2)			(100)	(55)	(82)

aln C₆H₆. bVs. saturated calomel electrode; for complexes 3 in deaerated DMF, 0.2 M LiCl; for complexes 5 in CH₃CN, 0.2 M LiClO₄. For complexes 3: M^+ , M^+ - Ci, M^+ - Ci - X; for complexes 5 M^+ , M^+ - X, M^+ - 2X.

Scheme I

tallocycles. The structures and physicochemical properties of these complexes have been deduced from various spectroscopic methods including an X-ray analysis for one of them. 12 These results definitively establish the existence of their Fe-O-C-C-N metallocycles which derive formally from the insertion of a O-C-C moiety of 1 into iron-nitrogen bonds of the starting iron-porphyrin complexes.

The formation and properties of this new class of iron-N-alkylporphyrin complexes are discussed in relation to the complexes formed during the suicidal inactivation of cytochrome P-450 by terminal monosubstituted 1-alkenes.

Results

I. Reaction of C₆H₅I=X, 1, with Iron(II)-Porphyrins. Upon addition under argon of a CH_2Cl_2 solution of $Fe^{II}(TPP)^{13}$ to a 20-fold excess of the iodonium ylide, $^{14}C_6H_5I=X$ (1), the red color of the porphyrin rapidly turned green, and a new complex characterized by a Soret band at 462 nm was formed quantitatively within 0.5 h at 50 °C. After purification by column chromatography and crystallization from CH₂Cl₂-pentane, complex 5a was obtained as a green powder with an almost quantitative yield (>95%).

Identical reactions were performed with Fe^{II}(TpClPP)¹³ or $Fe^{11}(TTP)^{13}$ and led to the corresponding complexes 5b and 5c.

However, the reaction of Fe^{II}(TMP)¹³ with 1 failed to give a similar complex of type 5 (vide infra, part II). See Scheme I.

Characteristics of Complexes 5. In mass spectrometry, all complexes 5 exhibit molecular peaks corresponding to Fe(porphyrin)X₂ and fragments corresponding to the successive loss of one and two X moieties and indicating that the iron-porphyrin ring remains intact after these fragmentations (Table I). Accordingly, the elemental analyses (C, H, N) of complexes 5 correspond to the formula Fe(porphyrin)(C₈H₁₀O₂)₂·2H₂O, indicating the formal incorporation of two X moieties into the Fe^{II}-porphyrin complexes and a residual solvation by two water molecules (see Experimental Section).

X-ray Analysis of Complex 5a. Crystals of complex 5a suitable for an X-ray structure determination were obtained by slow crystallization from a CH₂Cl₂-pentane solvent mixture.

The asymmetric unit of the crystals contains two independent half-molecules of bis N-alkylated iron(II) porphyrin which are located on an inversion center and hence presents at least 1 symmetry. Table II gives the atomic positional parameters for all non-hydrogen atoms. Figure 1 shows one of these centrosymmetric molecular units together with the labeling scheme used. Table III gives selected bond distances and angles and their mean values.

Complex 5a (Figure 1) presents an unusual bismetallocyclic structure in which two opposite pyrrole nitrogens are alkylated by two X groups. These groups are further ligated axially to iron by one oxygen atom. The two alkylated pyrrole nitrogens, the two porphyrinato nitrogens, and the metal are necessarily coplanar. Whereas the mean value of the two Fe-Np bond distances is 2.082 (5) Å, the Fe-N(alkyl) average bond is 2.242 (5) Å. The axial Fe-O average bond distance is 1.998 (5) Å.

In all the mono-N-alkylated porphyrin complexes whose structures are known so far,15 the metal-N(alkyl) bond distances

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of mesotetraphenyl-, mesotetra(parachlorophenyl), mesotetra(4-methylphenyl)-, mesotetra(2,4,6-trimethylphenyl)porphyrin, PPIXDME and DPIXDME for those of proto- and deuterioporphyrin IX dimethylester. (14) Karele, B.; Neilands, O. Zh. Org. Khim. 1968, 4, 1818-1822.

Table II. Atomic Positional Parameters for All Non-Hydrogen

Atoms for	Complex 5a ^a		•	
atom	X	У	z	$B(Å^2)$
Fel	0.000	0.000	0.000	2.75 (4)
N1	0.0270 (4)	-0.0936 (4)	0.1255 (5)	3.0 (2)
C11	-0.0530 (6)	-0.1337 (5)	0.1463 (6)	3.2 (2)
C12 C13	-0.0099 (6) 0.0948 (7)	-0.1827 (6) -0.1711 (6)	0.2510 (7) 0.2955 (7)	4.2 (2) 4.1 (2)
C13	0.1192 (6)	-0.1157 (5)	0.2171 (6)	3.2 (2)
C5	0.2232 (6)	-0.0946 (5)	0.2245 (6)	3.2 (2)
N2	0.1640 (4)	0.0343 (4)	0.0925 (5)	2.8 (2)
C21	0.2420 (6)	-0.0394 (5)	0.1506 (6)	3.0 (2)
C22 C23	0.3369 (6)	-0.0489 (6)	0.1104 (7)	4.0 (2) 3.8 (2)
C24	0.3203 (6) 0.2134 (6)	0.0110 (6) 0.0629 (5)	0.0262 (7) 0.0114 (6)	3.0 (2)
C6	-0.1653 (6)	-0.1194 (5)	0.0803 (6)	3.1 (2)
C51	0.3214 (6)	-0.1405 (6)	0.3114 (7)	3.7 (2)
C52	0.3423 (7)	-0.2285 (6)	0.3291 (8)	5.2 (3)
C53	0.4308 (8)	-0.2676 (7)	0.4136 (9)	6.8 (3)
C54 C55	0.4989 (7) 0.4845 (7)	-0.2187 (9) -0.1326 (8)	0.4809 (8) 0.4648 (8)	7.7 (3) 6.5 (3)
C56	0.3929 (6)	-0.0927 (6)	0.3772 (7)	4.8 (2)
C61	-0.2432 (6)	-0.1561 (5)	0.1290 (7)	3.2 (2)
C62	-0.2602 (7)	-0.1326 (6)	0.2327 (8)	4.7 (2)
C63	-0.3383 (7)	-0.1614 (7)	0.2711 (8)	5.8 (3)
C64	-0.3978 (7)	-0.2150 (6)	0.2065 (8)	5.5 (3)
C65 C66	-0.3838 (7) -0.3037 (7)	-0.2412 (6) -0.2105 (6)	0.1043 (9) 0.0635 (7)	5.4 (3) 4.3 (2)
C1D1	0.1258 (5)	0.0988 (5)	0.1707 (6)	3.1 (2)
C2D1	0.0149 (6)	0.1153 (5)	0.1724 (6)	3.1 (2)
C3D1	-0.0180 (6)	0.1717 (6)	0.2629 (8)	4.4 (2)
C4D1	0.0726 (7)	0.1740 (6)	0.3736 (7)	4.2 (2)
C5D1 C6D1	0.1699 (7) 0.2073 (7)	0.1945 (6) 0.1359 (6)	0.3354 (8) 0.2500 (7)	4.9 (2) 4.3 (2)
C7D1	0.1036 (9)	0.0883 (7)	0.4369 (9)	6.8 (3)
C8D1	0.0335 (8)	0.2419 (7)	0.4508 (8)	5.7 (3)
OlDi	-0.0607 (4)	0.0862 (3)	0.1020 (4)	3.3 (1)
O2D1	0.3056 (4)	0.1208 (4)	0.2487 (6)	6.4 (2)
Fe2 N3	0.500 0.4217 (5)	0.500 0.4366 (5)	0.000 0.0926 (6)	4.39 (5) 4.7 (2)
C31	0.4217 (3)	0.4626 (6)	0.1016 (8)	5.2 (3)
C32	0.2933 (7)	0.4033 (6)	0.1754 (8)	5.7 (3)
C33	0.3853 (7)	0.3419 (6)	0.2097 (8)	4.9 (2)
C34	0.4653 (7)	0.3632 (6)	0.1606 (8)	4.4 (2)
C7 N4	0.5697 (6) 0.6597 (5)	0.3118 (6) 0.4135 (5)	0.1676 (8)	4.2 (2)
C41	0.6397 (3)	0.4133 (3)	0.0804 (6) 0.0043 (8)	4.8 (2) 4.9 (2)
C42	0.7626 (7)	0.3080 (6)	-0.0055(8)	5.0 (3)
C43	0.7136 (7)	0.2672 (6)	0.0601 (8)	5.0 (3)
C44	0.6452 (7)	0.3297 (6)	0.1091 (8)	4.4 (2)
C8	0.7551 (7)	0.4588 (6)	-0.0577 (8)	5.1 (3)
C71 C72	0.6030 (6) 0.5394 (8)	0.2295 (5) 0.1664 (5)	0.2293 (8) 0.2022 (9)	4.2 (2) 5.9 (3)
C73	0.5714 (7)	0.0900 (6)	0.2589 (9)	5.7 (3)
C74	0.6669 (8)	0.0757 (6)	0.3470 (9)	5.9 (3)
C75	0.7315 (8)	0.1350 (7)	0.3746 (8)	5.7 (3)
C76	0.6979 (7)	0.2123 (6)	0.3149 (8)	5.0 (3)
C81 C82	0.8599 (7) 0.9574 (8)	0.4301 (6) 0.4017 (7)	-0.0972 (8) -0.0147 (9)	5.4 (3) 6.3 (3)
C83	1.0538 (8)	0.3761 (7)	-0.053 (1)	7.8 (4)
C84	1.0528 (9)	0.3801 (7)	-0.163 (1)	8.2 (4)
C85	0.9581 (9)	0.4066 (8)	-0.243 (1)	8.7 (4)
C86	0.8586 (8)	0.4324 (7)	-0.209 (1)	7.0 (3)
C1D2 C2D2	0.6984 (7) 0.6294 (7)	0.4653 (6) 0.5389 (6)	0.1859 (8) 0.2080 (8)	4.5 (2) 4.6 (2)
C2D2	0.6668 (8)	0.5389 (8)	0.2080 (8)	6.9 (3)
C4D2	0.7420 (8)	0.5258 (7)	0.4181 (8)	6.2 (3)
C5D2	0.8402 (9)	0.4790 (7)	0.3731 (9)	6.9 (3)
C6D2	0.8056 (8)	0.4348 (7)	0.2623 (8)	5.7 (3)
C7D2	0.682 (1)	0.4622 (9)	0.454 (1)	10.1 (4)
C8D2 O1D2	0.787 (1) 0.5398 (5)	0.5799 (8) 0.5707 (4)	0.519 (1) 0.1381 (6)	8.6 (4) 6.1 (2)
O2D2	0.8724 (6)	0.3732 (5)	0.2376 (7)	8.2 (2)
			en in the form o	

