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Effects of Structural Deformations on Optical Properties of Tetrabenzoporphyrins: Free-Bases and Pd Complexes

Artem Y. Lebedev,[†] Mikhail A. Filatov,[‡] Andrei V. Cheprakov,^{*,‡} and Sergei A. Vinogradov^{*,†}

Department of Biochemistry and Biophysics, University of Pennsylvania, Philadelphia, Pennsylvania 19104, and Department of Chemistry, Moscow State University, Moscow 119899, Russia

Received: May 16, 2008; Revised Manuscript Received: June 12, 2008

A recently developed method of synthesis of π -extended porphyrins made it possible to prepare a series of tetrabenzoporphyrins (TBP) with different numbers of *meso*-aryl substituents. The photophysical parameters of free-bases and Pd complexes of *meso*-unsubstituted TBP's, 5,15-diaryl-TBP's (Ar_2TBP 's) and 5,10,15,20-tetraaryl-TBP's (Ar_4TBP 's) were measured. For comparison, similarly *meso*-arylsubstituted porphyrins fused with nonaromatic cyclohexeno-rings, i.e. Ar_n -tetracyclohexenoporphyrins (Ar_nTCHP 's, $n = 0, 2, 4$), were also synthesized and studied. Structural information was obtained by ab initio (DFT) calculations and X-ray crystallography. It was found that: 1) Free-base Ar_4TBP 's are strongly distorted out-of-plane (saddled), possess broadened, red-shifted spectra, short excited-state lifetimes and low fluorescence quantum yields ($\tau_{\text{fl}} = 2\text{--}3$ ns, $\phi_{\text{fl}} = 0.02\text{--}0.03$). These features are characteristic of other nonplanar free-base porphyrins, including Ar_4TCHP 's. 2) Ar_2TBP free-bases possess completely planar geometries, although with significant in-plane deformations. These deformations have practically no effect on the singlet excited-state properties of Ar_2TBP 's as compared to planar *meso*-unsubstituted TBP's. Both types of porphyrins retain strong fluorescence ($\tau_{\text{fl}} = 10\text{--}12$ ns, $\phi_{\text{fl}} = 0.3\text{--}0.4$), and their radiative rate constants (k_{r}) are 3–4 times higher than those of planar H_2TCHP 's. 3) Nonplanar deformations dramatically enhance nonradiative decay of triplet states of regular Pd porphyrins. For example, planar PdTCHP phosphoresces with high quantum yield ($\phi_{\text{phos}} = 0.45$, $\tau_{\text{phos}} = 1118$ μs), while saddled PdPh₄TCHP is practically nonemissive. In contrast, both ruffled and saddled PdAr_nTBP's retain strong phosphorescence at ambient temperatures (PdPh₂TBP: $\tau_{\text{phos}} = 496$ μs , $\phi_{\text{phos}} = 0.15$; PdPh₄TBP: $\tau_{\text{phos}} = 258$ μs , $\phi_{\text{phos}} = 0.08$). It appears that π -extension is capable of counterbalancing deleterious effects of nonplanar deformations on triplet emissivity of Pd porphyrins.

Introduction

Porphyrins extended by way of annealing the pyrrole residues with external aromatic rings are commonly referred to as π -extended porphyrins. Symmetrically π -extended porphyrins, e.g., tetrabenzoporphyrins and tetranaphthoporphyrins, possess high absorption in the near-infrared region of the spectrum, and consequently draw attention in photomedicine,¹ biomedical sensing and imaging.² In addition, π -extended porphyrins have shown promise in up-conversion of noncoherent infrared light,³ optical power limiting and electrooptical applications⁴ as well as in material science.⁵ Such broad range of applicability warrants in-depth studies of structure–property relationships in the family of π -extended porphyrins, especially in the area of their photophysics, which is central to many of their applications.

Starting 1960s, influence of the π -extension on optical properties of porphyrins has been addressed by a number of researchers.^{6–8} As early as in 1961, Gouterman applied his four-orbital model to qualitatively explain differences between the electronic absorption spectra of regular porphyrins and tetrabenzoporphyrins.^{6a} Soon after that his group⁶ and the group from Belarus⁷ published first reports on the luminescence of tetrabenzoporphyrin complexes. In more recent years, several in-depth theoretical studies⁹ provided insights into the electronic structure of π -extended tetrapyrroles, focusing mainly on their

ground-state absorption spectra. For example, combined influence of π -extension and *meso*-tetraarylation on the spectra of tetrabenzo- and tetranaphthoporphyrins has been examined.¹⁰

In spite of these advances, effects of π -extension on the excited-state properties of porphyrins have been studied only scarcely, notwithstanding the fact that many potential uses of π -extended porphyrins rely on their singlet and triplet excited states. One particular question, much discussed in relation to regular porphyrins, is the interplay between the macrocycle structural deformations and its photophysics.^{11–13} Although fluorescence and phosphorescence of tetrabenzoporphyrins^{2,1a–c,7,8,10,14} and tetranaphthoporphyrins¹⁵ has been mentioned in a number of publications, relationships between structural features and excited-state properties of these porphyrins have never been studied systematically. Such structure/property analysis could not be done because structural information on π -extended porphyrins was extremely limited,^{16,17} in part due to their poor synthetic availability.

Over the past several years, a general method of synthesis of tetrabenzoporphyrins has been developed, based on the oxidative aromatization of porphyrins, annealed with nonaromatic hydrocarbon rings.¹⁸ This method provides an access to *meso*-tetraaryl-tetrabenzo-^{18a,b} and tetranaphthoporphyrins (Ar_4TBP 's and Ar_4TNP 's)^{18c,d} as well as to *meso*-unsubstituted tetrabenzo- and tetranaphthoporphyrins (TBP's and TNP's).^{18e} A recent modification of the method, based on the use of 4,7-dihydroisoindole,^{18f} also opened a pathway to 5,15-diaryl-tetrabenzoporphyrins (Ar_2TBP).^{18g} In all these cases, various functional

* To whom correspondence should be addressed. E-mail: vinograd@mail.med.upenn.edu (S.A.V.); avchep@elorg.chem.msu.ru (A.V.C.).

[†] University of Pennsylvania.

[‡] Moscow State University.

groups can be added to the TBP macrocycles, increasing their solubility and facilitating crystal growth for X-ray structure determination. As a result, originally scarce structural information became more complete, enabling structure/property studies in the family of π -extended porphyrins.

In this work we analyzed photophysical properties of tetrabenzoporphyrins substituted with different numbers of *meso*-aryl groups. *meso*-Arylation is known to induce structural deformations in β -substituted porphyrins,¹⁹ and, therefore, it can be used to study effects of macrocycle distortions on tetrabenzoporphyrin photophysics. In particular, we focused on fluorescent free-base TBP's, Ar_2TBP 's and Ar_4TBP 's and the corresponding phosphorescent Pd complexes, which to us present a special interest as near-infrared optical probes for biological imaging of oxygen.²

Effects of nonplanarity on the photophysics of regular porphyrins were systematically studied by Holten and co-workers,¹³ who elucidated deactivation pathways of singlet excited $^1(\pi, \pi^*)$ states (S_1) of nonplanar free-bases,^{13a,b,d,f,h} Zn porphyrins,^{13d} Ni porphyrins^{13c,e} and porphyrin dication.^{13g} Formation and deactivation of triplet $^3(\pi, \pi^*)$ states (T_1) in nonplanar porphyrins has also been addressed,²⁰ including recent studies of nonplanar Pt and Pd porphyrins.²¹ In the present paper we compared tetrabenzoporphyrins Ar_nTBP ($n = 0, 2, 4$) with porphyrins annealed with nonaromatic cyclohexeno-rings, i.e. tetracyclohexenoporphyrins (Ar_nTCHP 's, $n = 0, 2, 4$). These were chosen among other β -alkyl-substituted porphyrins because of their close structural similarity to Ar_nTBP 's. This comparison enabled us to directly evaluate π -extension as a factor influencing photophysics of nonplanar porphyrins.

