

Highly Sensitive Halide Ions Recognition with Diprotonated Porphyrin

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The titration of the diprotonated form of 3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin with the series of halide ions has been done in the acetonitrile solution. It was shown that the stable complexes of 1:1 and 1:2 structures with halide ions have been formed. The influence of complexation with halide ions on the porphyrin fluorescent properties has been studied. The strong quenching of the porphyrin fluorescence has been found. It was established that both static quenching upon the nonfluorescent complex formation due to internal heavy atom effect and dynamic diffusion-controlled quenching took place. Based on the analysis of dependencies of fluorescence intensity and lifetime on the iodide ions concentration in solution, it was proposed to use the diprotonated form of the porphyrin as a basic compound for design of the fluorescent molecular receptor for the halide ions.

Introduction

The design of new efficient means for real-time detection of chemical and biochemical species through the use of sensors is among the most significant fields in modern science.^[1] The nature of the molecular sensing agent used to generate the chemical diagnostic signal upon the recognition of the given substance is central to determining the overall performance of any chemical sensor. The way of transduction of this chemical signal into appropriate analytical signal is also of great importance since it determines the ease of production and exploitation of the sensing device. The optical transduction seems to be the most attractive for several reasons which include inherent safety, less noise pickup in signal transmission over long distances, and the possibility of obtaining much more comprehensive information from a single probe.

The representatives of the tetrapyrrolic compounds family can serve as good candidates for optical transduction. They are known to have rather strong luminescence (fluorescence and/or phosphorescence) that can be tuned in the appropriate spectral range by modification of the molecule. The sensitivity of the rate constant and quantum yield of porphyrin luminescence to the properties of molecular microenvironment provides the possibility to use tetrapyrrolic molecules not only as optical transducers but as sensing agents also. Design of the molecular sensor can be simplified in such a case, since the same molecule combines both functions.

Anions play a fundamental role in a wide range of chemical and biological processes, and numerous efforts have been devoted to the development of abiotic receptors for anion species,^[2] but they are still very limited. The detection of different anions in the solution can be achieved with using of tetrapyrrolic compounds. The diprotonated forms of tetrapyrrolic compounds are able to form adducts with the organic and inorganic acid residues^[3-5] and to form

complexes with salt anions.^[6,7] The large efforts has been made in the elucidation of the molecular conformation,^[3,5,8-16] acid-base equilibria involved (^[17-21] and Refs. therein), ground and excited states properties^[4,15,20-24] of the diprotonated porphyrins. The essential progress has been achieved in this area. It was found, that upon protonation of the macrocycle core the nonplanar conformation is adopted.^[3,5,8-16] The type and extent of macrocycle nonplanar distortions have been extensively discussed in terms of molecular flexibility,^[5,10] pattern of the peripheral substitution,^[5,8,10,15,16,22] strength of intermolecular interactions with acid residues.^[3,5,11,12,15] The relationship between the molecular structure and macrocycle core acidity-basicity equilibrium has been studied,^[18-21] and the intermediate monoprotonated form has been characterized for several tetrapyrrolic compounds.^[20,21,25-27] The dramatic changes in the rates and channels of the excitation energy deactivation have been documented in going from the free base to diprotonated compounds,^[4,20-23] and different mechanisms involved in these changes were proposed.^[4,15,22-24]

One of the most interesting issues from point of view of possible practical applications is the interaction of diprotonated macrocycle with anions. The X-ray studies,^[3,5] NMR, IR, Raman and UV-vis spectroscopies,^[6,7,25,28] theoretical calculations^[11,12,15,24] have been applied to study this problem. It was established that both in solid state and in liquid organic solutions the hydrogen bonding between the positively charged core protons and anions occurs,^[3-5,15,16] and the properties of this bond has been extensively discussed. The most of these studies have focused mainly on the structural aspects^[3,5,15] or the consequences of this hydrogen bond formation for the photophysical properties of diprotonated compounds.^[3,22,24] The detailed studies of the complexation constants and the relationship between amount of the salt added and the spectral response of porphyrin have received much less attention.^[7,30-32] In this

work we report the experimental results on the recognition of halide ion salts in solution with the diprotonated form of 3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin (Figure 1). The fluorometric and spectropotentiometric titration experiments have been done. The mechanisms of the diprotonated porphyrin fluorescence quenching upon complexation with halide ions are studied. We discuss the possibility of using of diprotonated porphyrins as the basic compounds in the design of new efficient fluorescent molecular sensors for recognition of halide ions and measurements of halide ion salts concentration in the solutions.

