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Spectroscopic Study of the Dimerization Process of Iron Protoporphyrin IX

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The commercial protoporphyrin IX, iron-ferritoporphyrin IX-chloride and synthesized iron porphyrin μ -oxo-dimers were examined by UV/vis absorption and fluorescence, Fourier transformed infrared spectroscopy, resonance Raman, X-ray absorption, Mössbauer spectroscopy and SQUID. The evidence of Fe-O-Fe antiferromagnetic coupling concluded from SQUID and Mössbauer in the case of samples containing dimerized forms confirmed the presence of the oxo-bridges. In this paper the results of UV/vis, fluorescence, Fourier transform infrared FTIR and Raman spectroscopies are reported and discussed. The study is based on the comparison of the free-base protoporphyrin IX, Fe-PPIX-Cl and the synthesized dimerized specimen. The vibrational modes in two energy regions i.e. 330–650 cm^{-1} and 750–900 cm^{-1} , reportedly characteristic of the existence of Fe–O–Fe bridges, are discussed. A significant photoluminescence emission, strongly Stokes shifted from the Soret band, absent in the protoporphyrin IX and the iron-ferritoporphyrin IX-chloride, is observed. The strong Stokes shift and the mismatch of the excitation spectrum to the Soret band suggest that it does not have origin in the de-excitation of the porphyrin moiety and that it could have origin in an Fe-O-Fe charge transfer state.

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1. Introduction

Metalloporphyrins, which contain Fe, Mg, Mn, Co, Cu, Ni or Zn suitably nested between the four nitrogens of pyrrole rings play an important role in biological systems as functional groups of heme proteins, oxidases, chlorophylls and cobalamins [1–3]. Porphyrin derivatives have also recently attracted attention because of potential clinical applications [4, 5]. As photosensitizing agents they have proven, for example, to be very useful photosensitizers in the photodynamic therapy (PDT) of cancer [6, 7]. Dimerization and oligomerization of iron porphyrins have been studied in various model systems for more than 30 years. However, the mechanism of interaction between the haem centers remains unclear, especially in the most important porphyrin IX. The most often investigated systems are μ -oxo-bridged dimers of iron porphyrins. The schematic structure of the iron porphyrin molecule and one of its possible dimer forms is shown in Fig. 1.

Iron porphyrin μ -oxo-dimers were synthesized according to the modified Adler procedure. The results of a study of the local dynamical properties of the iron atoms in the porphyrin monomers and μ -oxo-dimers have been reported previously [8]. The strong antiferromagnetic behavior observed in both Mössbauer and SQUID measurements confirms a strong coupling between iron centers characteristic of dimerization. In this study the characterization of the material using UV/visible and mid

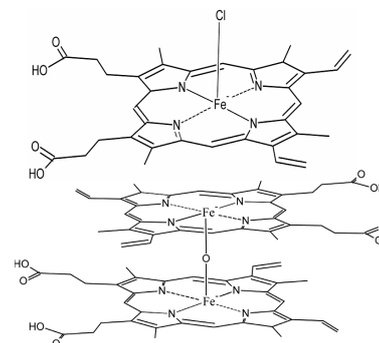


Fig. 1. Molecular structure of the iron porphyrin monomer (top) and iron porphyrin dimer (bottom).

infrared absorption, fluorescence, and Raman spectroscopies has been carried out. In all cases the spectra are referenced to those of the protoporphyrin and the metal monomer.

2. Experimental

Iron porphyrin dimers were prepared by insertion of iron atoms from $^{57}\text{FeCl}_3$ into protoporphyrin IX (PPIX) (Sigma) and finally precipitated from water-*N,N*-dimethylformamide (DMF) solution, according to the modified Adler procedure [9, 10].

According to some modifications during the preparation, final samples containing different amounts of the

μ -oxo dimers were obtained. For the final analysis, a sample of $(\text{FeP})_2\text{O}$ consisting of around 70% of the dimer as determined by SQUID and Mössbauer measurements was chosen [11]. The impurities within this sample consist of the ^{57}Fe -PPIX-Cl and trace amounts of PPIX.