Results and Discussion

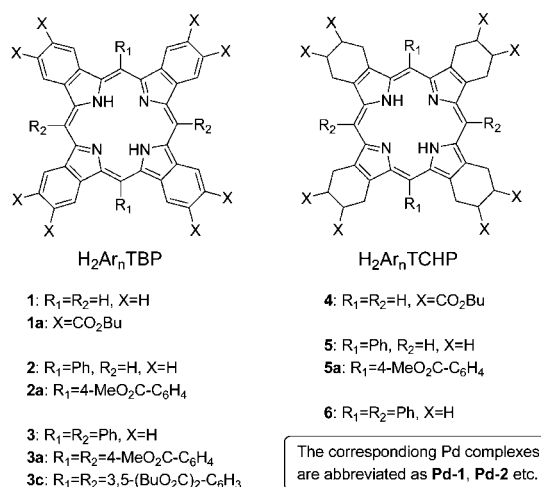
In the beginning we briefly summarize relevant facts about the photophysics and structures of conformationally distorted porphyrins. For detailed discussion see refs.11, 13, 19.

The most common way to induce structural deformations in synthetic porphyrins¹⁹ is to position several bulky substituents at the macrocycle periphery. Steric repulsion between the β -pyrrole groups and *meso*-substituents leads to deviations of the macrocycle from its highly symmetrical D_{2h} (free-base porphyrins) or D_{4h} (metalloporphyrin) form. Macrocycle deformations fall into two main categories: in-plane (ip) and out-of-plane distortions (oop). There are several ways to quantify these distortions; however, the most general method is to decompose the macrocycle structure in the basis of distortions corresponding to the normal vibrational modes. The Normal mode Structural Decomposition (NSD) analysis, developed by Shelnutt and co-workers,²² quantifies not only the total in-plane and out-of-plane displacements (D_{ip} and D_{oop}), but distinguishes between individual contributions of different distortion modes,^{22,19} which are known to affect the photophysics of nonplanar porphyrins differently.^{13a,b}

Distorted porphyrins typically exhibit mixtures of distortion modes; however, each structure is usually dominated by a single prevalent type of distortion. For example, out-of-plane distorted porphyrins are usually either mostly saddled (B_{2u} -deformation, e.g., octaethyltetraphenylporphyrin (OETPP) or dodecaphenylporphyrin (DPP)) or ruffled (B_{1u} -deformation, e.g., *meso*-tetra-*tert*-butylporphyrin (*T*(*t*-Bu)P)).^{23,19}

Compared to planar porphyrins, such as octaethylporphyrin (OEP) or tetraphenylporphyrin (TPP), out-of-plane distorted porphyrins exhibit red-shifted and broadened optical spectra. The origin of red shifts in the spectra of nonplanar porphyrins has been extensively discussed in the literature.^{11,12f-h} In *meso*-

CHART 1



tetraarylporphyrins the red shifts at least in part are caused by an increase in the conjugation between the macrocycle and *meso*-aryl substituents.^{10,24} In nonplanar porphyrins, the *meso*-aryl groups are rotated to a less than 90° angle with respect to the macrocycle average plane.

Nonplanar deformations of porphyrins also strongly affect properties of excited states. Both saddling and ruffling broaden the emission spectra and induce progressively large Stokes shifts with increasing degree of nonplanarity.¹³ Saddle-shaped free-base porphyrins (e.g., H_2OETPP and H_2DPP) typically exhibit S_1 lifetimes (τ_f) of about 0.5–1.0 ns and fluorescence quantum yields (ϕ_f) in the order of 0.005 at room temperature.^{11,13a} Ruffled free-base porphyrins have even shorter singlet lifetimes (e.g., $\tau_f = 50$ ps for $\text{H}_2\text{T}(t\text{-Bu})\text{P}$) and negligible emission yields ($\phi_f < 0.0001$).^{13b} For comparison, singlet lifetimes and fluorescence quantum yields of the planar analogues (e.g., H_2OEP or H_2TPP) are 10–15 ns and 0.1–0.2, respectively, while their internal conversion quantum yields are in the range $\phi_{ic} \approx 0.1$.^{13,25,26} Short S_1 lifetimes of nonplanar porphyrins were shown to be affiliated with enhancements of both $S_1 \rightarrow S_0$ internal conversion and $S_1 \rightarrow T_1$ intersystem crossing.^{12c} Polar solvents further increase the Stokes shifts, shorten the lifetimes and decrease the fluorescence quantum yields of nonplanar porphyrins,^{13h} while lowering the temperature has the opposite effect, especially in the case of ruffled porphyrins.^{13d}

Holten and co-workers rationalized these findings by considering increased conformational flexibility of nonplanar porphyrins in their S_0 and S_1 states.¹³ Broadening of the absorption and fluorescence spectra and large fluorescence Stokes shifts were associated with the existence of multiple excited-state conformations, easily accessible *via* low-energy out-of-plane vibrations. Some of these conformations can form so-called *funnels*²⁷ on the S_1 potential energy surface, separated from the global energy minimum by small kinetic barriers. Rapid internal conversion occurring in the funnels is due to the small S_1 – S_0 energy gaps and favorable Franck–Condon factors, corresponding to out-of-plane vibrations. In addition, intersystem crossing near the funnels also can be enhanced. On the contrary, fluorescence occurs away from the funnels (e.g., near the global S_1 minimum), where the S_1 – S_0 gaps are quite large.

The photophysical properties of tetrabenzoporphyrins are discussed below within the framework of this model.

Free-Base Ar_nTBP 's and Ar_nTCHP 's ($n = 0, 2, 4$). The structures of porphyrins studied in this work are summarized in Chart 1 (see Supporting Information, II for details of synthesis).

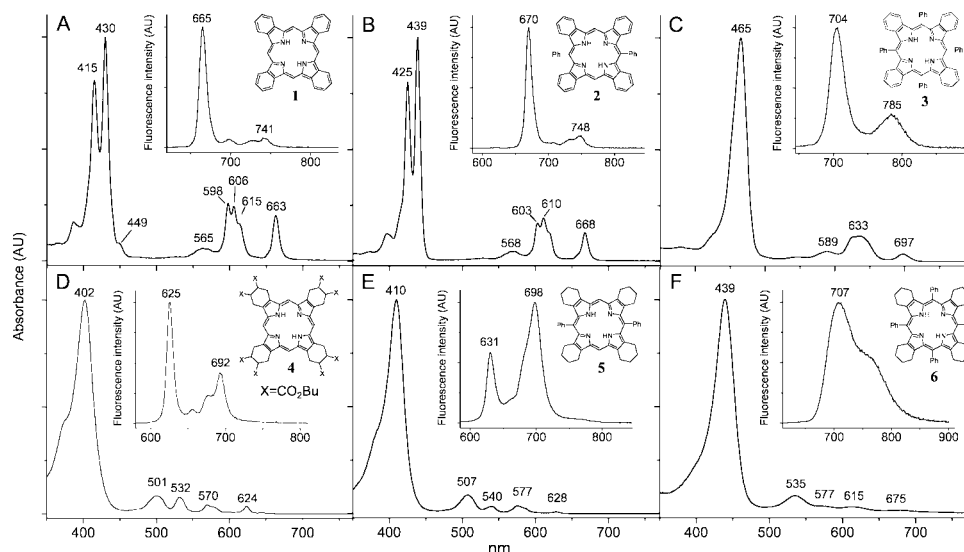


Figure 1. Absorption and fluorescence (insets) spectra of H₂TBP (**1**, A) (H₂TBP was first dissolved in pyridine, and a drop of this solution was added to toluene.), H₂Ph₂TBP (**2**, B), H₂Ph₄TBP (**3**, C) and H₂TCHP(CO₂Bu)₈ (**4**, D), H₂Ph₂TCHP (**5**, E) and H₂Ph₄TCHP (**6**, F) (Small amount of TMEDA was added to prevent dication formation.) in toluene at 295 K.

TABLE 1: Photophysical Properties of Free-Base Porphyrins H₂Ar_nTBP and H₂Ar_nTCHP (*n* = 0, 2, 4) in Toluene at 295 K and in Dimethylacetamide (DMA, Noted)^a

free-base porphyrin	fluorescence λ_{\max} (λ_{ex}), nm	Stokes shift (cm ⁻¹)	ϕ_{fl}^b	τ_{fl} (ns) ^c
TBP				
1	665, 741 (571)	45.4	0.35	9.8
1 (in DMA)	663, 740 (571)	45.6	0.47	—
2	670, 746 (571)	44.7	0.34	10.3
2 (in DMA)	668, 744 (517)	45.0	0.41	—
2a	668, 744 (517)	45.0	—	9.6
3	705, 786 (635)	162.8	0.03	2.8 ^c
3 (in DMA)	707, 791 (635)	223.5	0.02	—
3a	712, 792 (594)	220.4	0.03	2.9 ^c
TCHP				
4	625, 692 (531)	25.6	0.14	14.1
5	631, 698 (507)	75.7	0.08	11.8
6	707 (535)	670	0.01	2.6 ^c
6 (in DMA)	720 (635)	882	<0.01	—

^a A drop of H₂TBP solution in pyridine was added to toluene.

