

Formation of heterobinuclear μ -nitrido complexes with a Mn-N–Fe moiety by reaction of nitrido(octaphenyltetraazaporphyrinato)manganese(V) with iron(III) porphyrins

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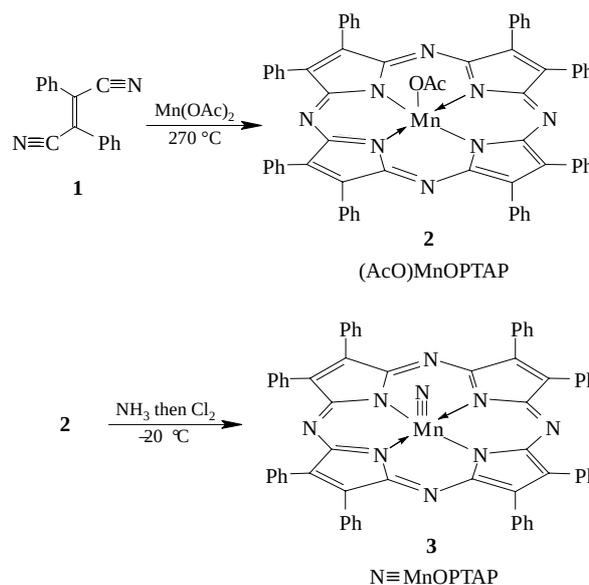
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Nitrido(octaphenyltetraazaporphyrinato)manganese(V) ($N\equiv MnOPTAP$) obtained by treatment of chlorine on acido(octaphenyltetraazaporphyrinato)manganese(III) [(X)MnOPTAP; X = Cl, AcO, HSO₄] in chloroform solution saturated with ammonia, reacts with acido(octaphenyltetraazaporphyrinato)iron(III) [(X)FeOPTAP; X = Cl, Br, AcO] forming the heterometallic μ -nitrido complex μ -(MnNFe)(OPTAP)₂; μ -nitrido complexes with a dissimilar porphyrin ligand on each metal have also been prepared.

Binuclear μ -nitrido iron complexes of porphyrins,¹ tetraazaporphyrins² and phthalocyanine^{3,4} can be easily obtained by thermolysis of the corresponding azidoiron(III) complexes. Heating of azido(tetraphenylporphyrinato)iron(III) [(N₃)FeTPP] in the presence of iron or ruthenium phthalocyanines (FePc, RuPc) results in the formation of mixed binuclear μ -nitrido complexes (TPP)Fe(μ -N)Fe(Pc) and (TPP)Fe(μ -N)Ru(Pc).⁵ It was supposed^{1,6} that nitrido(tetraphenylporphyrinato)iron(V) ($N\equiv FeTPP$) is a reactive intermediate in the formation of μ -N(FeTPP)₂ from (N₃)FeTPP. However, due to the extremely high instability of the nitridoiron(V) complexes [$N\equiv FeTPP$ was detected⁷ as a photolysis product of (N₃)FeTPP only below 150 K] their role in μ -nitrido complex formation has not received any direct confirmation as yet. Unlike nitridoiron(V) complexes nitridomanganese(V) complexes of porphyrins⁸ and phthalocyanines⁹ are very stable. In order to throw some light on the mechanism of the μ -nitrido complex formation, we have obtained stable nitrido(octaphenyltetraazaporphyrinato)manganese(V) ($N\equiv MnOPTAP$ **3**; Scheme 1) and attempted to use it in the synthesis of the heterometallic μ -nitrido complexes (Scheme 2).