Experimental

Materials and Methods

3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin (Figure 1) was prepared using method described elsewhere.^[33] The identity of the structure was checked by spectroscopic methods and was found to correspond to the literature data.^[34] The halide salts and perchloric acid solutions were prepared with carefully dried acetonitrile. Acetonitrile was purified with dynamic drying on the column with neutral aluminium oxide and with following rectification.^[35] Potassium iodide (KI) and sodium iodide (NaI) were dried under vacuum (1.33 Pa) at 393 K during 48 hours until the constancy of mass has been achieved. The perchloric acid as 8.75 M water solution was used for the preparation of the 0.01 M HClO₄ solution in dried acetonitrile (water content in dried acetonitrile was no more than 0.01 M). The total amount of water added to the solutions in the most acidic solution (pH = 4.5) was calculated to be about 0.011 M. The value was estimated by the method of Fischer titration. Tetraethylammonium bromide (Et₄NBr) was recrystallized twice from acetonitrile and dried in vacuum-desiccator over NaOH during 24 hours at 293 K. Tetraethylammonium chloride (Et₄NCl) was recrystallized twice from acetonitrile and dried under vacuum (1.33 Pa) during 24 hours at 293 K. The halide salts were chosen taking into account their dissociation degrees in acetonitrile.^[36-39]

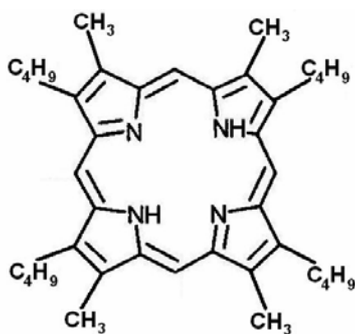


Figure 1. Structure of 3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin.

The spectrophotometric titration of the diprotonated species H₄P²⁺ with halide ion salts was carried out at pH = 6.5, which has been maintained with microaddition of 0.05 N solution of perchloric acid in acetonitrile. At the each titration step the analytical concentrations of reagents were corrected for dilution according to the equation $A_T = A_T^0 / (1 + \sum \Delta V / V_0)$, where V_0 is an initial solution volume, $\sum \Delta V$ is a sum of titrants, A_T is a current optical density value at the analytical wavelength taking into

account the titrants added, A_T^0 is a measured current optical density at the analytical wavelength.

For the fluorescent experiments the porphyrin powder were dissolved in the acetonitrile that contained the chloric acid in the amount required for the complete transformation of all the porphyrin molecules from free base form into diprotonated form (pH = 6.5).^[31] Titration of the porphyrin samples with potassium iodide was done with addition of the small amounts of the concentrated potassium iodide solution that were prepared in the same acidified acetonitrile as the porphyrin sample.

The concentration of diprotonated porphyrin in solutions was determined spectrophotometrically with known extinction coefficients ($\epsilon = 8710 \text{ M}^{-1}\text{cm}^{-1}$ at 585 nm, $\epsilon = 28180 \text{ M}^{-1}\text{cm}^{-1}$ at 543 nm, $\epsilon = 102330 \text{ M}^{-1}\text{cm}^{-1}$ at 403.7 nm^[34]) and was $\sim 2 \cdot 10^{-6}$ and $1.5 \cdot 10^{-5} \text{ M}$ in the fluorometric and spectropotentiometric titration experiments, respectively. The cell for the studies of the acid-base equilibria and the procedure of the spectropotentiometric titration in acetonitrile were described in details.^[30] Electronic absorption spectra were measured with CARY 500 Scan (Varian).