^b Quantum yields were measured relative to the fluorescence of H₂TBP in deoxygenated C₆H₆ ($\phi_{\text{fl}} = 0.11$ in deoxygenated benzene).²⁵ Error in determination of yields is no more than 10%.

Purging samples with Ar was shown to increase the fluorescence quantum yields by no more than 5% both in toluene and DMA.

^c Inhomogeneous multiexponential decays with broad lifetime distributions were detected. Average lifetime value is reported in the Table 1.

The optical absorption and fluorescence spectra of free-base Ar_nTBP's (*n* = 0, 2, 4) are shown in Figure 1 (upper row). The spectra of the similarly substituted *meso*-aryl tetracyclohexenoporphyrins (Ar_nTCHP's, *n* = 0, 2, 4) are shown in the bottom row, and all the photophysical data are summarized in Table 1. All the measurements were performed in air-equilibrated toluene or dimethylacetamide (DMA) (noted) solutions at 295 K, except that: 1) to overcome solubility problems, H₂TBP (**1**) was first dissolved in pyridine, and a drop of this solution was added to about 5 mL of toluene; 2) a small amount of TMEDA (tetramethylethylenediamine) was added to the solution of H₂Ph₄TCHP (**6**) to avoid protonation, since H₂Ar₄TCHP's are known to possess very high basicities.²⁸ Due to its improved solubility, H₂TCHP (**4**), substituted with eight butoxycarbonyl groups, was used in our experiments instead of unsubstituted

H₂TCHP. The effect of the butoxycarbonyl groups on the macrocycle π -system in H₂TCHP (**5**) is anticipated to be insignificant, since these substituents are separated from the macrocycle by several σ -bonds. In addition to free-bases shown in Figure 1, porphyrins with peripheral substituents (Chart 1) were used in some of the experiments (vide infra).

The optical absorption spectrum of planar H₂TBP (**1**) (Figure 1A) shows a well-resolved vibronic structure in the Q-band region and a large splitting, 841 cm⁻¹, of the Soret band. This splitting is caused by the strong mixing of the B and Q states and relatively high intensity of the Q-bands.^{6a,9b,c} The lowest energy Q-band (663 nm) is red-shifted by 39 nm (943 cm⁻¹) relative to the corresponding transition of the also planar (vide infra)²⁹ H₂TCHP (**4**) (Figure 1D) due to the effect of extended π -conjugation.^{9,10}

Both H₂TBP (**1**) and H₂TCHP (**4**) exhibit small fluorescence Stokes shifts (45 cm⁻¹ and 25.6 cm⁻¹), indicating rigid planar structures. In good agreement with the previously published data,^{1b,8c} H₂TBP (**1**) shows a very high for porphyrins fluorescence quantum yield ($\phi_{\text{fl}} = 0.35$ in toluene), while the yield of H₂TCHP (**4**) is significantly lower ($\phi_{\text{fl}} = 0.14$), nearly matching that of H₂OEP ($\phi_{\text{fl}} = 0.16$).²⁶ The S₁ lifetime of H₂TBP (**1**) (9.8 ns) is shorter than that of H₂TCHP (**4**) (14.1 ns). The calculated values of the corresponding rate constants are presented in Table 2. An almost 3.6 times higher value of the radiate rate constant (k_{r}) for H₂TBP (**1**) agrees well with much higher oscillator strength of its Q-band transition (log ϵ (663 nm) = 4.22 for H₂TBP (**1**) vs log ϵ (624 nm) = 3.70 for H₂TCHP (**4**)).^{18c}

The quantum yield of the nonradiative decay ($\phi_{\text{nr}} = \phi_{\text{ic}} + \phi_{\text{isc}}$; ϕ_{ic} , internal conversion; ϕ_{isc} , intersystem crossing) for H₂TCHP (**4**) ($\phi_{\text{nr}} = 0.86$) is higher than that for H₂TBP (**1**) ($\phi_{\text{nr}} = 0.65$), but the corresponding nonradiative rate constant ($k_{\text{nr}} = k_{\text{ic}} + k_{\text{isc}}$) for H₂TCHP (**4**) is slightly lower (Table 2). The faster nonradiative decay of H₂TBP (**1**) might be due to the higher rate of the internal conversion alone, since the S₁–S₀ gap for H₂TBP (**1**) is almost 1000 cm⁻¹ narrower than that in H₂TCHP (**4**); however, it may also reflect the altered rate of intersystem crossing.

The optical properties of H₂Ph₄TBP (**3**) (Figure 1C) (strongly saddled structure, vide infra) in general follow the pattern

TABLE 2: Deactivation Rate Constants for S₁ State of Free-Base Porphyrins H₂Ar_nTBP and H₂Ar_nTCHP (*n* = 0, 2, 4) in Toluene at 295 K

compound	$k = 1/\tau_{\text{fl}}$ (s ⁻¹)	k_r (s ⁻¹) ^a	k_{nr} (s ⁻¹) ^b
TBP			
H ₂ TBP (1)	1.02×10^8	3.57×10^7	6.63×10^7
H ₂ Ph ₂ TBP (2)	9.71×10^7	3.30×10^7	6.41×10^7
H ₂ Ph ₄ TBP (3)	3.57×10^8	1.07×10^7	3.46×10^8
TCHP			
H ₂ TCHP (4)	7.09×10^7	9.93×10^6	6.10×10^7
H ₂ Ph ₂ TCHP (5)	8.47×10^7	6.78×10^6	7.80×10^7
H ₂ Ph ₄ TCHP (6)	3.85×10^8	3.85×10^6	3.81×10^8

^a k_r , radiative decay rate constant, calculated as $k_r = \phi_{\text{fl}}k$, where ϕ_{fl} is the fluorescence quantum yield. ^b k_{nr} , nonradiative decay rate constant ($k_{\text{nr}} = k_{\text{ic}} + k_{\text{isc}}$), calculated as $k_r = k - k_{\text{nr}}$.

predicted by the Holten's model.¹³ Upon going from H₂TBP (1) to H₂Ph₄TBP (3), the fine vibronic structure of the Q-band disappears, and both bands shift to the red: Q-band by 736 cm⁻¹, Soret band by 2163 cm⁻¹. Similar trends are seen in the absorption spectrum of also saddled (vide infra) H₂Ph₄TCHP (6) (Figure 1F) vs planar H₂TCHP (4) (Figure 1D). The two latter spectra closely resemble the published spectra of H₂OETPP and H₂OEP, respectively.^{13a,26}

The changes in the fluorescence of H₂Ph₄TBP (3) vs that of H₂TBP (1) in general also conform to the model (see Table 1). For example, the fluorescence quantum yield of H₂Ph₄TBP (3) is lower than that of H₂TBP (1) by about 10 times, almost matching the change in the pair H₂Ph₄TCHP (6)/H₂TCHP (4). The Stokes shift of H₂Ph₄TBP (3) (162.8 cm⁻¹) is somewhat lower than that of H₂Ph₄TCHP (6) (670 cm⁻¹), and upon changing the solvent to more polar DMA it increases by 36% (to 223 cm⁻¹). The Stokes shift for H₂Ph₄TCHP (6) increases in DMA by almost the same value (to 882 cm⁻¹).

Typical of nonplanar porphyrins,^{13h} the fluorescence quantum yield of H₂Ph₄TBP (3) decreases in more polar solvent and its emission spectrum broadens (see Supporting Information, III). Interestingly, the fluorescence quantum yields of planar H₂TBP (1) increases in DMA to 0.47—a change not normally observed for planar porphyrins.

The dynamic fluorescence data for the most part are consistent with the steady-state measurements. The fluorescence decays of both H₂Ph₄TBP (3) and H₂Ph₄TCHP (6) reveal broad underlying distributions of lifetimes (recovered by the Maximum Entropy Method (MEM),³⁰ Figure 2), which are indicative of multiple excited-state conformations in agreement with the model. The radiative rate constants (k_r) (Table 2) decrease by about 3.3 times on going from H₂TBP (1) to H₂Ph₄TBP (3) and by about 2.6 times on going from H₂TCHP (4) to H₂Ph₄TCHP (6). Therefore, low S₁ lifetimes (see Table 1) of both *meso*-tetraarylated porphyrins are due to the strong enhancement of the nonradiative processes. Indeed, the corresponding rate constants (k_{nr}) increase by 6.2 times from H₂TCHP (4) to H₂Ph₄TCHP (6) and by only slightly less, i.e. 5.2 times, from H₂TBP (1) to H₂Ph₄TBP (3). Notably, the nonradiative rate for H₂OETPP is significantly higher, i.e., $k_{\text{nr}} = 1.5 \times 10^9$ s⁻¹,^{13a} than for H₂Ph₄TCHP.