^a Anisotropically refined atoms are given in the form of the isotropic equivalent thermal parameter defined as (4/3)*[a2*B(1,1) + b2*B(2,2)] $+ c2*B(3,3) + ab(\cos \gamma)*B(1,2) + ac(\cos \beta)*B(1,3) + bc(\cos \alpha)*B$ (2,3)].

are significantly longer than the 2.242 (5) Å value found in complex 5a. However, in these mono-N-alkyl derivatives, the

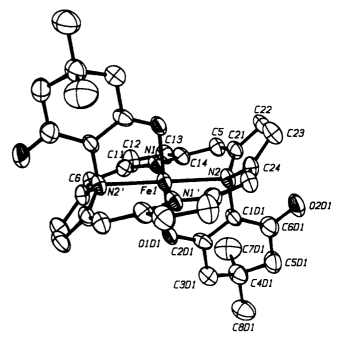


Figure 1. ORTEP plot of one centrosymmetric molecule of complex 5a together with the labeling scheme used. Primed atoms are related to independent atoms by the inversion center. Phenyl rings and hydrogen atoms are omitted for clarity. Ellipsoids are scaled to enclose 30% of the electronic density. The labeling scheme of the second half molecule of complex 5a starts with Fe2, the two pyrrole rings are numbered N3, C31 to C34 and N4, C41 to C44, respectively; C7 and C8 are the methine carbons, and the phenyl carbons are numbered from C71 to C76 and C81 to C86, respectively. The dimedon (X) group starts with C1D2 and goes

metal is always displaced with respect to the plane containing the three nonalkylated porphyrinato nitrogens and also displaced relative to the mean plane containing the alkylated pyrrole nitrogen and the three porphyrinato nitrogens. In contrast, in complex 5a, the iron atom lies in the plane containing the two alkylated and the two nonalkylated nitrogens.

The two Fe-Np bond lengths which are cis to the Fe-N(alkyl) bonds are slightly longer than in all the known six-coordinate low-spin iron(II) porphyrin derivatives (2.000 Å) 16 and even slightly longer than the average bond distance known for the few six-coordinate high-spin iron(II) derivatives (2.063 Å) in which, due to the high-spin state (S = 2) of iron, an expanded core is present.¹⁷ Consequently, bis-N-alkylation such as in complex 5a leads to macrocyclic derivatives in which the metal, the two alkylated nitrogens, and the two nonalkylated nitrogens can form a planar entity with an expanded core relative to normal six-coordinate porphyrin derivatives. Such an expanded core is compatible with the formation of six-coordinate high-spin iron(II)

Although the two individual pyrrole and the two N-alkylated pyrrole rings are almost planar, the mean value of the dihedral angles between their mean planes is 41.3° in the two centrosymmetric molecular units. The mean deviation of the nonalkylated pyrrole ring mean planes relative to the plane containing the four nitrogens and iron (4NFe) is 11.0°, whereas the mean deviation of the N-alkylated pyrrole ring mean planes with respect to the 4NFe plane is as large as 50.3° (the mean value corresponding to the two centrosymmetric entities present in the asymmetric unit). The latter distorsion from planarity is necessary to bring the lone pair of the N-alkylated nitrogens (essentially sp³ hybridized) into an appropriate orientation with iron.

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Table III. Selected Bond Lengths (Å), Angles (deg), and Averages with Their Estimated Standard Deviations for Complex 5a^c

		•		
Fe ₁ -N ₁	2.078 (6)	N_1 -Fe ₁ - N_2	88.7 (2)	
Fe_2-N_3	2.086 (8)	N_3 -Fe ₂ - N_4	89.6 (3)	
Fe_1-N_2	2.261 (5)	N_1 -Fe ₁ - O_1D_1	89.1 (2)	
Fe ₂ -N ₄	2.223 (6)	N_1 -Fe ₂ - O_1D_2	87.2 (3)	
$Fe_1 - O_1D_1$	2.002 (5)	N_2 -Fe ₁ -O ₁ D ₁	84.1 (2)	
Fe ₂ -O ₁ D ₂	1.995 (7)	N_4 -Fe ₂ -O ₁ D ₂	84.6 (3)	
$N_2 - C_1 D_1$	1.479 (10)	, , ,	` '	
$N_4-C_1D_2$	1.523 (11)			

Pyrrole Rings

Rings N	N ₁ and N ₃	Rings N2 and N4
N-C,	1.380 (4)	1.444 (4)
$C_{\alpha} - C_{\beta}$	1.431 (4)	1.418 (4)
$C_{\theta}^{u}-C_{\theta}^{u}$	1.351 (6)	1.372 (6)
$C_{\alpha}^{\nu}-C_{m}^{\nu}$	1.416 (4)	1.382 (4)
C"-N-C"	106.0 (4)	105.6 (4)
$N-C_{\alpha}-C_{\beta}$	109.6 (3)	108.0 (3)
$C_{\alpha}-C_{\beta}-C_{\beta}$	107.3 (3)	109.1 (3)
N-C _a -C _m	124.6 (3)	127.0 (3)