This synthesized product $(\text{FeP})_2\text{O}$, the commercial PPIX (Sigma) and ferriprotoporphyrin IX chloride (Fe-PPIX-Cl) (Alfa Aesar) were tested by UV/vis and Fourier transform infrared (FTIR) absorption spectroscopy as well as by fluorescence and Raman spectroscopy. In the case of UV/vis absorption and fluorescence spectroscopy the samples were examined in dimethylformamide (DMF) solution, while in the FTIR and Raman spectroscopies as KBr pellets (1% concentration).

UV/vis absorption spectra were taken with the use of the JASCO V-530 spectrometer with a spectral resolution of 2 nm. The fluorescence spectra were taken by a Hitachi F-4500 spectrophotometer. The emission slit was 2.5 nm while the excitation slit was set for 2.5 nm or 5 nm in the case of fluorescence emission or excitation measurements, respectively.

The FTIR measurements were performed with a Bruker "Equinox 55" interferometer, modified to work under vacuum. System parameters: FIR range ($10\text{--}700\text{ cm}^{-1}$), source: Hg lamp, detector: Si bolometer at 4.2 K; mid infrared (MIR) range ($600\text{--}5000\text{ cm}^{-1}$), source: globar (silicon carbide rod), detector: MCT. The spectral resolution was 4 cm^{-1} and the sample was a 1% by weight KBr pellet.

The Raman spectroscopy was carried out using an Instruments S.A. (Horiba Jobin-Yvon) Labram 1B operating at a wavelength of 514.5 nm. The power at the sample was $\approx 10\text{ mW}$ and a $10\times$ microscope objective was employed. To avoid photodegradation the powdered materials were mixed ($\approx 1\%$) with KBr and pressed into pellet form. Whereas samples were visibly degraded in their pure powder form, no degradation of the Raman signal was observable during prolonged exposure/measurement (minutes) and no visible degradation was observable microscopically in the KBr pellet form.

3. Results and discussion

UV/vis absorption spectroscopy was carried out in order to establish any signatures of electronic interaction between the porphyrin moiety and the bridging unit, and to identify the optimum excitation wavelengths for photoemission studies. Figure 2 shows the UV/vis absorption spectrum of PPIX, Fe-PPIX-Cl , $(\text{FeP})_2\text{O}$. The spectra in all cases are dominated by the characteristic porphyrin Q-bands ($\approx 500\text{--}620\text{ nm}$) and Soret band ($\approx 400\text{ nm}$). The abrupt step at $\approx 400\text{ nm}$ is an artefact of the spectrometer. The Soret band is largely unchanged in the dimer products. The doublet Q-band splitting of the protoporphyrin is removed in the Fe-PPIX-Cl due to the increased symmetry in the metal environment. In the $(\text{FeP})_2\text{O}$ sample, these bands have become broadened probably due to the incomplete reaction. There is

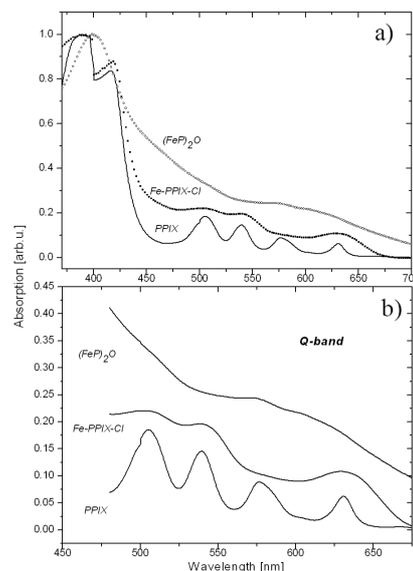


Fig. 2. (a) The UV/vis absorption spectra of all compounds (normalized at the Soret band for visual clarity). (b) The Q-band of the UV/vis of all compounds.

however no compelling evidence in the spectrum of the $(\text{FeP})_2\text{O}$ compound of specific signatures of the Fe-O-Fe bridge.