The main conclusion drawn from these comparisons is that S₁ states of nonplanar porphyrins are significantly less prone to nonradiative deactivation when these porphyrins are annealed with cyclic motifs (as in H₂Ph₄TCHP (6)) as opposed to β -substituted with pendant alkyl groups (as in H₂OETPP). In addition, if the cyclic motifs are in conjugation with the macrocycle (as in H₂Ph₄TBP (3)), the nonradiative decay

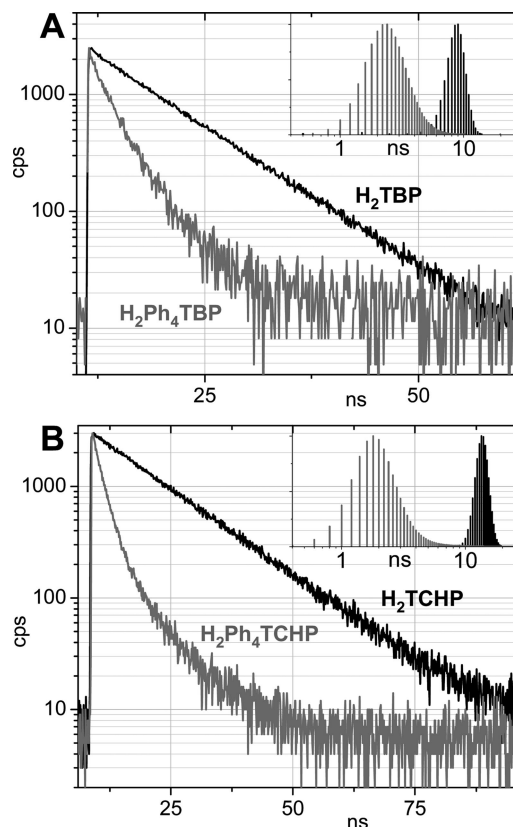


Figure 2. Fluorescence decays for H₂TBP (1), H₂Ph₄TBP (3) (A) and H₂TBP (4) and H₂Ph₄TCHP (6) (B) in toluene at 296 K and the underlying lifetime distributions, recovered by the Maximum Entropy Method.

appears to weaken even more, although within the measurement error. The differences between nonradiative rates of these three types porphyrins can be explained by taking into account that their S₀–S₁ gaps change in parallel with their rate constants k_{nr} , i.e., H₂Ph₄TBP \approx H₂Ph₄TCHP < H₂OETPP. However, the observed effect may also be related to higher rigidity of macrocycles annealed with exocyclic fragments, especially with benzo- rings.

It has been shown that photophysical properties of free-base porphyrins change gradually with increasing degree of the macrocycle nonplanarity.^{12a,13h} Therefore, one could expect that tetrabenzoporphyrins substituted with two, as opposed to four, *meso*-aryl groups would exhibit optical properties somewhat average to those of H₂TBP's and H₂Ar₄TBP's. In contrast to this expectation, the absorption spectrum of H₂Ph₂TBP (2) (Figure 1B) appears to be nearly identical to that of H₂TBP (1), with the Q and Soret bands only slightly shifted to the red, by 5 and 10 nm, respectively.³¹

The vibronic structure of the Q-band of H₂Ph₂TBP (2) and the shape of its Soret band are identical to those of H₂TBP (1), except for a small extra peak at 449 nm in the spectrum of the latter. This peak is likely to be caused by an impurity. A similar feature was observed in the spectrum of H₂TBP^{8d} synthesized by the template condensation method.³² The method is known to yield complex mixtures of products. On the other hand, the spectrum of well-soluble octabutoxy-derivative H₂TBP(CO₂Bu)₈ did not show this extra feature (see Supporting Information, III).

The fluorescence properties of H₂TBP (1) and H₂Ph₂TBP (2) are also very close: small Stokes shifts (~ 45 cm⁻¹), long

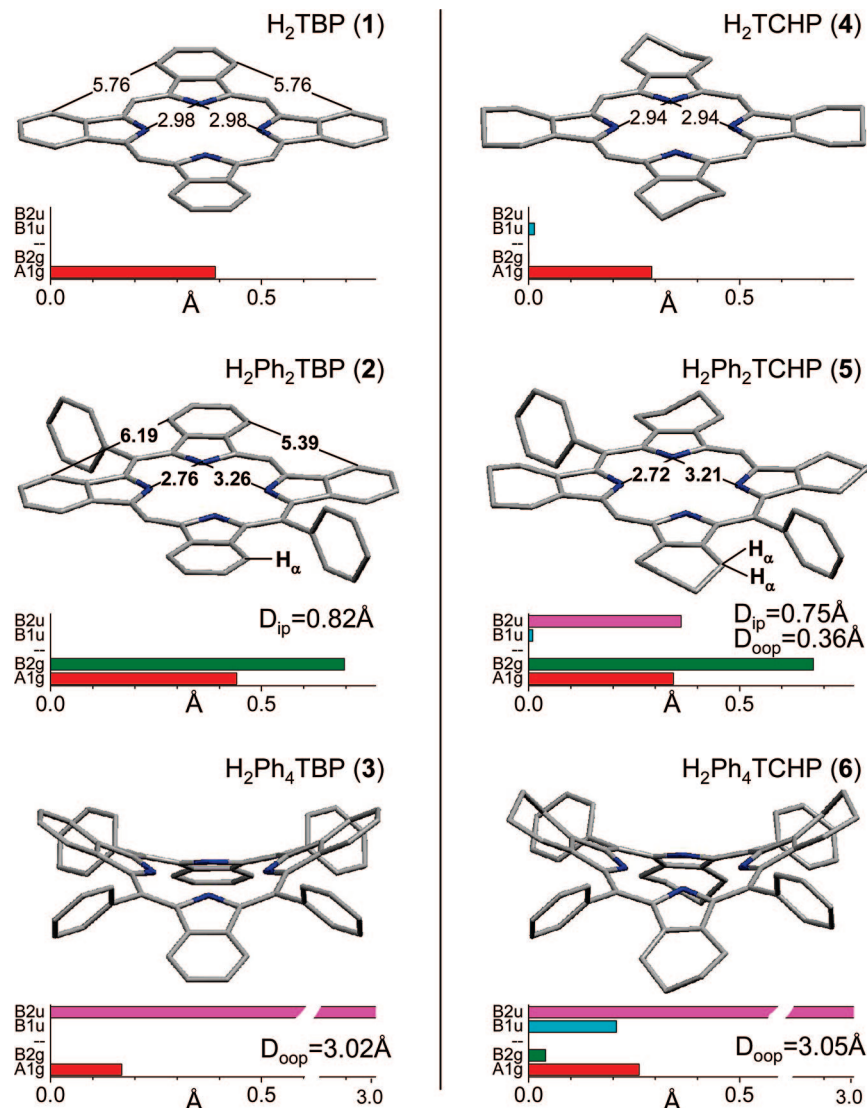


Figure 3. Computed structures (DFT B3LYP/6-31G(d)) of H_2Ph_nTBP 's (left column) and Ph_nTCHP 's (right column) ($n = 0, 2, 4$). Distances are in Å. Main distortion modes, recovered by the NSD analysis,²² are shown by the bar graphs. In-plane distortions: A_{1g} , red; B_{2g} , green. Out-of-plane distortions: B_{1u} (ruffling), cyan; B_{2u} (saddling), magenta. (For complete list of modes see Supporting Information, V). D_{oop} and D_{ip} designate total mean-square out-of-plane and in-plane displacements, respectively.

fluorescence lifetimes (9.8 and 10.3 ns) and high emission yields (0.35 and 0.34), increasing to 0.47 and 0.41, respectively, in DMA.

Similarities in the optical properties of H_2Ph_2TBP (2) and H_2TBP (1) suggest that the structures of these porphyrins should also be similar, i.e. that H_2Ph_2TBP (2) should be planar. On the other hand, based on the published data,^{16,18a,b,28} H_2Ph_4TBP (3) is expected to be strongly nonplanar, presumably saddled. The question arises as to why the interactions between *meso*-aryl rings and benzo-rings in H_2Ph_2TBP (2) do not affect its planarity, whereas the same interactions in H_2Ph_4TBP cause it to become strongly nonplanar?