Phenyl Rings 1.492 (5) C-C-C 1.387 (3)

Ligands C_1-C_2/C_1-C_6 1.392 (7) $C_2-C_3/C_3-C_4/C_4-C_5/C_5-C_6$ 1.506 (4) O_1-C_2/O_2-C_6 1.250 (4) Fe- O_1-C_2 112.6 (4)

The mean value of the axial Fe-O bond length in the two molecular units is 1.998 (5) Å. This distance is larger than the Fe-O(C-) bond length present in high-spin Fe(PPIXDME¹³)-(OCH₃) (1.842 (4) Å)¹⁸ but shorter than in the low-spin iron(II) carbonyl adduct Fe(DPIXDME¹³)(CO)(THF) (2.127 (4) Å)¹⁹ and much shorter than in the high-spin iron(II) bis-THF adduct, Fe(TPP)(THF)₂, where the Fe-O bond length is 2.351 (3) Å. 18 Despite the large distorsion of the porphyrin ring only the N-(alkyl)– C_2 bond distance is significantly lengthened (mean value 1.444 (4) Å) relative to the Np– C_α bond length (mean value 1.380 (4) Å) whereas the C_{α} – C_{m} bond is significantly shortened: C_{m} – C_{α} (alkylated pyrrole rings) = 1.382 (4) Å, instead of $C_m - C_\alpha$ (nonalkylated pyrrole rings) = 1.416 (4) Å (Table III). The mean value of the N-C(X group) bond distances of 1.501 (6) Å is similar to those present in other N-alkylated metalloporphyrins. 15

UV-Vis Spectroscopy of Complexes 5. The UV-vis characteristics of complexes 5 in C_6H_6 at 27 °C are reported in Table I. As expected from iron(II)-N-alkylporphyrins, ^{20,21} they all exhibit a red-shifted Soret peak (around 465 nm) with a relatively low molecular extinction coefficient. Three peaks around 560, 620, and 750 nm are also observed in the visible region.

Magnetic Susceptibility and EPR of Complex 5a. The magnetic susceptibility of complex 5a was recorded between 4 and 300 K. Above 40 K, the effective magnetic moment is constant and equal to 5.40 \pm 0.05 μ_B . Between 40 and 4 K, the moment decreases progressively. It is equal to 4.5 μ_B at 4 K. The compound behaves as expected for a quintet spin state $(S = 2)^{2}$. The decrease in the lower temperature range is characteristic of a zero field splitting effect which is usually observed for quintet states arising from high spin ferrous ions in distorted octahedral ligand fields.²³

No EPR transitions are observed in the temperature range

Fable IV. 1H NMR Characteristics of Complexes 3 and

									pher	phenyl groups	S					X moiety	iety	
		S.d.	pyrrole				para			meta			10	ortho	CH,	CH,	and	CH,
complex	2H	2H	2H		2H	2H	2H	4H	_	2H	7	2H 4	4H	4H	2H,	$2H^{\hat{L}}$	+	, Н9
3ab	97.6	82.5			-16.4	7.2	3.5	11.6	9	10.3	٦	6.6	ndç	-l-c	-28.5	6.5	3	-
(M, 1,7)d	(800)	(1100)	(1100)		(1020)	(20)	(50)		•	e		e	•		(2000)		b	
3a-d 30 b.		,				7	3.5	11.1	-	5	.45	7	6.7	4.9				
(x, y)						نه	(25)	(56)	-	_	(56)	-	ь	(34)				
. .	89.2	75.4	70.5		-14.2		•		_	10.3		8.6	pu		28	6.5	o S	-
3b-dg	89.5	75.7	70.5		-14.5													
b(c,1,w)	(20)	(55)	(55)		(55)													
ို့မွ	86	81.8	75		-16.7	9.78	4.98	11.9	6	10.3		10	ndç	<u>-</u>	-29.5	7.5	t	-:
E	93.5	81	8		<u>&</u>	3.5	2.8		6	9.3	ж	8.8	.pu	 	-26.5	7.7	đ	1.5
		4H		4H			4H		8H				4H		4H	4H	+	12H
5		28.4		-42.4			9.0		9				9.	5	35.8	7.6		6.1
b(1,1)		(135)		(350)			(17)		(30)				(190)	(0	(875)	٦		(120)4
S		29.3		-42.8					5.9				6	15	35.8	7.4		6.3
Ş		27.5		-41.8			0.2		5.9				9.5	5	34.6	7.8		6.2

as detected. "Half-width in Hz. "Not measurable because of an overlapping of the signals with another one. $^{1/4}$ H NMK in CH₂Cl₂, signal at 3.5 ppm 8 CH₃-para (6H). "Determined from complex 5a- d_{20} . "CH₃-para (12H). $^{1/4}$ H NMR in HCCl₃, 10% DCCl₃ has been added as internal reference. = not detected. "Half-width in Hz.

^a Numbering as in Figure 1; C_{α} and C_{β} are respectively α and β carbons of pyrrole rings, C_m meso carbons, and C_{phe} quaternary carbon of phenyl groups.

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and refences cited therein.

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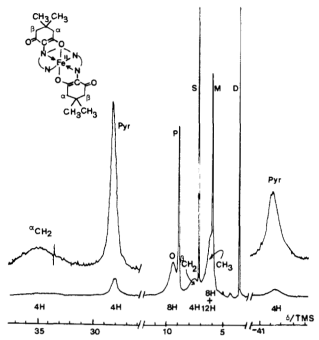


Figure 2. ¹H NMR spectrum of complex 5a (in DCCl₃, 20 °C, δ in ppm from Me₄Si). For the protons of the porphyrin ring: O = ortho, M = meta, P = para, Pyr = pyrrole. A known amount of dioxane (D) has been used as internal standard for proton integration. S = HCCl₃.

4.2-300 K. This fact confirms the quintet state as a ground state, with a zero field splitting larger than the X-band microwave frequency.

¹H NMR Spectra of Complexes 5. The ¹H NMR spectra of complexes 5 are also characteristic of their paramagnetic nature with the large range (36 to -43 ppm) and broad shapes of the signals. The assignments of these signals have been made by comparison of the spectra of complexes 5 with those of their analogues $5a-d_{20}$ and $5b-d_8$ prepared from the tetraarylporphyrin

selectively deuterated on the aryl groups (d_{20}) or on the pyrrole rings (d_8) . The ¹H NMR data are reported in Table IV.

In agreement with the centrosymmetric structure, the pyrrole protons of all complexes 5 appeared as two broad signals (4 H each), one of them appearing downfield around 28 ppm from Me₄Si and the other one upfield around -42 ppm. This assignment was confirmed by the absence of these two signals in the ¹H NMR spectrum of complex 5b-d₈. The para H of phenyl rings of complex 5a which appeared as a sharp signal ($\hat{W}_{1/2} = 17 \text{ Hz}$) at 9.0 ppm (4 H) was unambiguously identified since this signal was missing in the spectra of complexes 5b and 5c (Figure 2). Two others signals (8 H each) around 9 and 6 ppm are present in the spectra of all complexes 5 and are lacking in those of complex 5a-d₂₀. They were respectively assigned to the ortho and meta H of phenyl rings, on the basis of their half-widths. Effectively, the ortho-aryl protons which are closest to the paramagnetic iron center are more efficiently relaxed and then appear as a broad signal ($W_{1/2} = 160 \text{ Hz}$) while the meta-aryl proton signal is sharper $(W_{1/2} = 30 \text{ Hz}).$

The signals of protons of the X moiety were identified by comparison between the spectra of complexes 5a- d_{20} and 5b- d_8 . Only three signals at about 35 (4 H), 7.5 (4 H), and 6.1 ppm (12 H) were present in both spectra and in all complexes 5. On the basis of their integration, the latter signal was assigned to the dimethyl groups and the two others to the methylene groups of the two X moieties. The methylene groups which are closer to the paramagnetic center give a very broad signal ($W_{1/2} = 875$ Hz) at low field (around 35 ppm). The other methylene groups lead to a sharper signal at 7.6 ppm which is in part masked by the meta aryl protons signal.

