Unveiled optical properties of tetrapyrrolic pigments in cryogenic environments

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An unexpected phenomenon was revealed in the laser induced fluorescence spectra of free-base (H₂Pc) and zinc (ZnPc) phthalocyanines trapped in rare gas and nitrogen matrices under a moderate increase in the laser intensity. In all matrices the intensity of an emission band near 755 nm increased drastically when pumping the S₁ ← S₀ transition. This observation was assigned to stimulated emission in a four-level scheme involving a vibronic transition from the lowest electronic state to a vibrational level of the ground state. In the present work, we expose new similar results obtained with porphyrin molecules, i.e. tetra-benzoporphin (TBP). With free-base H₂TBP, stimulated emission was observed in Ar or N₂ matrices, but not in Xe matrices. A possible reason could be a fast inter-system crossing rate due to the heavy atom effect induced by Xe. We also report the observation of persistent burnt spectral holes, although the low efficiency of this process is not competitive with stimulated emission and no decrease of the stimulated emission with time was observed. With ZnTBP no stimulated emission could be recorded but the appearance of rather strong phosphorescence was noticed. Similar studies focusing on the stimulated emission phenomenon is that they should allow new insights into site effects and site selectivity. © 2010 American Institute of Physics [doi:10.1063/1.3432263]

I. INTRODUCTION

Recent luminescence work¹ on phthalocyanine-doped cryogenic matrices revealed the occurrence of stimulated emission for specific vibronic transitions. This phenomenon was characterized by a drastic intensity increase of a specific vibronic fluorescence line with only a moderate increase of the laser pump intensity. The most concerned transition was assigned as the one taking place between the first excited electronic state and a vibrational level of the ground state assigned as the one taking place between the first excited electronic state and a vibrational level of the ground state located at approximately 1550 and 1525 cm⁻¹ in H₂Pc and ZnPc, respectively.¹

One important question to address in the development of this work is whether the stimulated emission observed for the phthalocyanines in cryogenic matrices is a more general behavior occurring for other molecules in other environments. With this question in mind we are therefore in search for other systems, molecules and/or matrices, in which similar behavior might be observed. In present paper, we are looking for similar processes in the case of porphyrin molecules belonging to the tetrabenzoporphin family. Indeed, tetrabenzoporphin (TBP) molecules have the closest structure to the phthalocyanines (Pc), the most noticeable change being the replacement of the four bridging nitrogen atoms belonging to the internal ring by carbon atoms (Fig. 1). Therefore similar vibronic structures are expected between both species.

The first absorption spectra of matrix-isolated porphyrins and phthalocyanines were reported early by Bajema et al.²,³ More recently Waluk and co-workers reported on the spectroscopy⁴ and isolation⁵ of porphyrins and related molecules in rare gas matrices. Other groups⁶–⁸ analyzed the vibronic structure on the S₁ state of several porphyrins and phthalocyanines in Shpol’skii matrices. A complete spectroscopic study and vibrational analysis of H₂TBP in a Shpol’skii matrix of n-octane (n-C₈) was early reported by Arabei.⁹ The same author also studied the influence of temperature on the electronic spectra of H₂TBP in n-octane.¹₀ However, the behavior of this important class of molecules isolated in the solid rare gases needs still to be more thoroughly analyzed and characterized.

Laser-induced fluorescence spectroscopy is very well adapted for studying these molecules because most of their absorption bands in the visible region closely match the working range of tunable dye lasers. While undertaking such a spectroscopic study of matrix-isolated free-base phthalocyanine (H₂Pc) and zinc phthalocyanine (ZnPc), by means of emission-excitation spectra using pulsed dye lasers for excitation, an unusually intense vibronic band was observed with slightly increased laser power. This novel solid state effect, which had never been reported before, was the subject of our paper.¹ The band involved corresponded to the transition from the first excited electronic state to a vibrational level of
the ground state at approximately 1550 cm\(^{-1}\). The drastic intensity enhancement was attributed to the occurrence of stimulated emission (SE) on the basis of the following observations: 1) the laser power dependence of the phenomenon exhibits a well-defined threshold, 2) there is also a threshold as a function of the concentration of dopants, 3) the emission band is narrow compared to the spectral width of the standard fluorescence, even in the case of broad bands in the absorption spectrum, and 4) its lifetime is strongly shortened in comparison with other vibronic transitions of the emission spectrum and exhibits a perfect synchronism with the exciting laser pulses.\(^1\)