The fluorescence measurements was done in the standard quartz rectangular cells (1×1 cm, Hellma) in the air equilibrated solutions at $293 \pm 2 \text{ K}$. The deoxygenated solutions were used for the measurements of the fluorescence quantum yield Φ_f of the diprotonated form of 3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin (hereafter referred as H₄P²⁺). Deoxygenation of the solutions has been performed with argon bubbling during 20 min just before measurements. The fluorescence spectra and fluorescence excitation spectra were measured using of spectrofluorometer SFL-1211 (Solar). The fluorescence decay kinetics was measured with the using of photon counting system (Edinburgh Instruments). The fluorescence quantum yield Φ_f was determined using the standard sample method.^[40] The free base 5,10,15,20-tetraphenylporphyrin (H₂TPP) was taken as a standard sample ($\Phi_f = 0.11$ ^[41]).

Results and Discussions

Acid-base Equilibria of Macrocycle

Prior the studies of porphyrin interactions with halide ions the acid-base equilibria of 3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin have been studied with the use of the spectropotentiometric titration method. Titration was carried out in acetonitrile with 0.01 M perchloric acid solution. In this case the perchloric acid is completely dissociated and porphyrin protonation occurs by means of solvated proton ($\text{p}K_a^{298} = 2.8$ ^[42,43]). The corresponding titration curve reveals two steps (Figure 2).

This observation indicates that successive formation of the monoprotated H₃P⁺ and diprotonated species H₄P²⁺ takes place upon increase in the concentration of perchloric acid. The difference in the equilibrium constants for the formation of mono- and diprotonated species (K_{a1} and K_{a2}) is three orders of magnitude ($K_{a1} = (1.66 \pm 0.05) \cdot 10^{12} \text{ M}^{-1}$ and $K_{a2} = (1.66 \pm 0.05) \cdot 10^9 \text{ M}^{-1}$). Therefore for the given system “porphyrin – acetonitrile – perchloric acid” three pH ranges are available in which the studied molecules exist either exclusively in the free base H₂P (pH > 15) or diprotonated H₄P²⁺ forms (pH ≤ 7) or mainly in the monoprotated H₃P⁺ form (pH = 9 - 11). The electronic absorption spectra in the Q band range for each of the free base H₂P, monoprotated H₃P⁺ and diprotonated H₄P²⁺ forms are shown in Figure 3.

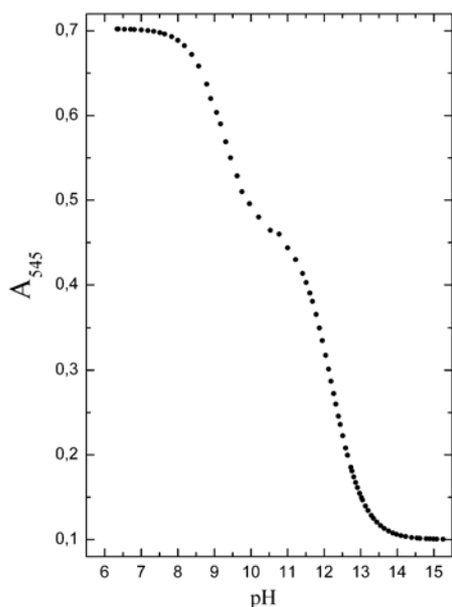


Figure 2. Absorption titration curve at $\lambda = 545$ nm for 3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin in acetonitrile at 298 K.

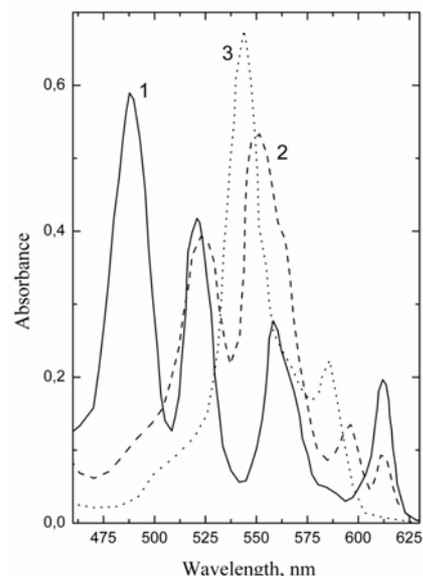
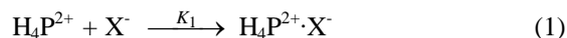


Figure 3. Electronic absorption spectra in the *Q*-band region: free base H_2P (1), monoprotonated H_3P^+ (2) and diprotonated H_4P^{2+} (3) forms. Porphyrin concentration is $1.5 \cdot 10^{-5}$ M.