Electrochemical Oxidation of Complexes 5. The cyclic voltammogram of 5a exhibits a single wave which can be described as a one-electron reversible step with $E_{1/2} = 0.41$ V vs. SCE (Figure 3A). Coulometry at a platinum gauze (controlled potential set to 0.6 V vs. SCE) gives 0.99 as number of electrons exchanged per molecule. Spectroelectrochemistry confirms the reversible transformation of complex 5a into a new red complex 6a. Indeed, upon oxidation of complex 5a at + 0.6 V, the visible spectrum of complex 6a exhibiting bands at 440 and 540 nm²⁴

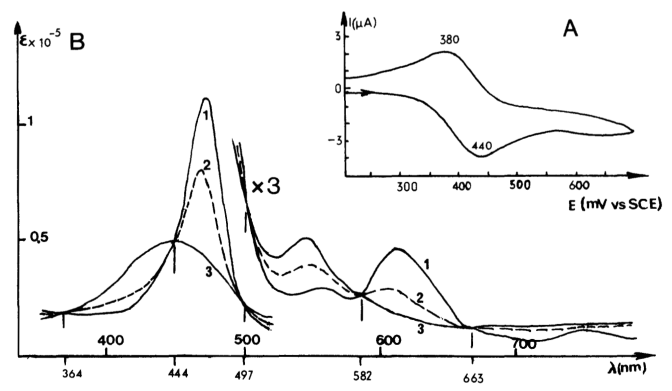


Figure 3. (A) Cyclic voltammogram of complex 5a (1 mM in CH₃CN, 0.2 M LiClO₄) at a glassy carbon electrode. Potential sweep rate 0.2 V/s, (B) spectroelectrochemical oxidation of complex 5a (0.4 mM in CH₃CN, 0.2 M LiClO₄) at a platinum minigrid electrode. Potential: 0.6 V. Spectra 1-3 were obtained after respectively 0, 5, and 10 min.

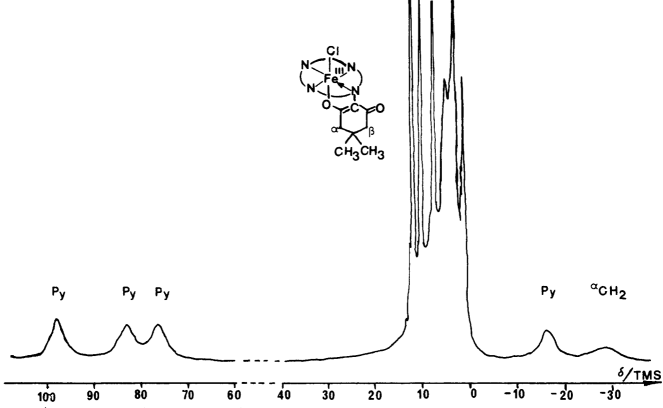


Figure 4. ¹H NMR spectrum of complex 3a (in DCCl₃, 20 °C); Py = pyrrole.

is obtained with five clean isosbestic points at 364, 444, 497, 582, and 663 nm (Figure 3B). Spectrum of complex 5a is regenerated upon reduction of complex 6a at 0 V with the same set of isosbestic points.

Although we did not try to isolate and completely characterize complex 6a, its electrochemical characteristics are compatible with a bismetallocyclic structure like complex 5a but with a Fe^{III} state of the iron (Scheme I). Similar results were obtained from complexes 5b and 5c which led to complexes 6b and 6c, respectively²⁴ (Table I).

II. Reaction of C₆H₅I=X (1) with Iron(III)-Porphyrins: Preparation of Monometallocyclic Iron-Porphyrin Complexes, 3. Although reaction of the iodonium ylide 1 with various Fe^{II} porphyrins has led almost exclusively to the bismetallocyclic Fe^{II} complexes 5 (for porphyrin TPP, TpClPP and TTP), we found that the reaction of Fe^{II}(TMP) with 1 failed to give a similar complex. Even the use of a very large excess of 1 (80 equiv) did not provide a bismetallocyclic Fe^{II} complex analogous to 5a, presumably because of the greater hindrance of the TMP derivatives. In fact, this reaction led to a monometallocyclic N-alkylporphyrin-Fe^{III} complex 3d (after aerobic HCl treatment of the crude reaction mixture) (see below structure evidence).

We then found that such monometallocyclic complexes 3 could be easily obtained in a more general manner upon reaction of 1 with Fe^{III}(porphyrin)(ClO₄) complexes generated in situ from Fe(porphyrin)(Cl) and AgClO₄ in deaerated CH₂Cl₂. For instance, reaction of Fe(TPP)(ClO₄) with 20 equiv of 1 led, after HCl treatment, to a 60% yield of 3a together with minor amounts of 5a (5%). By using a smaller excess of 1 (10 equiv), complex 3a was obtained almost quantitatively within 2 h at 20 °C. A similar procedure applied to Fe^{III}(TpClPP)(ClO₄) and Fe^{III}(TT-P)(ClO₄) led respectively to complexes 3b and 3c (70% yield). Complex 3d was prepared from reaction of 1 with Fe(TMP)-(ClO₄), but in that case an excess of 50 equiv was required to obtain a satisfactory yield (60%).

Structure and Spectral Properties of Complexes 3. The green-brown complexes 3a, 3b, and 3c exhibit UV-vis spectra (Table I) with a red-shifted Soret peak around 456 nm with a relatively small molar extinction coefficient and 4 peaks in the visible region around 525, 580, 655, and 730 nm. The spectrum of complex 3d is similar, but all its peaks are slightly blue-shifted. These spectra are slightly different from those of the classical N-methyl- or N-phenylporphyrin-iron(III) complexes,²¹ but they are similar to that of the only N-alkylporphyrin-iron(III) complex for which such a metallocyclic structure has been very recently proposed.25

Complexes 3 are high-spin Fe^{III} complexes $(S = \frac{5}{2})$ as shown for 3a and 3d by the value of their magnetic moment ($\mu_{eff} = 5.9$ μ_B in HCCl₃ solution at 20 °C) measured by the Evans method.²⁶ Their EPR spectra recorded on the complexes in the powder state at 4.2 K respectively exhibit signals at g = 4.168 and 4.056, each corresponding to the isotropic signal of a high-spin iron(III) complex with rhombic symmetry.

Structures of complexes 3 have been deduced from (i) their elemental analysis, (ii) their mass and ¹H NMR spectra and (iii) the complete transformation of complexes 3a, 3b, and 3c into complexes 5a, 5b, and 5c, respectively, upon reaction of the one-electron reduction products of complexes 3 with 1. Because of the steric hindrance caused by its ortho methyl groups, complex 3d fails to react with compound 1. However, its structure is assumed to be identical with those of other complexes 3 on the basis of its elemental analysis, its mass spectrum, and the great similarity of its UV-vis and ¹H NMR spectra with those of complexes 3a-3c.

The elemental analysis (C, H, N, Cl) of all complexes 3 corresponds to the formula $Fe(porphyrin)(C_8H_{10}O_2)(Cl),H_2O,$ indicating the formal incorporation of only one X moiety into Fe-(porphyrin)Cl and a residual solvation by one water molecule (see Experimental Section).