A complete analysis is complicated by the different possible sites occupied by the guest molecules in the crystalline host; however, one important point noticed was that the SE line appeared as a narrow line from the most populated family of sites. In the case of H\(_2\)Pc/N\(_2\), where different sites were clearly observed in the absorption band, SE exhibited several lines, correlated with the main sites. Excitation spectra highlighted energy transfers between the families of sites. In the case of H\(_2\)Pc/Ne, a spectral hole burning (SHB) phenomenon was also present and competed with SE, resulting in a decrease of the SE intensity with the total irradiation time.\(^1\) These studies of the SE phenomenon should allow new insights into site effects and site selectivity. Possible reasons for the different observations made depending on the different hosts studied have still to be discussed. To extend the range of guest molecules beyond those belonging to the phthalocyanine family, it is easy to be convinced of the strong similarity, both in terms of the molecular structure and the spectroscopy, between phthalocyanines and tetrabenzoporphyrids (Fig. 1). The aim of the present paper is to present new and preliminary results obtained with molecules belonging to benzoporphyrids in different cryogenic environments.

**II. EXPERIMENTAL DETAILS**

The tetrabenzoporphyrids, free-base H\(_2\)TBP as well as ZnTBP, studied in this work were not commercial products. However, we can mention that Frontier Scientific Inc. can also make these molecules available. The molecules we used were synthesized according to the procedure described in Ref. 11 and carefully controlled by chromatography. For sub-

**III. RESULTS AND DISCUSSION**

The results obtained for H\(_2\)TBP isolated in Ar, Xe, and N\(_2\) and for ZnTBP isolated in Ar and N\(_2\) are presented and discussed in this paper.
A. Absorption spectra

Following the procedure previously mentioned, we can ascertain that the absorption spectrum of H₂TBP/Ar shown in Fig. 3a, clearly corresponds to well-isolated guest molecules. The two main bands appearing at about 654 nm (15284 cm⁻¹) and 584 nm (17132 cm⁻¹) belong to absorptions of the split Q band, frequently labeled as the Qₓ and Qᵧ bands, respectively, in order of increasing energy. The two corresponding bands in the H₂Pc/Ar absorption spectrum are located at about 677 nm (14771 cm⁻¹) and 635 nm (15748 cm⁻¹). The shift of the absorption bands observed between H₂TBP and H₂Pc results from the change in energy of the molecular orbitals due to the replacement of the interpyrrolic carbons Cₘ by the nitrogen atoms (Fig. 1). As a consequence, the H₂Pc Q bands are shifted to the red while the B bands in the Soret’s region are oppositely blue shifted.

From an analysis of the different absorption spectra that we recorded for benzoporphyrin in several hosts we can extract the following results about the Qₓ−Qᵧ splitting. For H₂TBP, we report values of about 1848 cm⁻¹, 1655 cm⁻¹ (average value for the two main sites observed), and 1851 cm⁻¹ for Ar, Xe and N₂ hosts, respectively, as compared to the splitting values of 982 cm⁻¹, 958 cm⁻¹, and 1090 cm⁻¹ found for H₂Pc in the same hosts, respectively. It is important to mention that the values of the Qₓ−Qᵧ splitting are calculated by simply taking the difference between the maximum of the intense Qₓ band associated to the localization of the S₂←S₀ transition. Indeed, this is not entirely true as in doing this, the Fermi resonance⁹,¹⁴ which influences the exact position and intensity of the Qᵧ(0-0) band is not considered here. Therefore the position as well as the intensity of the Qᵧ(0-0) band are not precisely defined in this work.

Another comparison with previously reported spectra of H₂Pc in the same host allows us to make the following comments: absorption lines recorded for H₂Pc/Ar under similar conditions looked narrower that those reported in Fig. 3. The broadening observed is probably not due to a strong contribution from aggregates, even if we cannot completely exclude it. Indeed, H₂TBP/Ar spectra show some structured features that very likely arise from a greater number of separated sites in the matrix. From a more detailed view of the Qₓ absorption band of H₂TBP/Ar, at least seven sites can be distinguished. This point will be more explicitly clarified in a later paragraph concerning the observation of spectral hole burning. Note that the presence of multiple sites would also explain the width of the bands.