Melting of (*E*)-1,2-diphenyl-1,2-dicyanoethylene **1** with anhydrous manganese(II) acetate in a 1:1 molar ratio at 270 °C gave acetato(octaphenyltetraazaporphyrinato)manganese(III) [(AcO)MnOPTAP **2**; yield 85%].[†] $N\equiv MnOPTAP$ **3** was obtained by bubbling chlorine through a cooled (-20 °C) solution of **2** in chloroform saturated with gaseous ammonia (yield 61%).[‡] Other acidomanganese(III) complexes (X)MnOPTAP (e.g. X = Cl, HSO₄) can also be used as a starting material. Complex **2** can bind ammonia as an axial ligand forming hexacoordinated ammine complexes of manganese(III), (AcO)(H₃N)MnOPTAP, or on more prolonged exposition of manganese(II), (H₃N)₂MnOPTAP. Coordination of ammonia to **2** and subsequent conversion of the ammine complexes to $N\equiv MnOPTAP$ under action of chlorine can be followed by UV/VIS spectroscopy (Figure 1). Whereas the spectrum of (AcO)(H₃N)MnOPTAP is almost identical with that of **2** [Figure 1(a)], additional characteristic bands appear at 830 and 886 nm for (H₃N)₂MnOPTAP [Figure 1(b)]. These ammine complexes can be chlorinated to give, we suppose, unstable intermediate complexes containing *N,N*-dichloro-



Scheme 1

amidomanganese(III), (Cl₂N)MnOPTAP, or *N*-chloronitrenomanganese(IV), ClN=MnOPTAP, which easily split off chlorine forming stable nitridomanganese(V) complex **3** [Figure 1(c)]. In the absence of ammonia complex **2** is oxidized irreversibly with chlorine producing colourless products. A substantial hypsochromical shift of the $\pi \rightarrow \pi^*$ transitions of the macrocycle observed for **3** as compared with **2** is in agreement with strengthening of the π -donation effect expected for d² complexes such as $N\equiv MnOPTAP$.

The structure of **3** is confirmed by the presence in the mass spectrum of a molecular ion peak of $N\equiv MnOPTAP^+$ at $m/z = 990.4$ and by the appearance of the Mn \equiv N stretching vibration as a weak band at 1054 cm⁻¹ in the IR spectrum and as a medium-strong line at 1058 cm⁻¹ in the resonance Raman spectrum.

The heterometallic μ -nitrido complex **5**, μ -(MnNFe)(OPTAP)₂, was obtained by reaction of the nitridomanganese(V) complex **3** with chloroiron(III) complex **4**, (Cl)FeOPTAP,¹⁰ in a boiling benzene solution (**3**:**4** = 1:1.5 molar ratio) (Scheme 2). The μ -nitrido complex **5** was chromatographically purified and separated from an admixture of μ -oxodimer O(FeOPTAP)₂, formed partially from **4** (neutral Al₂O₃, eluent CHCl₃, yield of **5** 23%).[§] The Mn/Fe ratio determined by a flame photometry

[†] Spectral data for **2**: Found (%): C, 76.28; H, 4.54; N, 10.49. Calc. for C₆₆H₄₃N₈O₂Mn (%): C, 76.59; H, 4.19; N, 10.83. UV/VIS [CHCl₃, λ_{max} /nm (log ϵ): 285 (4.66), 331sh, 414 (4.56), 475 (4.39), 613sh, 665 (4.60). Camenzind and Hill¹¹ have reported the preparation of octaphenyltetraazaporphyrinatomanganese(II) from **1** and manganese powder with low yield (5.7%); no spectral data have been reported for this compound.

[‡] Spectral data for **3**: Found (%): C, 77.76; H, 4.18; N, 12.55. Calc. for C₆₄H₄₀N₉Mn (%): C, 77.65; H, 4.07; N, 12.73. FD-MS m/z : $N\equiv MnOPTAP^+$ (990.4, 12%); MnOPTAP⁺ (975.3, 100%). UV/VIS [CHCl₃, λ_{max} /nm (log ϵ): 270sh, 345sh, 361 (4.65), 447 (4.22), 577 (4.12), 601sh, 629 (4.91).