⁽²⁴⁾ UV-vis (C₆H₆, 27 °C, λ(nm), ε (mM⁻¹·cm⁻¹)) **6a**: 440 (50.2), 540 (17), 580 (sh); **6b**: 440 (68.3), 540 (19.5), 575 (sh); **6c**: 450 (71.5), 543 (22.4), 583 (sh).

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The mass spectra (impact direct inlet) of complexes 3a, 3b, and 3c do not exhibit the molecular peak but only peaks corresponding to the loss of the axial chlorine ligand (i.e., M^+-Cl) and to the loss of the X group (i.e., M^+-Cl-X) (Table I); whereas the mass spectrum of complex 3d (desorption chemical ionization technique, I^- , CH_4) exhibits a molecular cluster peak at m/e 1011 and 1009 (3%) with an isotopic ratio characteristic of the presence of a chlorine atom and fragments at m/e 975 ($M^+ + 1 - Cl$) (100%) and 836 ($M^+ - Cl - X = Fe(TMP)$) (18%). All the mass spectra of complexes 3 exhibit a fragment corresponding to Fe(porphyrin) indicating that the porphyrin ring remains intact in complexes 3 and disfavors any structure which would have derived from an addition of the X carbene moiety on a pyrrole double bond or from its insertion into a C-H pyrrolic or C-aryl simple bond.

The reduction of complexes 3a, 3b, 3c, and 3d by sodium dithionite in a biphasic medium (H_2O, CH_2Cl_2) gives new complexes, respectively 4a, 4b, 4c, and 4d which are stable under these reductive conditions but are rapidly reoxidized to the starting complex in the presence of O_2 and Cl^- . The reversible transformation $3 \rightleftharpoons 4$ is confirmed by cyclic voltammetry in DMF solution $(E_{1/2}$ between -0.3 and -0.1 V vs. $SCE)^{27}$ (Table I). The addition of a methylene chloride solution of 4a, 4b, or 4c to an excess of compound 1 gives immediately and quantitatively the corresponding bismetallocyclic iron(II) complex 5a, 5b, or 5c whereas 4d fails to react with compound 1 (Scheme I).

Taken altogether, these data support the monometallocyclic structure of complexes 3, indicated in Scheme I, where a O-C-C moiety derived from the X group is inserted into a Fe-N bond of the starting ferric porphyrin. The ¹H NMR spectra of complexes 3 are in complete agreement with this structure.

¹H NMR Spectra of Complexes 3. As for complexes 5, the ¹H NMR spectra of complexes 3 are also characteristic of paramagnetic compounds with broad signals appearing over a 130 ppm range. For a complete assignment of the signals it has been necessary to compare the spectra of complexes 3 with those of their analogues 3a-d₂₀ (3a deuterated on the phenyl groups) and 3b-d₈ (3b deuterated on the pyrrole rings) and to study the ²H NMR spectra of the two latter complexes. The ¹H NMR characteristics of complexes 3 are listed in Table IV, and the ¹H NMR spectrum of complex 3a is shown in Figures 4 and 5.

In agreement with the C_s symmetry of complexes 3, their pyrrole protons appear as four signals (2 H each), three of them at very low field, between 98 and 76 ppm downfield from Me₄Si, the fourth one at high field, between -8 and -16 ppm (Figure 4). This assignment, made on the basis of the absence of these signals in the HNMR spectra of complex 3b-d₈, was confirmed by a study of the ²H NMR of complex 3b-d₈ which exhibits only four signals of equal intensity at the same chemical shifts. The para H of the phenyl groups of complex 3a appear as two signals at 7.2 (2 H) and 3.5 (2 H). These two signals are absent in the spectra of complexes 3b and 3d. They are respectively replaced by two signals at 3.5 (2 \times 3 H) and 2.8 (2 \times 3 H) ppm in complex 3d which are assigned to the para methyl protons. The signals of the meta H protons of the phenyl groups appear as a set of three signals (4 H, 2 H, and 2 H) between 11.6 and 8.8 ppm. They are present in the spectra of all complexes 3 and missing in complex 3a-d₂₀.

Because of the large line width of the signals and the presence of other broad signals in the same region, the resonances of ortho H protons could not be directly determined on the ¹H NMR spectra of 3a, 3b, or 3c. However their presence between 1 and 12 ppm was shown by integration of the signals in this region. Similarly, the presence of the signals of the ortho CH₃ groups of complex 3d in the 1-12 ppm region could only be detected by integration of the signals in this region. Finally, the ortho H protons of the aryl groups have been localized only on the ²H NMR spectrum of complex 3a-d₂₀ where they appear as two very broad signals around 7.9 and 4.9 ppm. Since the signal at 7.9 ppm overlaps with those of para ²H, only the half-width of the

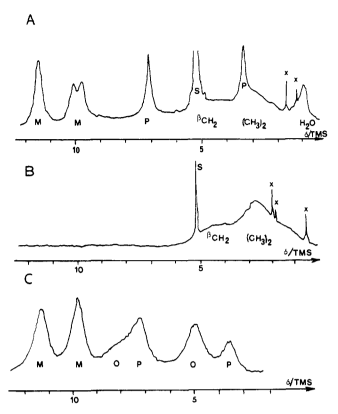


Figure 5. (A) ¹H NMR spectrum of complex 3a (in DCCl₃, 20 °C) (0–12 ppm region). (B) ¹H NMR spectrum of complex 3a- d_{20} (in DCCl₃, 20 °C). (C) ²H NMR spectrum of complex 3a- d_{20} (in D₂CCl₂, 20 °C). O = ortho, M = meta, P = para, S = solvent, x = impurity, H₂O (disappears upon addition of D₂O).

signal at 4.9 ppm could be measured ($W_{1/2} = 34 \text{ Hz}$) (Figure 5C). This would lead to an extrapolated half-width of the corresponding resonances in ¹H NMR around 1400 Hz which would explain why these resonances of the ortho H phenyl protons could not be directly detected in the ¹H NMR spectra of complex 3a (Figure 5A).

As far as the signals of group X are concerned, a very broad resonance ($W_{1/2} = 2000 \text{ Hz}$) was observed around -28 ppm (Figure 4) in the spectra of all complexes 3. This signal (2 H) is assigned to the methylene group of the X moiety which is closest to the paramagnetic center. The protons of the other methylene group and of the two methyl groups appear in the spectrum of complex 3a-d₂₀ as two very broad signals (Figure 5B) between 6.5 and 1 ppm and centered at 4.5 and 2.9 ppm. In the spectra of nondeuterated complexes 3, they are superimposed to those of the ortho H phenyl groups (Figure 5A). The ¹H NMR spectra of complexes 3 are similar to those reported for [Fe^{III}(N-alkyl-TPP)(Cl)|+ cationic complexes.^{21a} They do not allow one to exclude a [FeIII(N-XH-TPP)(Cl)]+ structure for complexes 3 derived from the formula indicated in Scheme I by dissociation of the Fe-O bond and protonation of the oxygen atom. However, conductimetry measurements performed on complex 3a (conductivity of 3a 10⁻³ M in DMF almost equal to that of pure DMF) are completely inconsistent with such a cationic structure. Moreover, the great stability of complexes 3 at room temperature and their EPR signals (g around 4.1) indicative of a rhombic symmetry are in agreement with the bridged structure Fe(N-X-TPP)(Cl) but not with a cationic structure [Fe(N-XH-TPP)(Cl)]+, since previously described [Fe(N-alkyl-TPP)(Cl)]+ complexes, were found stable only below -40 °C and exhibit g = 5.6 and 2.1EPR signals indicative of an axial symmetry.²¹ All these data clearly indicate the bridged structure of Scheme I for complexes

Discussion

The aforementioned results show that the reaction of the io-

⁽²⁷⁾ UV-vis (DMF, 27 °C, λ (m), ϵ (mM⁻¹·cm⁻¹) 4a: 461 (141), 640 (14.7); 4b: 467 (169), 650 (18); 4c: 463 (141), 637 (13.8); 4d: 449 (128), 558 (7.9), 622 (6.2), 673 (3.6).