In order to perform structural comparisons we carried out DFT (B3LYP/6-31G(d)) calculations of porphyrins depicted in Figure 1. The calculated structures are shown in Figure 3 together with their normal distortion modes, as revealed by the NSD analysis.²² It should be mentioned that the X-ray structures of H_2TBP (1) cations (planar),¹⁷ unsubstituted H_2TCHP (anomalously planar dication)²⁹ and $H_2Ph_4TBP(CO_2Me)_8$ (saddled)²⁸ were published in the literature, providing the required reference points. However, for the purpose of the comparative analysis, computed structures may in fact present an advantage, since

X-ray structures are potentially affected by crystal packing forces, which may or may not affect the macrocycle's planarity.

It is seen from the figure that the macrocycle of H_2Ph_2TBP (2) is as planar as that of H_2TBP (1). The total out-of-plane displacements (D_{oop}) are practically zero for both porphyrins. The phenyl rings in H_2Ph_2TBP (1) are strictly orthogonal to the macrocycle plane and, as a result, their effect on the macrocycle electronic system is minor. Indeed, the absorption bands of H_2Ph_2TBP (2) are only slightly red-shifted vs those of H_2TBP (1), and the substituents in the *meso*-phenyl rings, e.g., *para*-methoxycarbonyl groups in porphyrin **2a** (Chart 1), have negligible effect on the peak positions (see Supporting Information, III).

As expected, the structure of H_2Ph_4TBP (3) is strongly nonplanar ($D_{oop} = 3.02$ Å), with B_{2u} -saddling mode being the dominant deformation. H_2Ph_4TCHP (6) is also saddled, but with a contribution of ruffling (B_{1u}). B_{1u} -mode is known to especially strongly enhance nonradiative decay of the S_1 state,^{13b} suggesting that the steeper increase in k_{nr} (vide supra) upon going from H_2TCHP (4) to H_2Ph_4TCHP (6) vs the pair H_2TBP (2)/ H_2Ph_4TBP (3), may be associated with this particular distortion. The *meso*-phenyl rings in H_2Ph_4TBP (3) are tilted with respect

to the macrocycle, and, therefore, are partially drawn into conjugation. As a result, *para*-methoxycarbonyl groups in **3a** shift its spectrum by 4–5 nm to the red relative to H₂Ph₄TBP (**3**) (see Supporting Information, III).

The intrinsic asymmetry of free-base porphyrins manifests itself in all the calculated structures by the A_{1g}-distortion, i.e., the elongation in the direction of the axis connecting the two opposite NH nitrogen atoms. In addition to that, H₂Ph₂TBP (**2**) and H₂Ph₂TCHP (**5**) exhibit strong B_{2g} mode, which is associated with the in-plane rotation of the isoindole fragments, leading to the change in the distances between the opposite pairs of pyrrole N-atoms and yielding more space for the *meso*-aryl substituents. This rotation appears to lift the steric hindrance between the *meso*-aryl groups and the α -hydrogen atoms (H _{α}) of the benzo (H₂Ph₂TBP (**2**)) or cyclohexeno rings (H₂Ph₂TCHP (**5**)). For example, the edges of the rectangular, drawn through the corresponding α -carbon atoms of benzo-rings in H₂Ph₂TBP (**2**), are 5.41 and 6.12 Å ($D_{ip} = 0.82$ Å), while in symmetrical H₂TBP (**1**) the corresponding distances are equal (5.76 Å).

Relief of the steric strain by way of in-plane as opposed to out-of-plane deformation appears to be more energetically favorable, suggesting that it might be a more effective way to preserve the macrocycle aromaticity. Interestingly, the structure of H₂Ph₂TCHP (**5**) in addition to being distorted in-plane ($D_{oop} = 0.75$ Å) also shows a small contribution of saddling (B_{2u}) ($D_{oop} = 0.36$ Å). This distortion is completely absent in the structure of H₂Ph₂TBP (**3**), even though the distance between the H _{α} -atom of the benzo-ring and the plane of the *meso*-phenyl ring (Figure 3) is actually shorter than the average distance between the *meso*-phenyl ring and the H _{α} -atoms of the cyclohexeno-ring in H₂Ph₂TCHP (**5**). Higher resistance of H₂Ph₂TBP (**2**) to nonplanar distortion may be another manifestation of the increased rigidity of TBP macrocycle. In this regard, in a recent theoretical study²⁴ the propensity of *meso*-aryl-substituted porphyrins to undergo nonplanar deformations was considered to be an effect of two counterbalancing forces: one acting to preserve planarity and aromaticity and another striving to bring the *meso*-aryl group into conjugation with the tetrapyrrole system. From this point of view, the former force in H₂Ph₂TBP (**2**) appears to be stronger than in H₂Ph₂TCHP (**5**).

Remarkably, even a small out-of-plane deformation of H₂Ph₂TCHP (**5**) immediately reveals itself in its photophysical behavior. The absorption bands of H₂Ph₂TCHP (**5**) (Figure 2E) are broadened and red-shifted relative to H₂TCHP (**4**), the fluorescence spectrum of H₂Ph₂TCHP (**5**) has a somewhat different shape, and its fluorescence yield (0.08) and lifetime (11.8 ns) are considerably lower than those of H₂TCHP (**4**). The nonradiative rate constant for H₂Ph₂TCHP (**5**) (Table 2) increases by about 30% relative to k_{nr} of H₂TCHP (**4**), and the radiative constant decreases by about 50% relative to k_r of H₂TCHP (**4**). In contrast to these changes, the in-plane distortion in H₂Ph₂TBP (**2**) has practically no influence on the S₁ decay parameters optical properties as compared to H₂TBP (**1**). These results strongly suggest that out-of-plane deformations play the dominant role in the excited-state properties of distorted porphyrins, whereas the role of in-plane deformations is insignificant.

To the best of our knowledge, the X-ray structure of H₂Ph₂TCHP (**2**) is not known. To verify that the computations adequately represented its structural features, we compared the calculated (B3LYP/6-31G(d)) structure of another 5,15-diphenyl-octalkylporphyrin with its published X-ray structure³³ (see Supporting Information, V). The results confirmed that all the main features of the experimental structure were adequately

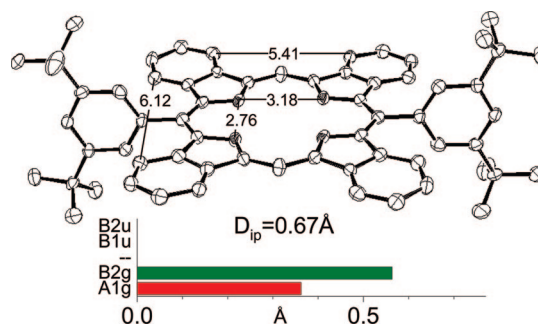


Figure 4. X-ray crystal structure of H₂Ar₂TBP (Ar = 3,5-*t*Bu₂C₆H₃) and its NSD analysis. Distances are in Å. Hydrogen atoms are omitted for clarity.

reproduced by the calculations. It also should be mentioned, that most of the published X-ray structures of free-base β -alkyl-5,15-diarylporphyrins (obtained from Cambridge Structural Database) (see ref 33 and Supporting Information, IV) reveal certain degree of out-of-plane deformations, although in some cases these could be caused by structural elements (e.g., strapping substituents) capable of inducing out-of-plane distortions independent of *meso*-aryl/ β -alkyl interactions.

The experimental evidence of virtually ideal planarity of H₂Ar₂TBP as well as of its strong in-plane distortion was obtained by X-ray crystallography (Figure 4).

The crystal structure of H₂Ar₂TBP (Ar = 3,5-*t*Bu₂-C₆H₃) shows the essentially flat tetrapyrrole skeleton, practically orthogonal *meso*-aryl rings and significant A_{1g} and B_{2g} deformations ($D_{ip} = 0.67$ Å). The N₄-core is asymmetrical with adjacent edges of 3.18 and 2.76 Å. Overall, the structure is in excellent agreement with the calculations.