The absorption spectrum of ZnTBP/Ar at 8 K is shown in Fig. 3b. ZnTBP belongs to the D₄h point group, while H₂TBP belongs to Dₒ₅h and for symmetry reasons the two bands Qₓ and Qᵧ collapse into only one Q band located at about 613 nm in the argon host, i.e. in between the positions of the Qₓ and Qᵧ bands of H₂TBP. From spectroscopic measurements of ZnTBP in supersonic expansion,¹⁵ the S₁ state was located at 16579 cm⁻¹ (603.2 nm) above the ground state, a value slightly blue-shifted compared to 16316 cm⁻¹ (612.9 nm) found here at 8 K. The present matrix value is in perfect agreement with the previously reported one as 16313 cm⁻¹ for ZnTBP/Ar at 20 K.²

While the absorptions in nitrogen matrices are quite similar to those in argon hosts, the absorption spectra of H₂TBP in Xe clearly exhibits «doublets» structures probably corresponding to two major sites. A comparison between the energy locations of the S₁←S₀ and S₂←S₀ transitions for H₂TBP and ZnTBP in different environments at low temperatures is presented in Table I.

B. Emission spectra

1. α-fluorescence

Singlet emission spectra recorded for H₂TBP and ZnTBP in the Ar host are shown in Fig. 4. Both fluorescence spectra shown for H₂TBP and ZnTBP are presented with the same horizontal scale with the position of the 0-0 region made coincident to allow comparison of their vibronic structure. For H₂TBP the non-resonant fluorescence spectrum is shown (Fig. 4a). The higher energy band shown at about 654.2 nm corresponds to the 0-0 zero-phonon origin band. It’s structure suggests the presence of several marked sites as also noticed from the absorption spectrum. Besides the rather intense electronic origin region, the most intense band which appears in the H₂TBP/Ar spectrum is that centered at 731.9 nm, revealing, like in the case of H₂Pc, the existence...
of stimulated emission. Stimulated emission observed in the case of H$_2$TBP will be discussed in detail in a paragraph to follow.

The fluorescence emission from ZnTBP/Ar is shown in Fig. 4b. A complete analysis of this fluorescence spectrum is complicated by the probable presence of impurities. However, undoubtedly, the major band at 681.2 nm is assigned to the same active mode at approximately 1636 cm$^{-1}$ which gives rise to stimulated emission in H$_2$TBP/Ar. Nevertheless, no evidence of stimulated emission was observed in that case irrespective of the laser intensity. A negative result concerning the observation of stimulated emission was also obtained in the case of ZnTBP/N$_2$.

### 2. $\beta$-stimulated emission

As previously indicated the intense and narrowed line at 731.9 nm (13663 cm$^{-1}$) is due to stimulated emission. While collecting data about this stimulated emission we noticed a qualitatively dependence upon a threshold effect even if we did not characterize it quantitatively. This stimulated emission was also observed in N$_2$ matrices around 731 nm under pulsed laser excitation of Q$_X$ and/or Q$_Y$, but not in Xe crystals. Such a situation is quite similar to what we reported already in the case of free-base phthalocyanine, for which stimulated emission was also observed in Ne, Ar, Kr but not in Xe hosts. Stimulated emissions in H$_2$Pc were assigned to specific vibronic transitions involving a vibration mode at about 1550 cm$^{-1}$ assigned to the stretching vibration C$_{m}$N bonds of the C$\equiv$N-C bridges of the central ring,$^{16,17}$ whereas the corresponding mode in the case of H$_2$TBP is around 1620 cm$^{-1}$. As was noted in Refs. 9 and 18 the 1620 cm$^{-1}$ vibration of H$_2$TBP corresponds to the A$_g$ stretching vibration of the C$_{m}$C$_{\alpha}$ bonds, i.e. to the vibration which is localized in the C$_{m}$C$_{\alpha}$C$_{\alpha}$ bridges in H$_2$TBP (C$_{\alpha}$—carbon atom in the $\alpha$-position of the pyrrole rings, C$_{m}$—carbon atom in the meso- position (C$_{m}$C$_{\alpha}$C$_{\alpha}$ bridges) as shown in Fig. 1).

When the pump power intensity was increased by a factor of about ten, from tens of $\mu$J/pulse to hundreds of $\mu$J/pulse, the emission band at about 732 nm gained enormously in intensity while the others remained unchanged or diminished slightly. Moreover, the linewidth of this emission band was reduced considerably while its intensity increased. The width of the band is approaching that of the exciting laser, decreasing from approximately 8 cm$^{-1}$ to 2 cm$^{-1}$, a value surely limited by the resolving power of the monochromator.