Scheme II

donium ylide C₆H₅I=X, a carbon equivalent of C₆H₅I=O, with iron(II)- or iron(III)-porphyrins provides an easy access to porphyrin complexes exhibiting a new type of metallocyclic Fe-O-C-C-N structure. Both monometallocyclic and bismetallocyclic complexes could thus be prepared, the iron being either in the ferric or ferrous state.

Although its mechanism is not yet known, the formation of the five-membered F-O-C-C-N metallocycle can be very formally written as an insertion of a O-C=C+ moiety of the carbene X derived from C₆H₅I=X into a Fe-N bond of the starting iron-porphyrin (Scheme II).

The ferric monometallocyclic complexes 3 have been isolated and completely characterized. The reversible passage to the corresponding ferrous complexes has been shown by cyclic voltammetry and spectroelectrochemistry. The latter complexes, 4, have been so far only characterized by UV-vis spectroscopy mainly because of their instability toward dioxygen compared to complexes 3. A more complete study of their structures, spectral characteristics, and reactivity is underway. However, one of their properties particularly relevant to Scheme I which is demonstrated by this study is the fast and complete reaction with $C_6H_5I=X$, leading to the bismetallocyclic complexes 5. On the contrary, complex 3 fails to react with C₆H₅I=X under identical conditions. The ferrous bismetallocyclic complexes 5 have been also completely characterized. They are very stable in aerated solutions, unlike to the ferrous monometallocyclic complexes 4. The ferric bismetallocyclic complexes 6 have been only characterized so far by UV-vis spectroscopy. The reversible passage 5 = 6 has been shown by cyclic voltammetry and spectroelectrochemistry. The fact that the more stable state for the monometallocyclic complexes is the ferric state, whereas the more stable state for the bismetallocyclic complexes is the ferrous state, is in agreement with the redox potentials of the 3 = 4 (-100 to -300 mV) and 6 =5 (+300 to +600 mV) reactions.

The high reactivity of complexes 4 toward either dioxygen or C₆H₅I=X when compared to complexes 3 and the very difficult oxidation of complexes 5 (see redox potentials, Table I) into complexes 6 explain why complexes 3 and 5 were the only final complexes isolated from reactions between iron-porphyrins and C₆H₅I=X. In reactions between Fe^{III}(porphyrin)(ClO₄) and C₆H₅I=X, the ferric complexes 3 are formed almost quantitatively. In reactions performed between Fe^{II}-porphyrins and C₆H₅I=X under anaerobic conditions, it is likely that complex 4 is formed in steady-state concentrations and reacts very rapidly with C₆H₅I=X leading to complex 5 which is almost the only product isolated under these conditions. In the particular case of Fe^{II}(TMP), the proposed intermediate complex 4d would fail to react with C₆H₅I=X because of the steric hindrance caused by the ortho-methyl groups of the meso phenyl substituents. Instead, this intermediate should be rapidly oxidized to complex 3d, explaining why it was the only product isolated from the reaction.

It is noteworthy that complexes 3 and 4 have a different redox behavior when compared to iron-N-alkylporphyrin complexes in general. The latter complexes have been reported to be stable in the ferrous state^{15d} and very unstable in the ferric state²¹ in agreement with their higher redox potentials (from +500 to -60 mV). 15d,21b,28 The much greater reactivity of complexes 4 toward

Scheme III

dioxygen when compared to Cl-Fe^{II}-N-alkyl-TPP complexes^{15d} could be due at least in part to the much more difficult access of iron in the latter where the two axial positions are hindered by the chloride ligand on one side and by the N-R group on the other side. On the contrary, in complexes 4, one axial position is readily accessible (Chart I) explaining its great reactivity not only toward dioxygen but also toward $C_6H_5I=X$.

The aforementioned results completely establish the existence of metallocyclic iron-porphyrin complexes exhibiting a Fe-O-C-C-N structure and give structural data for one of them. They also give spectroscopic data and reactivity characteristics which should be important for the isolation and study of analogous metallocyclic complexes which have been postulated as intermediates in the formation of green porphyrin pigments in the liver of animals treated by alkenes²⁹ (Scheme III). The formation of such green N-substituted porphyrins has been recently described upon model reaction of alk-1-enes with iodosoarenes in the presence of iron-tetraarylporphyrins.^{25,30} The iron complexes which are formed in these model reactions and which lead to the isolated N-substituted porphyrins by acid demetalation have been characterized so far by UV-vis and EPR spectroscopy.25 Their UV-vis characteristics are similar to those of the metallocyclic ferric complexes described in this paper. Moreover, contrary to ferric-N-alkylporphyrin complexes described previously, 21 they are stable at room temperature. 25 Our results showing the exceptional stability of the Fe^{III}-O-C-C-N metallocyclic complexes allow one to understand this property.

Experimental Section

Materials. Compound 1, PhI=X ($X = C_8H_{10}O_2$), was prepared according to ref 14. TPPH2, TTPH2, and TpClPPH2 were synthesized according to Alder's method³¹ and TMPH₂ according to Badger's.³² TPP- d_{20} and TpClPP- d_8 were synthesized according to ref 33. were made chlorine free by Smith's procedure,34 and they were metalated by FeCl₂·4H₂O in DMF.35

Physical Measurements. UV-vis spectra were obtained in toluene or in CH₂Cl₂ by using a Kontron Uvikon 810 or an Aminco DW 2 spectrophotometer.

Mass spectra were recorded on a VG 70-250 double-focussing in-

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strument (from VG Analytica, Manchester, UK) equipped with a fastatom bombardment gun (from Ion Tech Ldt, Teddindton, UK). The scanning acquisition parameters were the following: accelerating voltage, 4 KV; scan time, 15 s/decade; interscan time, 2 s. The FAB gun was operated with xenon (N45, Air Liquide, France) at 7.5 KV and 1.2 mA. The apparatus was calibrated with cesium iodide, and different matrices were used: glycerol and thioglycerol or dimethyl sulfoxide. All chemical reagents used were of the best available quality.

Elemental analyses were done by the Service de Microanalyse du C.N.R.S. at Gif sur Yvette.

¹H NMR spectra for complexes 10⁻² M in CDCl₃ or CD₂Cl₂ were run at 20 °C on a Bruker WH 250 spectrometer operating at 250 MHz; chemical shifts are reported in ppm downfield of Me₄Si (sweep width 5000 Hz, 256-1024 scans, 60 pulses, 3.0 Hz/pt resolution, 32K point memory blocks, acquisition time 0.32 s, repetition delay 1 s). ²H NMR spectra were run at 38.4 MHz (sweep width 5680 Hz, 20000-30000 scans, 60 pulses, 0.71 Hz/pt resolution, 16K point memory blocks, acquisition time 1.4 s with broad band of ¹H decoupling, repetition delay 1 s). DCCl₃ or D₂CCl₂ (10%) was used as internal reference

Magnetic susceptibility measurements were carried out with a Faraday type magnetometer equipped with a continuous flow cryostat in the temperature range 4.2-300 K. The polycrystalline powder samples weighed about 3 mg. The applied magnetic field was in the range 0.1-0.7 T. The independence of the susceptibility against the magnetic field was checked at room temperature. Mercuritetrakis(thiocyanato)cobaltate-(III) was used as a standard susceptibility. The magnetic data were corrected for the molecular diamagnetism estimated as -860×10^{-6} cm3-mol-1

The EPR properties were checked with a Bruker ER 200 D spectrometer equipped with a continuous flow cryostat and working in Xband, in the temperature range 4.2-300 K.