Spectropotentiometric Titration with Halide Ions

The diprotonated form H_4P^{2+} was titrated with three halide ion salts. Electronic absorption spectra (Figure 4) indicate that increase of halide ion concentration $[X^-]$ leads to the red shift of the porphyrin absorption bands indicating that the complexation takes place. The two sets of the isobestic points during the transformation of the absorption spectra were identified for each of the studied halide ions. These isobestic points are related to the equilibria (1) and (2) respectively:



where the constant K_1 is referred to the formation of porphyrin-halide ion complex with 1:1 ratio, and the constant K_2 is related to the complex with 1:2 ratio. The isobestic points for the equilibrium (1) have been measured as 526 and 553 nm, 548 and 590 nm, 549 and 590 nm in case of iodide, bromide and chloride ions, respectively. For equilibrium (2) the isobestic points are 528 and 555 nm, 550 and 592 nm, 552 and 593 nm, in case of iodide, bromide and chloride ions, respectively.

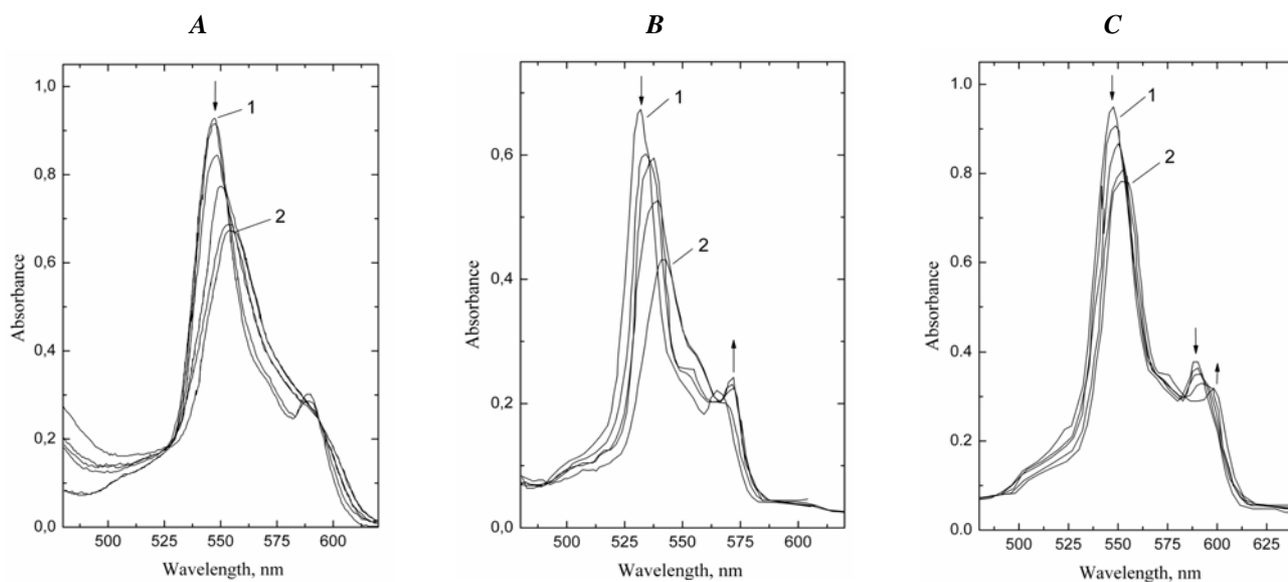


Figure 4. The transformation of the electronic absorption spectra of H_4P^{2+} upon the formation of complexes with halogenide ions in acetonitrile at $pH = 6.5$: A - titration with NaI, 1 - H_4P^{2+} , 2 - $H_4P^{2+} \cdot 2I^-$; B - titration with Et_4NBr , 1 - H_4P^{2+} , 2 - $H_4P^{2+} \cdot 2Br^-$; C - titration with Et_4NCl , 1 - H_4P^{2+} , 2 - $H_4P^{2+} \cdot 2Cl^-$. Arrows indicate the direction of the absorbance changes upon formation of the complexes.