Figure 3 shows the FTIR spectra of the products compared with the Fe-PPIX-Cl monomer and PPIX. Of interest is the feature at $\approx 840\text{ cm}^{-1}$ which has previously been attributed to the asymmetric stretch of the Fe-O-Fe bridge [12–15]. The fact that both PPIX and Fe-PPIX-Cl exhibit absorption in this region is an indication that these vibrational modes originate from the porphyrin ring, however. The band at $\approx 840\text{ cm}^{-1}$ in PPIX is shifted to higher wave number in the Fe-PPIX-Cl by $\approx 14\text{ cm}^{-1}$, consistent with a stiffening of the ring by replacement of the central hydrogens by the metal. The band is softened in the $(\text{FeP})_2\text{O}$ compound, which is indicative of an increased effective mass of the central metal. While dimerization softens this mode it cannot be simply attributed to the Fe-O-Fe bridge. The FTIR spectrum of the porphyrin PPIX is extremely rich in this region and although many studies have assigned the strong bands in this region to direct evidence of Fe-O-Fe bridging [16–19] no identifiable independent signature of such a bridge is evident in Fig. 3.

Upon excitation at 600 nm, all compounds show characteristic porphyrin-like $S_0\text{--}S_1$ fluorescence emission features at $\approx 630\text{ nm}$. Excitation into the Soret band at 405 nm also gives rise to $S_0\text{--}S_1$ emission. While the PPIX and Fe-PPIX-Cl showed no indication of an $S_2\text{--}S_0$ emission, $(\text{FeP})_2\text{O}$ exhibits a notable emission at $\approx 480\text{ nm}$ (Fig. 4). Zinc porphyrins have been reported to exhibit $S_2\text{--}S_0$ emission although they are normally characterized by a small Stokes shift of $\approx 5\text{ nm}$ associated with the macromolecular ring relaxation [20, 21]. The emission in Fig. 4 has a significantly larger Stokes shift of \approx

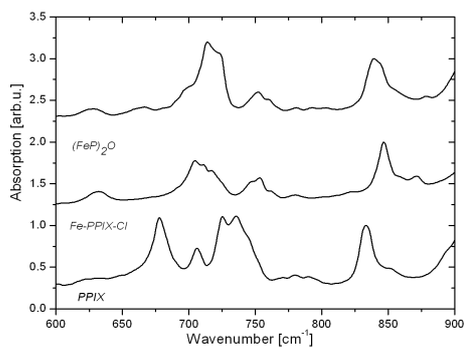


Fig. 3. The FTIR spectra of all compounds (normalized to the peak around 840 cm^{-1}). The spectra are offset for clarity.

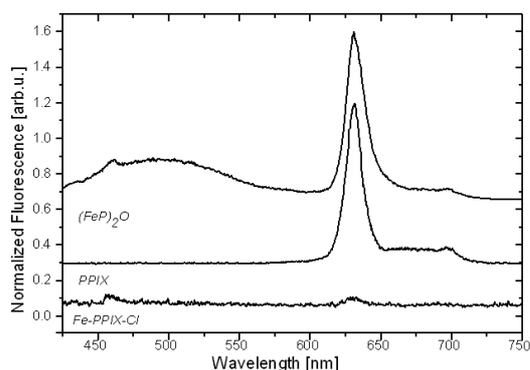


Fig. 4. The fluorescence spectra for all compounds (excitation wavelength 405 nm , normalized at 631 nm). The spectra are offset for clarity.

100 nm (eV) suggesting that it is not associated with the π -conjugated ring. Fe-O-Fe bridges have been reported to exhibit UV/vis absorbances at $360\text{--}380\text{ nm}$ which have been attributed to charge transfer states [15, 22–25]. Although they are not evident in Fig. 2, such an absorption could be present in the region of the Soret band. Figure 5 shows the excitation spectrum of the emission of

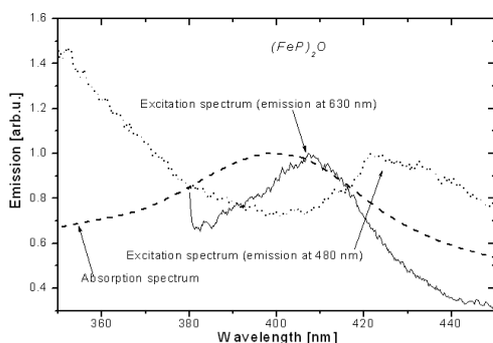


Fig. 5. Excitation spectra of $(\text{FeP})_2\text{O}$. The spectra are offset for clarity.

$(\text{FeP})_2\text{O}$ at 630 nm compared to that at 480 nm . The emission at 480 nm is clearly of different origin than the porphyrin based emission at 630 nm , whose excitation spectrum compares well with the porphyrin based Soret band emission.