Cyclic Voltammetry and Coulometry. Experiments were carried out with a Princeton Applied Research Model 173 potentiostat, monitored with an Apple //e microcomputer and a P.A.R. Model 276 GPIB interface. A three electrode system was used with a platinum disk or a glassy carbon disk working electrode, a platinum wire counter electrode, and a saturated calomel electrode (filled with NaCl instead of KCl to prevent crystallization of KClO₄). The reference electrode was separated from the bulk of the solution by a fritted glass bridge and filled with the solvent and supporting electrolyte at appropriate concentration. For coulometry experiments the counter electrode was separated by a bridge with a Nafion membrane, filled with the same supporting electrolyte solution. The spectroelectrochemical device (analogue to Lexa's³⁶) consisted in a three electrodes cell, with as compartment of electrolysis a quartz cell (optical length, 0.5 mm; inner volume, 0.08 mL). working electrode was a platinum minigrid. Except for the oxidation of 5a, all solutions were deaerated by an argon stream before experiments, then protected by a blanket of argon. Dimethylformamide was distilled under reduced pressure; acetonitrile (quality: HPLC grade S), purchased from Rathburn Chem. Limited, and reagent grade quality LiClO4 and LiCl, purchased from Fluka Co., were used without further purification.

X-ray Analysis. Suitable single crystals of 5a were obtained by slow evaporation of solvents from CH2Cl2-pentane solutions at room temperature.

A systematic search in reciprocal space using a Philips PW 1100/16 automatic diffractometer showed that crystals of 5a belong to the triclinic system. The unit-cell dimensions and their standard deviations were obtained and refined at room temperature with Cu K_{α} radiation (λ = 1.5405 Å) by using 25 carefully selected reflections and the standard Philips software. Final results for C₆₀H₄₈N₄O₄Fe were the following: mol wt 944.9, a = 12.785 (3) Å, b = 15.963 (4) Å, c = 12.104 (3) Å, $\alpha = 91.77$ (2)°, $\beta = 105.46$ (2)°, $\gamma = 78.71$ (2)°, V = 2334 Å³, Z = 2, $d_{\text{calcd}} = 1.344 \text{ g} \cdot \text{cm}^{-3}, d_{\text{obsd}} = 1.32 \pm 0.02 \text{ g} \cdot \text{cm}^{-3}, \mu = 30.11 \text{ cm}^{-1}, F_{\infty}$ = 988, space group P1.

A parallelepipedic crystal of 0.32 × 0.18 × 0.04 mm was sealed in a Lindemann glass capillary and mounted on a rotation free goniometer head. All quantitative data were obtained from a Philips PW 1100/16 four circle automatic diffractometer, controlled by a P 852 computer, by using a graphite monochromator and standard software. The vertical and horizontal apertures in front of the scintillation counter were adjusted so as to minimize the background counts without loss of net peak intensity at the 2σ level. The total scan width in the $\theta/2\theta$ flying step-scan used was $\Delta\theta = 0.9 + 0.143$ tg (θ) deg with a step width of 0.04 deg and a scan speed of 0.015 deg·s⁻¹. A total of 6529 hkl, $h\bar{k}l$, $h\bar{k}l$, and $h\bar{k}l$ reflections were recorded (5° $< \theta <$ 57°). The resulting data set was transferred to a PDP 11/60 computer, and for all subsequent computations with the

exception of a local data-reduction program, the Enraf-Nonius SDP/V18 package was used. $^{\rm 37}$

Three standard reflections measured every hour during the entire data-collection period showed no significant trend. The raw step-scan data were converted to intensities by using the Lehmann-Larsen method³⁸ and then corrected for Lorentz polarization and absorption factors, the latter computer by the numerical integration method of Busing and Levy³⁹ (transmission factors between 0.43 and 0.88). A unique data set of 3255 reflections having $(I) > 3\sigma(I)$ was used for determining and refining the structure.

The structure was solved by using the heavy atom method. After refinement of the heavy atoms, a difference-Fourier map revealed maxima of residual electronic density close to the positions expected for hydrogen atoms; they were introduced in structure factor calculations by their computed coordinates (C-H = 0.95 Å) and isotropic temperature factors of $B_{eqv}(C)$ + 1 Å² but not refined. Full least-squares refinement converged to R(F) = 0.076 and Rw(F) = 0.095 ($\sigma(F^2) = (\delta_{count}^2 + (pI)^2)$). The unit weight observation was 1.82 for p = 0.07. A final difference map revealed no significant maxima. The scattering factors coefficients and anomalous dispersion coefficients come respectively from ref 40 and

General Procedure for Preparation of Complexes 3: Fe(P)(C₈H₁₀-O₂)(Cl). To a deaerated solution of 0.2 mmol of Fe¹¹¹(porphyrin)(Cl) in 80 mL of CH₂Cl₂ was added 0.3 mmol of anhydrous AgClO₄. The solution was stirred under argon for 0.5 h at room temperature and then added to 2 mmol of compound 1 and 0.3 mmol of AgClO4 previously deaerated by a bubbling of argon. After stirring for 2 h at 20 °C, the solution was filtered and stirred during 0.25 h in the presence of 5 mL of concentrated HCl dissolved in 20 mL of CH₃OH. The organic layer was washed with water and dried over Na_2SO_4 . After filtration and evaporation of solvents, the crude product was purified by a SiO₂ column chromatography (CH₂Cl₂/ethyl acetate, 95:5 as eluent) and precipitated by addition of pentane to a CH₂Cl₂ solution.

Complexes 3a, 3b, and 3c were obtained as brown powders in 70-90% yield. Complex 3d which did not crystallize even in pure pentane was dissolved in C₆H₆ and lyophilized under high vacuum (60% yield). Complex 3a $Fe(TPP)(C_8H_{10}O_2)(Cl)\cdot H_2O$: Anal. Calcd (found) for C₅₂H₄₀N₄FeClO₃: C, 72.60 (72.39); H, 4.65 (4.42); N, 6.51 (6.86); Cl, 4.13 (4.50). Complex 3b Fe(TpClPP)($C_8H_{10}O_2$)(Cl)· H_2O : Anal. Calcd (found) for $C_{52}H_{34}N_4FeCl_5O_3$: C, 62.55 (62.47); H, 3.61 (3.59); N, 5.72 (6.01); Cl, 17.80 (18.00). Complex **3c** Fe(TTP)($C_8H_{10}O_2$)(Cl)· H_2O : Anal. Calcd (found) for $C_{56}H_{48}N_4FeClO_3$: C, 73.40 (72.95); H, 5.28 (5.17); N, 6.11 (6.44); Cl, 3.87 (3.92). Complex **3d** Fe- $(TMP)(C_8H_{10}O_2)(Cl)\cdot H_2O$: Anal. Calcd (found) for $C_{64}H_{64}N_4FeClO_3$: C, 74.74 (74.80); H, 6.23 (6.44); N, 5.45 (5.05); Cl, 3.45 (2.94)

General Procedure for Preparation of Complexes 5: Fe(P)(C₈H₁₀O₂)₂ (1) from Fe^{III}(porphyrin)(Cl). To a deaerated solution of 0.6 mmol of Fe^{III}(porphyrin)(Cl) in 70 mL of C₆H₆ or CH₂Cl₂ was added 25 mL of a saturated aqueous solution of S₂O₄Na₂. The reaction mixture was stirred, under argon, for 0.25 h during which time the solution changed from brown to red. An electronic spectrum of a sample (diluted in C₆H₆) was recorded to confirm the reduction of Fe^{III}(porphyrin)(Cl) to Fe^{II}-(porphyrin). The organic layer was added to 1.2 mmol of complex 1 previously deaerated by bubbling argon and stirred for 1 h at 50 °C. The red color turned progressively green. After solvent evaporation, the crude product was purified by a column chromatography (SiO₂, CH₂Cl₂/ CH₃COCH₃, 8:2) and crystallized from a CH₂Cl₂-pentane mixture. Complexes 5 were obtained as green crystals in excellent yields (>95%).