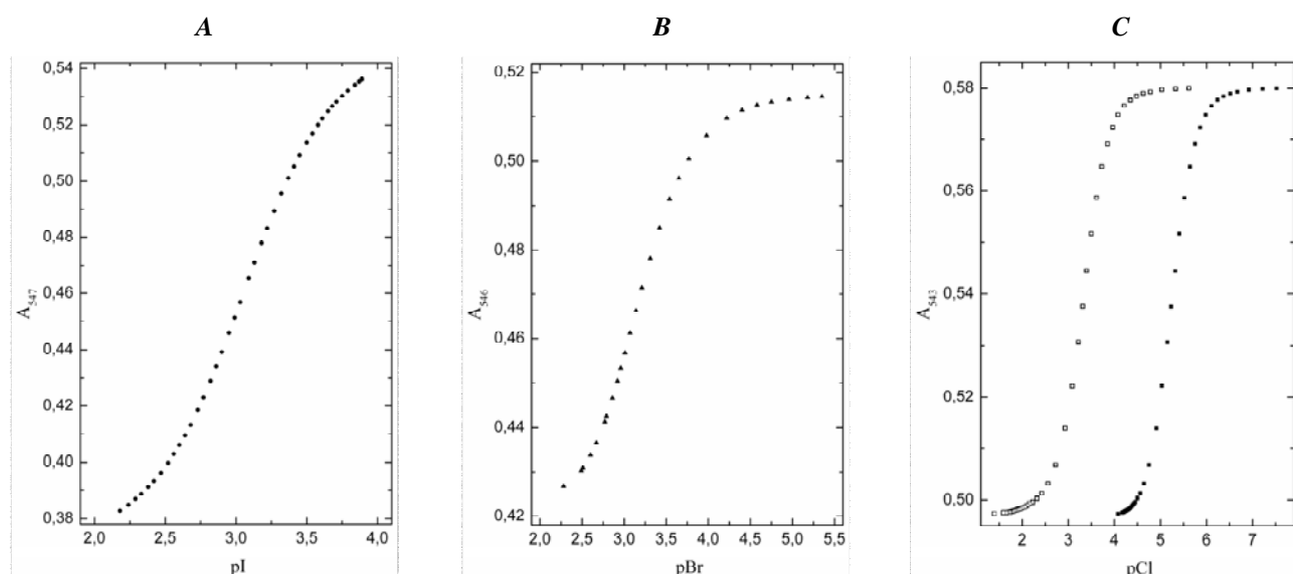


Figure 5. The dependence of the absorbance of the H_4P^{2+} on the concentration of the halide salts in acetonitrile at 298 K: A - NaI, B - Et_4NBr and C - Et_4NCl . Open and closed symbols in case of Et_4NCl denote the total concentration of salt and the concentration of nonaggregated solvated chloride ions, respectively.

The formation of complexes of diprotonated porphyrin with halide ions was analyzed with the titration curves (Figure 5). There are no two sharp well separated steps at the titration curves which correspond to the equilibria (1) and (2). Such behaviour is characteristic when the constants K_1 and K_2 of the conjugated equilibria differ less than two orders of magnitude.^[44] However, the values of the

$$A = \frac{A_{H_4P^{2+}} + A_{H_4P^{2+} \cdot X^-} \cdot [X^-] \cdot K_1 + A_{H_4P^{2+} \cdot 2X^-} \cdot [X^-]^2 \cdot K_1 \cdot K_2}{1 + [X^-] \cdot K_1 + [X^-]^2 \cdot K_1 \cdot K_2} \quad (3)$$

equilibrium constants K_1 and K_2 have been calculated with the nonlinear least-square method for the equation (3)

where $A_{H_4P^{2+}}$, $A_{H_4P^{2+} \cdot X^-}$, $A_{H_4P^{2+} \cdot 2X^-}$ are the optical densities measured for the solutions of uncomplexed diprotonated species H_4P^{2+} , monoligated $H_4P^{2+} \cdot X^-$ and diligated $H_4P^{2+} \cdot 2X^-$ complexes, respectively. During least-square fitting $A_{H_4P^{2+}}$ and $A_{H_4P^{2+} \cdot 2X^-}$ values were taken as they have been measured experimentally, and $A_{H_4P^{2+} \cdot X^-}$ value was the variable parameter. The obtained values for the equilibrium constants are presented in Table.