The excitation spectrum has a maximum of 430 nm , considerably shifted from both that of the 630 nm emission ($\approx 407\text{ nm}$) and the absorption spectrum, also shown in Fig. 5. The presence of this emission band, the associated Stokes shift and the relative positioning of the excitation maximum suggest that it is not associated with the porphyrin ring but rather with an Fe-O-Fe bridge charge transfer state.

The Raman spectra of the compounds recorded at 514.5 nm are shown in Fig. 6. Similar to the FTIR spectra of Fig. 3, they are rich in the fingerprint regions $400\text{--}1700\text{ cm}^{-1}$. The complexity of the spectra makes it difficult to identify any features uniquely attributable to the dimer and consequently the Fe-O-Fe bridge. While the asymmetric stretch of bridged porphyrin dimers has been reported to lie at $\approx 850\text{ cm}^{-1}$, the symmetric stretch, which should be Raman active, has been reported to lie at $\approx 450\text{ cm}^{-1}$ [14, 15, 25–28]. In the spectra of Fig. 6, however, there is no evidence of characteristic features in this region. $(\text{FeP})_2\text{O}$ does however uniquely show a feature at 895 cm^{-1} which could be a candidate for the asymmetric stretch which may be swamped by the strong ring absorptions in the FTIR spectra. Previous studies have demonstrated that the symmetric stretch is weak compared to the overall compound spectrum and becomes prominent only upon resonant excitation at $\approx 400\text{ nm}$. Excitation is into the Fe-O-Fe bridge charge transfer state indicated above giving a clear signature of the Fe-O-Fe bridge. Such resonant excitation was not available for this study however and at 514.5 nm $(\text{FeP})_2\text{O}$ shows significant absorption associated with the porphyrin implying that the ring modes are resonantly enhanced.

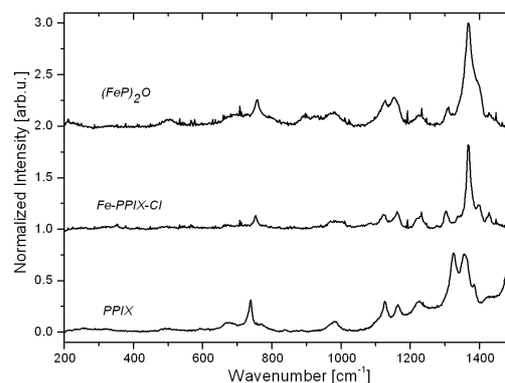


Fig. 6. Raman spectra of all compounds (normalized to the peak around 1630 cm^{-1}). The spectra are offset for clarity.

4. Summary and conclusion

The spectroscopy of the porphyrin based species in all cases is dominated by the π -conjugated macrocycle.

No unique signature of the dimer in (FeP)₂O, akin to the magnetic signatures of SQUID and Mössbauer spectroscopy, has been identified although strong supporting evidence is provided.

The UV/vis absorption spectra of all compounds are dominated by the strongly absorbing π -conjugated ring. The FTIR spectra are very complex and shifts of porphyrin ring modes, consistent with dimerization, can be observed. Features often assigned in literature to Fe-O-Fe bridges clearly have origin in the porphyrin ring, however. It is possible that the bridge modes underlie the very much stronger ring modes. Fluorescence spectroscopy provides more compelling evidence for a bridge based charge transfer state. The (FeP)₂O compound shows a significant emission at ≈ 480 nm, strongly Stokes shifted from the Soret band which has a significantly different excitation spectrum to the porphyrin emission, consistent with the presence of a charge transfer absorption feature in this region. Such an absorption feature in the UV/vis spectrum would be swamped by the strong Soret band, however. The Raman spectrum shows no evidence of the predicted Fe-O-Fe symmetric stretch mode at ≈ 450 cm⁻¹ although the product (FeP)₂O exhibits a feature at 895 cm⁻¹ which may be the asymmetric stretch which is swamped by the ring modes in the FTIR spectrum. The excitation wavelength employed is resonant with the ring absorptions and predictably the spectrum is dominated by the porphyrin based bands. Resonant excitation into the bridge charge transfer states at ≈ 400 nm should be employed to yield a clear signature of the dimer bridge.

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