[§] Spectral data for **5**: Found (%): C, 77.84; H, 3.95; N, 12.03; Mn, 2.72; Fe, 2.6. Calc. for C₁₂₈H₈₀N₁₇MnFe (%): C, 78.16; H, 4.10; N, 12.11; Mn, 2.79; Fe, 2.84. FD-MS m/z : $N\equiv MnOPTAP^+$ (990.4, 2.9%), FeOPTAP⁺ (976.3, 89%), MnOPTAP⁺ (975.3, 100%). UV/VIS [CHCl₃, λ_{max} /nm (log ϵ): 345 (4.81), 437sh, 583 (4.55), 642 (4.66).

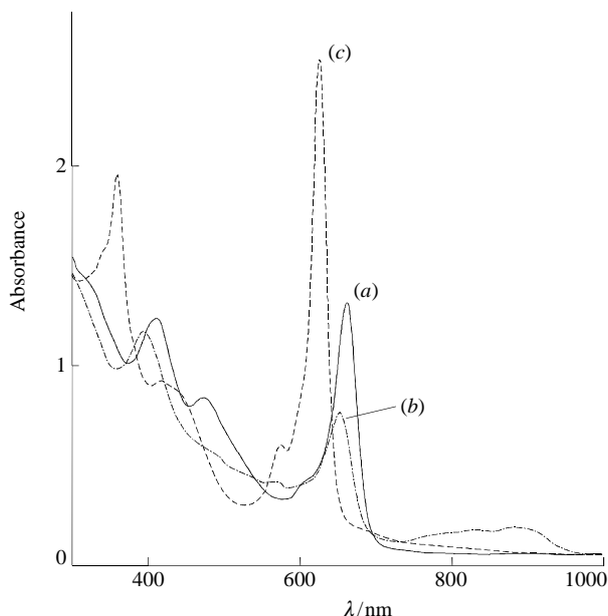


Figure 1 UV/VIS spectra of (a) (AcO)MnOPTAP in chloroform (3.5×10^{-5} M); (b) after saturation with ammonia at -20 °C; (c) after bubbling of chlorine.

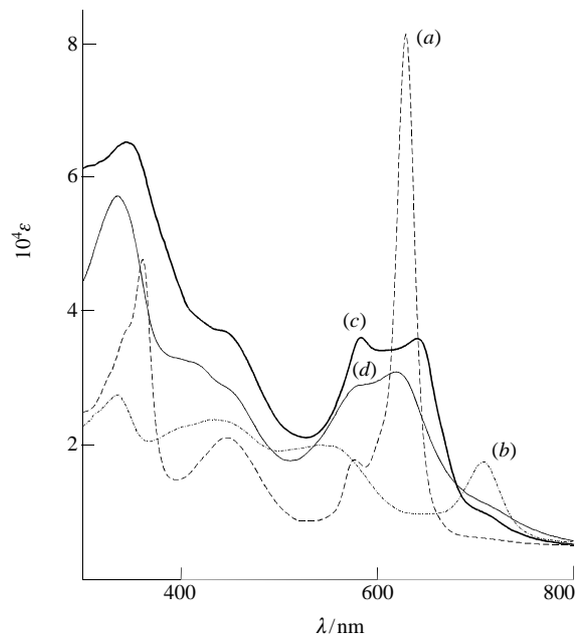


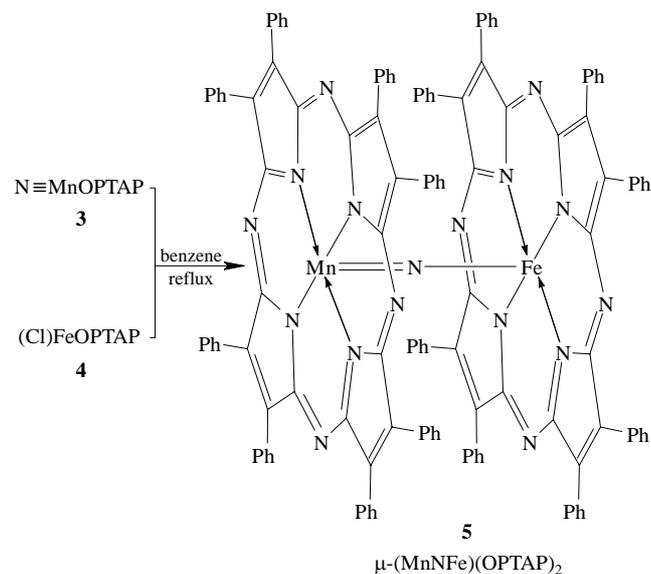
Figure 2 UV/VIS spectra of (a) $N \equiv MnOPTAP$; (b) $(Cl)FeOPTAP$; (c) $\mu-(MnNFe)(OPTAP)_2$ and (d) $\mu-N(FeOPTAP)_2$ in chloroform.