Table. The equilibrium constants K_1 and K_2 for the formation of the $H_4P^{2+} \cdot X^-$ and $H_4P^{2+} \cdot 2X^-$ complexes in acetonitrile at 298 K.

Halide ion	K_1, M^{-1} ^a	K_2, M^{-1} ^a
I ⁻	$1.20 \cdot 10^3$	$1.07 \cdot 10^3$
Br ⁻	$2.46 \cdot 10^3$	$1.23 \cdot 10^3$
Cl ⁻	$4.08 \cdot 10^3$	$1.27 \cdot 10^3$
	$2.51 \cdot 10^5$ ^b	$2.01 \cdot 10^5$ ^b

^a The error in determination of the K_1 and K_2 values is 3-5%.

^b The values are calculated taking into account the aggregation of chloride ions (see text).

As for the possible complexation of ClO_4^- with porphyrin, it can be excluded since the experiments

evidence that the complexation constant is very low and the large excess of acid is required to achieve the complexation.^[5] In our case the concentration of acid was very low and seems to be insufficient for complexation.

The measured equilibrium constant is strongly influenced by the state of the given ion in the solution. In the case of incomplete dissociation of salts or due to the aggregation phenomena (ion pair formation^[45]) the measured equilibrium constants values may deviate strongly from the actual ones. The aggregation of the halide ions has been studied in the separate experiments using the specific ion electrodes Crutur 52-27, 35-27, 17-27. The results of this study indicate that in the acetonitrile solution at $pH = 6.5$ the formation of multiple H_nCl_m aggregates (namely, HCl , H_2Cl^+ , HCl_2^- and HCl_3^{2-}) takes place. The special procedure has been used to evaluate the true chloride ion concentration.^[31] In the Table both the apparent (uncorrected for the aggregation) and true equilibrium constants are given for the $H_4P^{2+} \cdot 2Cl^-$ complex.

The presented data indicate that the stability of both $H_4P^{2+} \cdot X^-$ and $H_4P^{2+} \cdot 2X^-$ complexes in acetonitrile increases when the halide ion radius decreases. The complexation constants for the chloride (Cl^-) complexes are about 100 times higher than those for bromide (Br^-) and iodide (I^-) complexes. Low complexation constants for two last halide ions are likely due to their larger ionic radii as compared with those for the former. These results are in line with the conclusions on the relationship between the size of halide ion and the contribution from orbital and electrostatic interactions energy into the hydrogen bond formation within $H_4P^{2+} \cdot 2X^-$ complexes.^[15] With the relation to the application of the diprotonated form of porphyrin for the ions detection in the solution, the ions aggregation leads to decrease in the sensitivity of measurements. This is because of the porphyrin recognizes the free solvated ions only, and the ionic aggregates (mentioned above) are not bound by diprotonated porphyrin molecule.

It is significant to note that the monoprotonated H_3P^+ molecules do not form stable complexes with halide ions in the same conditions. We have observed the transformation of the H_3P^+ absorption spectrum into spectrum characteristic for $H_4P^{2+} \cdot 2X^-$ during titration of H_3P^+ with halide salts. The calculation of current concentrations of the H_2P , H_3P^+ , H_4P^{2+} , $H_4P^{2+} \cdot X^-$ and $H_4P^{2+} \cdot 2X^-$ are in good agreement with the results of the H_3P^+ titration by halide salts. The anion-induced protonation of monoprotonated species^[7] takes place first and the newly formed diprotonated species bind anions.