(2) From Complexes 3. To a deaerated solution of 0.1 mmol of complex 3a, 3b, or 3c in 30 mL of C₆H₆ was added 10 mL of a saturated aqueous solution of Na₂S₂O₄. The reaction mixture immediately turned light green. The organic layer was added to 0.4 mmol of compound 1 previously deaerated, and the color immediately turned dark green. The electronic spectrum of a sample diluted in C₆H₆ showed the characteristic spectra of the corresponding complexes 5. After a similar workup as in method 1, complexes 5a, 5b, or 5c were each obtained in quantitative yield. Complex 5a Fe(TPP)(C₈H₁₀O₂)₂·2H₂O: Anal. Calcd (found) for $C_{60}H_{52}N_4FeO_6$: C, 73.43 (73.54); H, 5.31 (5.21); N, 5.71 (5.72). Complex 5b Fe(TpClPP)(C₈H₁₀O₂)₂·2H₂O: Anal. Calcd (found) for C₆₀H₄₈N₄FeCl₄O₆: C, 64.40 (64.66); H, 4.29 (4.49); N, 5.01 (5.05); Cl,

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12.70 (12.96). Complexes **5c** Fe(TTP)($C_8H_{10}O_2$)₂·2 H_2O : Anal. Calcd (found) for $C_{64}H_{60}N_4$ FeO₆: 74.12 (74.92); H, 5.83 (5.70); N, 5.40 (5.51).

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Supplementary Material Available: Concerning the X-ray

structure of complex 5a: Table V, thermal parameters for non-hydrogen atoms; Table VI, positional parameters for hydrogen atoms; Table VIII, bond distances; Table IX, bond angles; Table X, least-squares mean planes of interest (13 pages); Table VII, calculated and observed structure factor amplitudes (×10) for all observed reflections (14 pages). Ordering information is given on any current masthead page.

A New Inorganic Ring System: Planar Fe₃(μ_2 -SR)₃ in [Fe₃(SR)₃X₆]³—Synthesis, Structures, and Solution Conformation and Equilibrium

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Abstract: Anaerobic reaction of equimolar FeX2 and RS- in acetonitrile affords the trinuclear complexes [Fe3(SR)3X6]3- (with X = Cl and R = Ph (4a), p-tolyl (4b), 2,6-Me₂C₆H₃ (4d), and X = Br, R = Ph (4c)) isolated as Et_4N^+ salts. These species constitute the fourth known type of discrete Fe(II) thiolates, the others being $[Fe(SR)_4]^{2^-}$, $[Fe_2(SR)_6]^{2^-}$, and $[Fe_4(SR)_{10}]^{2^-}$. The following crystallographic results were obtained (crystal system; space group; cell parameters; Z; unique data $(I > 3\sigma(I))$; R (%)). (Et₄N)₃ (4a): monoclinic; C_2/c ; a = 24.500 (6) Å, b = 13.488 (3) Å, c = 18.217 (5) Å, $\beta = 110.68$ (2)°; 4; 2679; 4.9. (Et₄N)₃ (**4d**)-3MeCN: monoclinic: $P2_1/n$; a = 15.07 (1) Å, b = 22.38 (3) Å, c = 20.46 (3) Å, $\beta = 91.5$ (1)°; 4, 3311; 9.7. (Et₄N)₃ (**4c**)-MeCN: hexagonal, $P\overline{6}2c$; a = b = 13.970 (4) Å, c = 18.274 (3) Å; 2; 890; 5.7. The reaction system FeBr₂ + NaSCH₂Ph in acetonitrile yields a different type of compound with the composition (Et₄N)₂[Fe₄(SCH₂Ph)₆Br₄] (5): triclinic; $P\bar{1}$; a = 12.874 (2) Å, b = 13.949 (3) Å, c = 43.272 (8) Å, $\alpha = 94.85$ (2)°, $\beta = 94.07$ (2)°, $\gamma = 114.98$ (1)°; 4; 7402; 6.7. Anions of type 4 contain a new type of inorganic ring system, $Fe_3(\mu_2-SR)_3$, distinguished by its nearly or exactly planar conformation and unusually large Fe-S-Fe angles of ~140°, the largest values observed for this angle type. The phenyl groups in 4a and 4c (which has imposed C_{3h} symmetry) are coplanar with the Fe₃S₃ rings. In 4d these groups are rotated by $\sim 90^{\circ}$ owing to the steric effects of the methyl groups. Distorted tetrahedral coordination at each Fe(II) atom is completed by two halide ligands. The planar structure is to be contrasted with the chair-type conformation of the rings in 5 and other species with adamantane-like stereochemistry. Ring conformations influence the extent of antiferromagnetic interactions in the solid state. Magnetic susceptibility data of planar 4a are fit well by two -J values at 19-22 cm⁻¹. For [Fe₄(SR)₁₀]²⁻, containing adamantane-like $Fe_4(\mu_2-SR)_6$ cages with $Fe_3(\mu_2-SR)_3$ rings in the chair conformation, best fits of the magnetic data give one -J value at 32 or 36 cm⁻¹ and one or two others at 42-66 cm⁻¹ depending on the coupling symmetry. MO calculations at the extended Hückel level on [Fe₃(SH)₃Cl₆]³⁻ suggest that the inherently stable conformation of the ring is planar. This structure is set in large measure by the stereochemical preference of the S-Fe-S angle and by transannular nonbonded repulsions. The decrease in Cl···Cl distances from 6.6 to 3.9 Å in changing from the planar to the chair conformation emphasizes the latter effect. In acetonitrile the equilibrium $2[Fe_3(SR)_3X_6]^{3-} \rightleftharpoons [Fe_4(SR)_6X_4]^{2-} + 2[FeCl_4]^{2-}$ has been approached in both directions and is shifted to the left with decreasing temperature. At $\lesssim 240 \text{ K}$ the formation of $[\text{Fe}_3(\text{SR})_3 \text{X}_6]^{3-}$ is essentially complete. In this temperature range the magnetic moment per Fe atom (μ_{Fe}) is 4.1-4.2 μ_{B} for **4a** in solution and as its crystalline Et₄N⁺ salt. This excellent agreement establishes a planar ring conformation in solution. Thus the results of three crystal structures, MO calculations, and solution and solid state magnetic measurements collectively demonstrate that the near or exact planarity of the Fe_3S_3 cycles in $[Fe_3(SR)_3X_6]^{3-}$ is an intrinsic property.

Previous investigations in this laboratory have demonstrated reactions 1-4 in Figure 1, affording the complexes $[Fe(SR)_4]^{2-}$ (1), $[Fe_2(SR)_6]^{2-}$ (2), $[Fe_4(SR)_{10}]^{2-}$ (3), and $[Fe_3(SR)_3Cl_6]^{3-}$ (4).²⁻⁷ Several additional examples of 2 have been prepared by

others.⁸ The set 1-4, containing species of varying nuclearity based on tetrahedral Fe(II) coordination, has a number of significant aspects. In reactions with elemental sulfur, 1-3 are precursors to a variety of Fe-S-SR clusters.^{2,4,5,9} These same complexes serve as structural prototypes of related thiolate complexes formed by M(II) ions with tetrahedral stereochemical preference. An expanding number of such complexes have been prepared and structurally characterized.^{10,11} Complex 4a (R =

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