method and the CHN elemental analysis data for **5** are in reasonable agreement with the proposed formula, $\mu-(MnNFe)(OPTAP)_2$. The mass spectrum of **5** obtained by a field desorption method contains mass peaks (in m/z) corresponding to the monomer constituents of the mixed μ -nitrido complex ($N \equiv MnOPTAP^+$ 990.4; $MnOPTAP^+$ 975.3; $FeOPTAP^+$ 976.3), but no molecular ion peak expected for $C_{128}H_{80}N_{17}MnFe$ at $m/z = 1965.6$ has been detected.

Formation of **5** can be easily monitored by UV/VIS spectroscopy: characteristic absorption bands of the initial complexes **3** (629 nm) and **4** (710 nm) disappear and a broad doublet (642, 583 nm) appears. Such splitting of the Q-band, being a result of excitonic interactions of the adjacent π -systems, is characteristic of binuclear single atom bridged complexes. The UV/VIS spectrum of the mixed Mn–Fe μ -nitrido complex **5** [$\mu-(MnNFe)(OPTAP)_2$, Figure 2(c)] is similar to the spectrum of the homobinuclear μ -nitridodiiron complex [$\mu-N(FeOPTAP)_2$, Figure 2(d)], but the maxima of the Q-band envelope are bathochromically shifted. No metal-axial ligand stretching vibrations characteristic of the initial complexes **3** and **4**

($\nu_{Mn \equiv N} = 1054 \text{ cm}^{-1}$ for **3** and $\nu_{Fe-C1} = 310 \text{ cm}^{-1}$ for **4**) are present in the IR spectrum of **5** and a medium-weak band at 918 cm^{-1} appears likely to be associated with the MnNFe bridge ($FeNFe$ absorbs at 920 cm^{-1}). The dinuclear identity of **5** has also been confirmed by its reaction with acids HX (HX = HCl, H_2SO_4 , AcOH, CCl_3COOH) which causes decomposition to the mononuclear complexes of Mn and Fe [$(X)MnOPTAP$ and/or $N \equiv MnOPTAP$ and $(X)FeOPTAP$], each of which can be separated by thin-layer chromatography and identified by UV/VIS spectroscopy. It has been verified that reaction of $N \equiv MnOPTAP$ with bromoiron(III), acetatoiron(III) and μ -oxodiiron(III) derivatives of octaphenyltetraazaporphine [$(Br)FeOPTAP$, $(AcO)FeOPTAP$ and $\mu-O(FeOPTAP)_2$] also leads to **5**. Mixed Mn–Fe μ -nitrido species containing dissimilar macrocyclic ligands bound to each metal can be obtained by coupling of the corresponding nitridomanganese(V) and iron(III) complexes, thus reaction of $N \equiv MnOPTAP$ with $(AcO)FeTPP$ gives $(OPTAP)Mn(\mu-N)Fe(TPP)$.

Whether the $\mu-(MnNFe)$ bridge has an asymmetrical bond distribution ($Mn^{III}N=Fe^{IV}$ or $Mn^{IV}=N-Fe^{III}$ as shown in Scheme 2) or its structure is more symmetrical ($Mn \equiv N \equiv Fe$) similar to the $(Fe \equiv N \equiv Fe)$ bridge in $\mu-N(FeOPTAP)_2$ is not yet clear. Physicochemical properties of mixed Mn–Fe μ -nitrido complexes of porphyrins as well as details of the mechanism of their formation are presently under investigation.



Scheme 2

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