It follows from the above results that the spectroscopic signature of *meso*-unsubstituted TBP is as a sensitive indicator of planarity of TBP-based systems. For example, it is likely that the structures of H₂ArTBP and 5,10-H₂Ar₂TBP, reported by Senge and Bischoff,³⁴ as well as the structure of H₂ArTBP (Ar = 4-MeO₂C-C₆H₄), reported by Berova et al.,³⁵ are planar. Similarly, *meso*-alkenyl-TBP's, synthesized by Ono et al.,³⁶ also are expected to be planar.

In summary of this section, planar free-base tetrabenzoporphyrins are strongly fluorescent and their radiative rate constants and quantum yields are 3–4 times higher than those of regular planar porphyrins. Nonplanar distortions affect the excited-state properties of free-base tetrabenzoporphyrins in a manner consistent with the model proposed for regular nonplanar porphyrins.¹³ In contrast to out-of-plane deformations, in-plane deformations, induced in tetrabenzoporphyrins by 5,15-diaryl-substitution, have practically no effect on their excited-state properties of tetrabenzoporphyrins.

PdAr_nTBP's and PdAr_nTCHP ($n = 0, 2, 4$). In Pd porphyrins, S₁(π, π^*) \rightarrow T₁(π, π^*) intersystem crossing occurs with almost unity quantum yield.³⁷ Because (*d, d*) transitions of Pd(II) ion are higher in energy than T₁(π, π^*) states, planar Pd porphyrins (e.g., PdOEP, PdTPP) strongly phosphoresce in solutions at room temperature.³⁸

Recently, Zenkevich and co-workers²¹ showed that saddling (B_{2u}) deformations in regular Pd porphyrins, such as PdOETPP,^{39,40} lead to a drastic decrease in their phosphorescence, whereas at low temperatures the phosphorescence is considerably regained ($\phi_{phos} = 0.25$ for PdOETPP at 77 K).^{21b} Having that in mind, we set out to examine how *meso*-aryl substituents in PdAr_nTBP's ($n = 0, 2, 4$) affect properties of their T₁ states. As in the case of free-base porphyrins, Pd complexes of structurally related Ar_nTCHP's ($n = 0, 2, 4$) (Chart 1) were used for comparison.

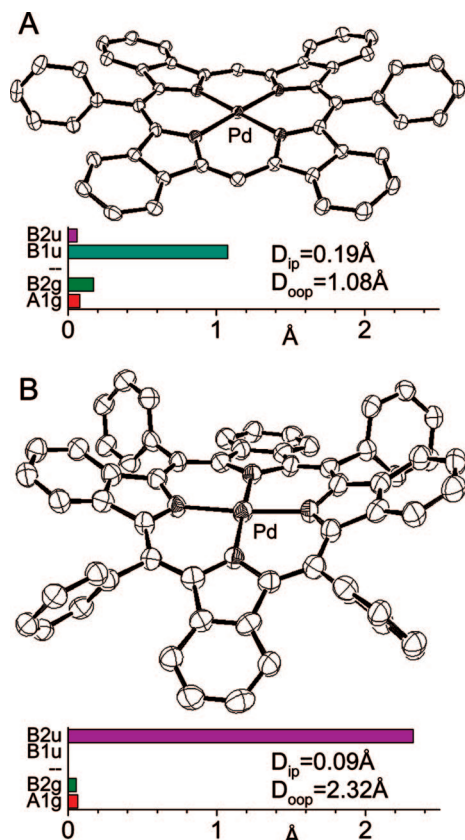


Figure 5. X-ray crystal structures of PdPh₂TBP (**Pd-2**) (A) and PdAr₄TBP (Ar = 3,5-(BuO₂C)₂C₆H₃) (B) and their NSD analyses. Hydrogen atoms and substituents in the *meso*-aryl rings in PdAr₄TBP (B) are not shown for clarity (see Supporting Information, IV for complete structure).

Structural information on Pd porphyrins is not as easily obtainable by *ab initio* methods as for free-bases. Nevertheless, previously published structural data on various types of metallated planar and nonplanar porphyrins allowed us to make the following conjectures. Similar to PdOEP,⁴¹ PdTCHP (**Pd-4**) is expected to have unperturbed planar geometry. Literature data on Cu, Co and Zn tetrabenzoporphyrins^{4b–d} suggest that PdTBP (**Pd-1**) also has planar geometry. Similar to Cu, Zn and Ni OETPP's,⁴⁰ PdPh₄TCHP (**Pd-6**) is expected to be saddled; while based on the existing X-ray data on metallated 5,15-diaryl β -octaalkylporphyrins,⁴² PdPh₂TCHP is expected to be predominantly ruffled.

By analogy with regular 5,15-diarylporphyrins, we expected that metalation of Ar₂TBP's would change the geometry of the N₄-core and, as a result, would lead to a closer contact between benzo and *meso*-aryl rings. Metallated β -alkyl 5,15-diarylporphyrins typically undergo nonplanar B_{1u}-type ruffling deformation,⁴² and the same was expected for PdPh₂TBP (**Pd-2**). The X-ray structure of PdPh₂TBP (**Pd-2**) (Figure 5A) confirmed this expectation. The predominant mode of its out-of-plane distortion (D_{oop} = 1.08 Å) is indeed ruffling (B_{1u}) with a small contribution of saddling (B_{2u}), while the in-plane distortion is much smaller (D_{ip} = 0.19 Å) than in the corresponding free-base H₂Ph₂TBP (**2**) (Figure 1 and 4).

PdAr₄TBP (Ar = 3,5-(BuO₂C)₂C₆H₃) (Figure 5B) is, on the contrary, strongly saddled. The value of the total out-of-plane deformation in PdAr₄TBP (D_{oop} = 2.32 Å) is similar to that determined for the earlier published structure of ZnPh₄TBP·THF (D_{oop} = 2.35 Å),¹⁶ but somewhat lower than in PtPh₄TBP (D_{oop} = 2.83 Å)^{4e} and especially in NiPh₄TBP(CO₂Me)₈ (D_{oop} = 3.43

Å).^{18b} These differences might be caused by the differences in ionic radii of the metals or by the crystal packing forces or by both.

The optical spectra of PdPh_{*n*}TBP's and PdPh_{*n*}TCHP's (*n* = 0, 2, 4) are shown in Figure 6, and the photophysical data are summarized in Table 3. The phosphorescence quantum yields and decays were measured in Ar-purged DMA solutions at 295 K, and the phosphorescence decays were found to be purely single-exponential. Porphyrin triplet states are extremely sensitive to various quenching processes, and that makes it difficult to compare the data on the phosphorescence of Pd porphyrins measured in different laboratories using different solvents and different deoxygenation methods. Therefore, in our experiments we attempted to maintain identical conditions (solvent, Ar purity, control of air displacement) for all the samples measured; however, the reported lifetimes and quantum yields should not be considered absolute.

Unsubstituted PdTBP (**Pd-1**) has very low solubility and shows tendency to aggregation. To account for possible effects of aggregation, a well-soluble PdTBP derivative substituted with eight butoxycarbonyl groups, PdTBP(CO₂Bu)₈ (**Pd-1a**), was also studied. It should be mentioned that insertion of Pd into Ar_{*n*}TCHP's often leads to the appearance of extra bands above 600 nm in the absorption spectra. These bands are most likely associated with partially oxidized products, i.e. mono-, di- and tribenzoporphyrins. Special care needs to be taken to avoid contamination by these side products, and only freshly prepared and purified samples of PdPh₂TCHP (**Pd-5**) must be used in photophysical measurements.

The changes in the ground-state absorption spectra of PdAr_{*n*}TBP's and PdAr_{*n*}TCHP's upon increase in *n* from 0 to 4 resemble the trends observed for the corresponding free-bases: red-shifting, broadening and, in the case of PdAr_{*n*}TCHP's, gradual change in the relative intensity of Q_x vs Q_y band. As expected, the spectrum of PdAr₄TCHP (**Pd-6**) was found to be similar to the published spectrum of PdOETPP.²¹ Noteworthy, there is a slight splitting of the Q-band (λ_{max} = 615 nm) of PdAr₂TBP (**Pd-2**), a feature absent in the spectrum of PdTBP (**Pd-1**).

The phosphorescence spectra in general follow the changes in the absorption. One exception is that the main peak of the phosphorescence of ruffled PdPh₂TBP (**Pd-2**) (λ_{max} = 796 nm) is located almost at the same position as that of saddled PdPh₄TBP (**Pd-3**) (λ_{max} = 797 nm), while the Q-bands shift gradually in the order PdTBP (**Pd-1**) → PdPh₂TBP (**Pd-2**) → PdPh₄TBP (**Pd-3**). As a result, the phosphorescence Stokes shift of PdPh₄TBP (**Pd-3**) is less by more than 200 cm⁻¹ compared to the two less substituted porphyrins.