Figure 5 shows a two-dimensional (2D) excitation/emission plot of the stimulated emission observed for free-base H$_2$TBP isolated in solid Ar at 8 K. The plot in Fig. 5a depicts the sites emission detected with excitation in the 0-0 region of the S$_1$←S$_0$ transition (Q$_X$ band) while Fig. 5b depicts that produced with excitation of the S$_2$←S$_0$ transition (Q$_Y$ band). Both 2D excitation/emission plots were recorded while monitoring the SE mode centered around 732 nm. For H$_2$TBP/Ar under pulsed excitation in the Q$_X$ region, around 654 nm, doublet SE lines (separated by 3 cm$^{-1}$) were recorded giving rise to similar excitation spectra (separated by about 5 cm$^{-1}$).

An analysis of the SE signals generated under selective excitation at 654.43 nm and 654.25 nm inside the broad Q$_X$...
absorption band of H₂TBP/Ar at 8 K revealed the existence of two stimulated emissions, which are assigned to two distinct sites (Fig. 5a). These two sites look independent. In contrast to the case of H₂Pc, no phonons are excited but a vibration at about 215 cm⁻¹ from the origin band is excited, as evidenced by looking at the absorption spectrum (dotted line) shown superimposed to the excitation spectra. Clearly, excitation spectra recorded for the SE signal provide greatly enhanced spectroscopic information over what can be extracted from either conventional fluorescence excitation or absorption spectroscopy.

The situation is more complicated in the case of H₂TBP/N₂ (not shown). Indeed, four sites giving rise to SE signals at 13678 cm⁻¹ (731.1 nm), 13687 cm⁻¹ (730.6 nm), 13694 cm⁻¹ (730.3 nm) and 13701 cm⁻¹ (729.9 nm) were identified, with possibilities for exciting several sites at one time because of possible transfers from one site to another as reported already for H₂Pc. In this case the excitation spectra of SE lines exhibit quite intense phonon bands allowing the simultaneous excitation of all sites.

Upon excitation of the Q₁ band of H₂TBP/Ar the observation of two sites is still revealed by the occurrence of stimulated emission, but the excitation spectra recorded were found to be more complex due to vibronic coupling between vibrational levels from the lower Qₓ state and the electronic Q₁ state. While monitoring one or the other site, the corresponding excitation spectra were revealed, demonstrating that they have the same structure simply shifted with respect to the one another by ~4 cm⁻¹, the exact energy spacing between the two sites. From this important observation we can conclude that the vibronic coupling between Q₁ and Qₓ is quite similar for the two sites, and that no phonon modes are contributing (Fig. 5b). For Qₓ in the case of H₂TBP/N₂, the situation (also not shown) needs to be clarified as disturbances due to excitation of phonon modes from the matrix are also assumed to contribute. Under Q₁ excitation, only the two main SE lines located at 13678 cm⁻¹ (731.1 nm) and 13687 cm⁻¹ (730.6 nm) are clearly observed, allowing the identification of two main sites separated by about ~9 cm⁻¹.

In the case of the fluorescence spectra of ZnTBP/Ar under pulsed laser excitation no stimulated emission enhancement could be observed irrespective of the incident laser intensity. A similar observation was also reported for H₂TBP/Xe. A possible explanation of the absence of stimulated emission in the case of ZnTBP and H₂TBP/Xe could be the heavy atom (Zn or Xe) effect which increases the probability of the intersystem crossing rate from the excited singlet S₁ state level to the lower lying triplet T₁ state. This non-radiative relaxation mechanism then competes with the fluorescence and thereby reduces the chance of the excited singlet state population for reaching the threshold condition. As a consequence of the enhanced intersystem crossing the triplet state is efficiently populated, leading to the observation of a phosphorescence signal.

3. c-phosphorescence

In the case of ZnTBP/Ar, for which no stimulated emission could be monitored, a rather strong phosphorescence signal was recorded with a lifetime longer than 30 ms. The corresponding phosphorescence spectrum is shown in Fig. 6.