The major difference between PdAr_{*n*}TBP's and PdAr_{*n*}TCHP's is in how their phosphorescence quantum yields and lifetimes respond to the changes in the macrocycle nonplanarity. Just like in the case of PdOETPP,²¹ saddling deformation in PdAr₄TCHP (**Pd-6**) leads to the complete loss of its emissivity. In contrast, saddled PdAr₄TBP (**Pd-3**) phosphoresces with quite high quantum yield (ϕ_{phos} = 0.08), which is only 1.6 lower than that of planar PdTBP (**Pd-1**) (ϕ_{phos} = 0.13). It is possible that the phosphorescence yield and lifetime of PdTBP (**Pd-1**) are affected (lowered) by aggregation. Indeed, the phosphorescence of much better soluble octabutoxycarbonyl-derivative **Pd-1a** was found to be significantly stronger (ϕ_{phos} = 0.35); but still, the difference between the quantum yields of **Pd-1a** and PdAr₄TBP (**Pd-3**) is not nearly as drastic as for PdPh₄TCHP (**Pd-6**) vs PdTCHP (**Pd-4**).

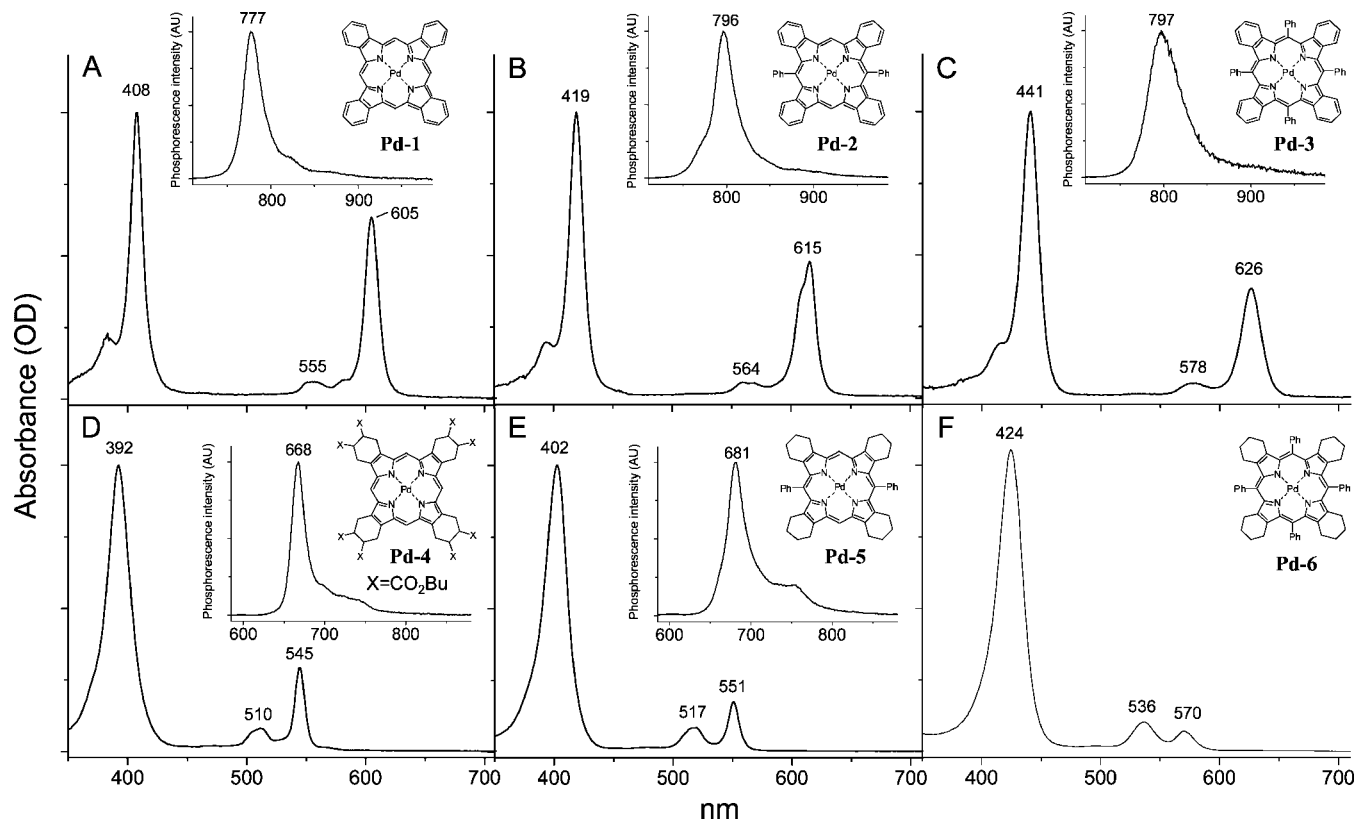


Figure 6. Absorption and phosphorescence (insets) spectra of PdTBP (**Pd-1**, A), PdPh₂TBP (**Pd-2**, B), PdPh₄TBP (**Pd-3**, C) and PdTCHP(CO₂Bu)₈ (**Pd-4**, D), PdPh₂TCHP (**Pd-5a**, E) and PdPh₄TCHP (**6**, F) in DMA at 295 K.

TABLE 3: Photophysical Properties of PdAr_nTBP and PdAr_nTCHP (*n* = 0, 2, 4) in Deoxygenated DMA at 295 K

Pd porphyrin	phosphorescence λ_{max} (λ_{ex}), nm	ϕ_{phos}^a	τ_{phos} (μs) ^b
TBP			
Pd-1	777 (605)	0.13	399
Pd-1a	770 (615)	0.35	496
Pd-2	796 (615)	0.15	423
Pd-3	797 (626)	0.08	258
TCHP			
Pd-4	668 (545)	0.45	1118
Pd-5	681 (550)	0.02	97
Pd-6	—	— ^c	— ^c

^a Measured relative to the fluorescence of H₂TTP in deoxygenated C₆H₆ (ϕ_{fl} = 0.11).²⁵ Error in determination of yields is no more than 10%. ^b Phosphorescence decays were nearly ideally single-exponential, as revealed by the Maximum Entropy analysis. ^c No signal could be detected.

Both radiative and nonradiative rate constants for PdTBP (**Pd-1a**) are somewhat higher than those of PdTCHP (**Pd-4**) (Table 4). In response to nonplanar deformation, the radiative rate in PdPh₄TBP (**Pd-3**) decreases slightly relative to that of PdTBP (**Pd-1a**), resembling the trend in the behavior of the corresponding free-bases. Remarkably, upon going from PdTBP (**Pd-1a**) to PdPh₄TBP (**Pd-3**) the nonradiative rate rises also by only about 2.7 times, whereas in PdPh₄TCHP (**Pd-6**) k_{nr} is apparently so large, that this porphyrin completely loses its phosphorescence.

For PdPh₂TBP (**Pd-2**) and PhPh₂TCHP (**Pd-5**) the differences are also quite striking. Upon going from planar PdTCHP (**Pd-4**) to presumably ruffled PdPh₂TCHP (**Pd-5**), the phosphorescence quantum yield decreases by as much as 22.5 times, while in the case of PdTBP (**Pd-1**) vs PdPh₂TBP (**Pd-2**) it seems to even slightly increase. Again, this can be related to the partial

TABLE 4: Deactivation Rate Constants for T₁ state of PdAr_nTBP and PdAr_nTCHP (*n* = 0, 2, 4) in Deoxygenated DMA at 295 K

compound	$k = 1/\tau_{\text{phos}}$ (s ⁻¹)	k_{r} (s ⁻¹) ^a	k_{nr} (s ⁻¹) ^b
TBP			
PdTBP (Pd-1)	2.51×10^3	3.26×10^2	2.18×10^3
Pd-1a	2.02×10^3	7.06×10^2	1.31×10^3
PdPh ₂ TBP (Pd-2)	2.36×10^3	3.55×10^2	2.00×10^3
PdPh ₄ TBP (Pd-3)	3.88×10^3	3.10×10^2	3.57×10^3
TCHP			
PdTCHP (4)	8.94×10^2	4.02×10^2	4.92×10^2
PdPh ₂ TCHP (Pd-5)	1.03×10^4	2.06×10^2	1.01×10^4

^a k_{r} , radiative decay rate constant, calculated as $k_{\text{r}} = \phi_{\text{phos}}k$, where ϕ_{phos} is the phosphorescence quantum yield. ^b k_{nr} , nonradiative decay rate constant ($k_{\text{nr}} = k_{\text{ic}} + k_{\text{isc}}$), calculated as $k_{\text{r}} = k - k_{\text{nr}}$.

aggregation of PdTBP (**Pd-1**); but even in comparison with octabutoxycarbonyl-derivative **Pd-1a**, the quantum yield of PdPh₂TBP (**Pd-2**) is only 4 times lower. Notably, this decrease in the quantum yield is largely due to the decrease in the radiative rate constant, while the increase in the nonradiative rate is only 1.5 times.

Overall, the photophysical data clearly show that π -extension strongly affects triplet state properties of Pd porphyrins. Nonplanarity, especially saddling, greatly promotes nonradiative decay of the T₁ states of regular nonplanar Pd porphyrins, rendering them completely nonemissive. On the contrary, either saddling (B_{2u}) or ruffling (B_{1u}) deformations affect the T₁ states of tetrabenzoporphyrins only slightly; both PdAr₂TBP's and PdAr₄TBP's remain strongly phosphorescent at ambient temperatures.

Disappearance of the phosphorescence in regular nonplanar porphyrins in principle can be caused by strong enhancement

of $S_1 \rightarrow S_0$ internal conversion, leading to the reduction in the yield of the T_1 state. However, literature data suggest that the enhancement of internal conversion in nonplanar porphyrins occurs in parallel with the enhancement of the intersystem crossing.^{12c} In the presence of Pd, $S_1 \rightarrow T_1$ decay should become even faster, and, as a result, it is likely to reach the rate at least as high as that of the internal conversion. Therefore, the key factor influencing phosphorescence of nonplanar Pd porphyrins is most likely the rate of $T_1 \rightarrow S_0$ decay.

The rates of radiative $T_1 \rightarrow S_0$ transitions are somewhat higher for planar PdTBP's than for planar PdTCHP's, and upon nonplanar deformations they slightly decrease (vide supra). The nonradiative $T_1 \rightarrow S_0$ rates increase sharply in response to nonplanar deformations in PdTCHP's, but only slightly in PdTBP's. The fact that the phosphorescence of nonplanar Pd porphyrins, e.g. of PdOETPP, is regained at 77 K²¹ suggests that the enhancement of the nonradiative decay is associated with vibrational activity. Thus, vibrations greatly enhance $T_1 \rightarrow S_0$ intersystem crossing in PdAr₄TCHP's, but have only small effect on the intersystem crossing in PdAr₄TBP's.

The rate of $T_1 \rightarrow S_0$ intersystem crossing is determined by the combination of three factors: spin–orbit coupling due to the heavy atom effect; interactions of Pd *d* orbitals with T_1 and S_0 states, resulting in their partial mixing; and Franck–Condon factors associated with the participating vibrational levels of the states involved in the transition.^{21,43} It is likely that some of the low-energy out-of-plane vibrations near the equilibrium point on the T_1 potential energy surface facilitate spin–orbit coupling and enhance the nonradiative decay.⁴³ Even if favorable for intersystem crossing conditions are not met near the equilibrium, there are likely to be conformers (local minima) on the T_1 surface characterized by higher magnitudes of spin–orbit coupling. High conformational flexibility of nonplanar Pd porphyrins is likely to be retained in their T_1 states,²⁰ and the conformers with increased magnitude of spin–orbit-coupling can be viewed as analogs of “funnels” proposed for nonradiative deactivation of S_1 states.¹³ Considering low $T_1 \rightarrow S_0$ radiative rates, the molecule has enough time to visit all these local minima, including the funnels, and undergo efficient nonradiative $T_1 \rightarrow S_0$ decay.

The described above experiments suggest that the conformations characterized by high rates of intersystem crossing are not as easily attainable by PdAr₄TBP's as by PdAr₄TCHP's. On the one hand, this can be a result of lower flexibility of the PdAr₄TBP system and, therefore, its limited ability to traverse the T_1 energy surface. However, the data on free-base Ar₄TBP's (vide supra) suggest that even if there is an increase in the rigidity of the TBP macrocycle compared to that of TCHP, the difference is not large.

Alternatively, it is possible that for all possible conformers of PdAr₄TBP's the magnitude of spin–orbit coupling is never as high as for some selected conformers of PdAr₄TCHP's. Should this indeed be true, the reason for much weaker effects of nonplanar deformations on the phosphorescence of PdAr₄TBP's vs PdAr₄TCHP's could be explained by the difference between the spatial distributions of b_1 (HOMO) and e (LUMO) orbitals of tetrabenzoporphyrins vs regular porphyrins.¹⁰ Indeed, b_1 HOMO and e_g LUMO orbitals in Zn and Pd Ph₄TBP's were shown to be partially moved out from the pyrroles into benzo-rings,¹⁰ away from the metal center. As a result, interactions of these orbitals with *d*-orbitals of Pd even in out-of-plane distorted macrocycle may be much weaker than in PdAr₄TCHP's.

An interesting observation was made that the triplet lifetimes of Pd porphyrins (both TBP's and TCHP's) are affected by ruffling much less than by saddling. Noteworthy, the structures of highly phosphorescent PdTPP⁴⁴ and Pd *meso*-tetracarboxyphenylporphyrin⁴⁵ apparently are also quite ruffled. In the case of S_1 states of free-base porphyrins, however, the situation is exactly the opposite, i.e. nonradiative decay in ruffled porphyrins is much faster than in saddled porphyrins (vide supra). In fact, the very fast internal conversion of ruffled free-base porphyrins^{13b} is perhaps one of strongest indications of the existence of funnels on their potential energy surfaces. Either analogous minima do not exist on T_1 surfaces of ruffled Pd porphyrins, or spin–orbit coupling in these minima is inefficient. A separate study will be required to understand this phenomenon.

Conclusion

The presented data provide initial experimental framework for understanding mechanisms of photophysical processes occurring in planar and nonplanar π -extended porphyrins. The performed phenomenological analysis can be useful for building empirical correlations between structures and photophysical properties of these molecules and forming the initial picture; however, comprehensive computational work, such as recently performed on free-base porphin and its Mg and Zn complexes,⁴³ will be required to fully delineate the effects of the π -extension on the excited properties of nonplanar tetrapyrroles.

From practical point of view, the described photophysical measurements allowed identification of a subset of π -extended porphyrins, namely 5,15-diaryltetrabenzoporphyrins, which combine powerful near-infrared absorption and strong emission from singlet and triplet states with ease of derivatization via substitution in the *meso*-aryl rings. Convenient functionalization not leading to alterations in photophysical properties is the key to many potential applications of these chromophores, such as biomedical imaging and PDT.

Experimental Section

All porphyrins used in this study were synthesized and purified as described previously,¹⁸ except for complexes **Pd-4**, **Pd-5**, **Pd-5a** and **Pd-6**. The latter were obtained by refluxing the corresponding free-base porphyrins with Pd(OAc)₂ in acetic acid and purified by column chromatography (see Supporting Information, II for details).

Absorption and emission spectra, fluorescence and phosphorescence quantum yields and lifetimes were measured using standard methods (see Supporting Information, I). Complete X-ray crystal structures are available in the Cambridge Crystallographic Data Centre (CCDC): 665410, 681173, 687685. For details of DFT calculations see Supporting Information, V. NSD analysis²² was performed using an online Java applet available at the Web site of Prof. John A. Shelnutt: <http://jasheln.unm.edu/jasheln/content/nsd/NSDengine/start.htm> (see Supporting Information, V for details).

Acknowledgment. Support of the grants EB007279 and HL081273 from the NIH USA and RFBR-04-03-32650 and RFBR-07-03-01121 from the Russian Foundation of Basic Research is gratefully acknowledged. We thank Dr. Thomas Troxler (Penn RLBL, NIH P41-RR001348) for assistance with fluorescence lifetime measurements, Drs. K. Lutsenko (INEOS RAN) and Patrick Carroll (Penn) for X-ray structure determination and Prof. Craig Medforth (University of New Mexico) for providing structural data on 5,15-diaryl- β -octaalkylporphyrins. SAV thanks Prof. Robin M. Hochstrasser for invaluable discussions.

Supporting Information Available: Description of synthesis, details of photophysical experiments, computations and X-ray structure determination, as well as characterization data for the new compounds. This material is available free of charge via the Internet at <http://pubs.acs.org>.

References and Notes

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JP8043626