The recorded spectrum is not easily understandable, and the situation is even unclear after attempting to analyze the associated excitation spectra (not shown). However, inspection of Fig. 6 shows that a «doublet» structure is clearly apparent in the phosphorescence spectra of ZnTBP/Ar and ZnTBP/N₂. A similar «doublet» structure was earlier reported for ZnTBP/n-C₈ by Arabei et al. The interpretation given by the authors was as follows: due to non-correlation between S₁–S₀ and T₁–T₀ transition energies in the site structure, a site selective excitation in the S₁–S₀ region can induce the phosphorescence from different sites with distinct T₁–S₀ transition energies. However several questions remain unsolved still and we consider that the phosphorescence spectra of ZnTBP in various cryogenic matrices requires further analysis. Amongst these, it is necessary to clarify whether or not efficient intersystem transfer, as ascertained

FIG. 5. 2D emission-excitation spectra of free-base H₂TBP/Ar at 8 K recorded under selective laser excitation. Both Qₓ and Q₁ regions were studied, emission spectra are shown on top, excitation spectra, together with the absorption spectra (dotted lines) are on the left. Analysis of the 2D spectrum under SE conditions of Q₁ region around 654 nm (a). Analysis of the 2D spectrum under SE conditions of Q₁ region around 584 nm (b).
by the observed phosphorescence, is the major factor for the non-observation of stimulated emission in ZnTBP.

C. Spectral hole-burning

With H₂TBP/Ar persistent spectral changes could be detected under selective laser excitation at 8 K. These spectral hole-burning changes were observed when looking for stimulated emission conditions under selective excitation of one or the other site. At low temperature the spectral hole-burning phenomenon is quite well known in the case of free-base porphyrins. It is then not unusual to observe it with H₂TBP. Figure 7 shows an expanded view of the absorption spectrum corresponding to the Qₓ absorption band before excitation and then, after excitation of one site and then of the other one. The inset of Fig. 7 presents a more expanded view to show the changes observed close to the band maximum. Typical burning times needed to observe this persistent spectral change were in the order of one hour for laser energy of ~40 μJ/pulse at 10 Hz.

Among the different sites which are spectrally identified in the absorption, only two specific lines showed selective persistent spectral changes. Burning at 654.25 nm induced a decrease of the absorption at this wavelength, while a similar change was observed when burning the other site at 654.43 nm. It is interesting to notice that quite similar spectral changes are observed on the vibronic band located at about 15500 cm⁻¹, assigned to the same vibration mode as the one which gives rise to SE under laser excitation. This phenomenon could be explained by tautomerization. In this model the two sites would be associated with the two tautomers of the porphyrin molecule, and the observed changes arise from the interconversion of one form to another, i.e. the flipping of the inner protons from one position to the other. In the case of H₂TBP/Ar this process has a rather low efficiency and cannot reduce SE significantly. Such a situation differs from what we reported for H₂Pc/Ne, for which very efficient hole-burning phenomenon was observed in the 0-0 S₁ ← S₀ absorption band with even weak dye laser excitation. Consequently, as the excited family of sites disappeared during excitation, SE was not sustainable for long periods in this solid.

IV. CONCLUSIONS

The results obtained in the present work confirm that the phenomenon of stimulated emission first observed for phthaloxyanines in cryogenic matrices under modest conditions of pulsed laser excitation is not an exception. With free-base H₂TBP stimulated emission was observed in Ar and N₂ matrices but not in Xe matrices. The vibronic transition involved is, like in the case of H₂Pc, the one involving a mode at about 1620 cm⁻¹, a particularly active mode corresponding to the stretching Aₙ vibration of the C₉C₉ bonds. We also reported the observation of persistent spectral hole-burning; however the quite low efficiency of the burning process can not affect SE. Finally, no stimulated emission was recorded with ZnTBP, but the appearance of rather strong phosphorescence was noticed. From all results we have obtained on SE in cryogenic matrices with H₂Pc, ZnPc, as well as H₂TBP we can reasonably predict that SE should also be observable for these molecules isolated in other solid materials that also produce narrow linewidths such as Shpol’skii matrices, like n-octane. Other possible interesting candidates could also be sol-gel or polymer hosts.

Another important remark is the following: compared to what is obtained in normal fluorescence excitation scans, greatly enhanced site selectivity is obtained for excitation scans recorded for the SE mode. This behavior stems from the increased resolution of individual sites whose lineshape change from featureless fluorescence excitation bands to a highly structured SE excitation band. The very well-resolved spectra available in SE excitation scans allow detailed exploration of site occupancies in a given solid and from one solid to another. Added to our previous, already published results, the results of the present study of SE should allow new insights into site effects and site selectivity.

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