Structure of Complexes with Halide Ions

The molecular structure of $H_4P^{2+} \cdot 2X^-$ complexes with halide ions was the subject of numerous experimental and theoretical investigations.^[3,5,8-16] It was shown^[3] that Cl^- ions are symmetrically aligned exactly at the porphyrin macrocycle symmetry axis at two sides of the porphyrin plane. The similar results have been obtained later for the complexes of several porphyrins with perchlorate ions ClO_4^- .^[5] The length of the hydrogen bond between the porphyrin core protons and Cl^- ions in the complexes $H_4TPP^{2+} \cdot 2Cl^-$ and $H_4TPyrP^{2+} \cdot 2Cl^-$ was found to be $\sim 3 - 4 \text{ \AA}$.^[3] The length of the hydrogen bond between protons and oxygen of perchlorate ion in complexes $H_4TPP^{2+} \cdot 2ClO_4^-$ and $H_4OEP^{2+} \cdot 2ClO_4^-$ is about 2 \AA .^[5]

The difference in the hydrogen bond length reflects the different value of the complexation constant due to dissimilarity of the properties of both porphyrin molecules and ligating ions. For the porphyrin under investigation, the complexation constant with halide ions decreases in the order $Cl^- > Br^- > I^-$ (see Table). Relatively low complexation constant for $H_4P^{2+} \cdot 2I^-$ complex may lead to the fluctuations of the hydrogen bond $H \dots I$ length in the solution as well the deviations from the symmetrical structure of the complex. The substantial broadening of the $H_4P^{2+} \cdot 2I^-$ complex absorption spectrum (Figure 4A and 7B) as compared with that of uncomplexed H_4P^{2+} molecule seems to be explained from that point of view. Thus, for the $H_4P^{2+} \cdot 2Br^-$ complex the broadening of the absorption spectrum is distinctly less (Figure 4B), and for the $H_4P^{2+} \cdot 2Cl^-$ complex it is vanishing (Figure 4C).

According to the X-ray data analysis for the diprotonated form H_2OEP with perchlorate ions ($H_4OEP^{2+} \cdot 2ClO_4^-$) the average value of dihedral angle θ between the planes of the pyrrolic rings and mean porphyrin plane is about 14° ,^[5] and the dihedral angle value depends on the interaction force between the ions in the complex.^[3,5,15] Therefore, the long wavelength shift in the electronic absorption spectrum upon the formation of $H_4P^{2+} \cdot 2I^-$ complex seems to be due to the increase in the tilting angle θ , *i.e.* results from an increase of the degree of the tetrapyrrolic macrocycle distortion upon complexation. This suggestion is in line with the results of quantum-chemical calculations, reported recently.^[12,15,16]

Steady State and Time-Resolved Fluorescence

As it was stated above, the formation of the $H_4P^{2+} \cdot 2I^-$ complexes upon increase of the KI concentration in the solution leads to the long wave-

length shift of Q -bands in the electronic absorption spectra and the changes of their spectral profiles. The absorption spectrum of H_4P^{2+} has two distinct Q -bands with maxima at 548 and 592 nm. The $H_4P^{2+} \cdot 2I^-$ complexes at 0.01 M concentration of KI show the diffusive spectrum with band maximum at 559 nm with a weak shoulder at 605 - 606 nm. Complex formation also leads to the substantial decrease in the intensity of the Soret band and small long wavelength shift of the maximum (1 nm at 0.01 M concentration of KI).

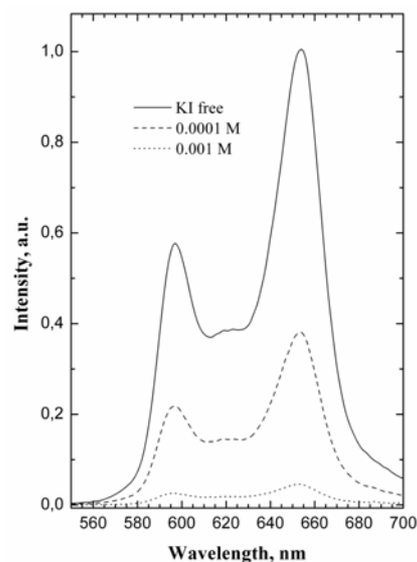


Figure 6. Fluorescence spectra of H_4P^{2+} as a function of KI concentration in acetonitrile; $\lambda_{exc} = 400 \text{ nm}$.

The fluorescence spectrum of the diprotonated H_4P^{2+} form has two sharp bands with maxima at 597 and 654 nm (Figure 6). The fluorescence quantum yield Φ_f of the diprotonated H_4P^{2+} form in the halide salts free solution is 0.11. The strong fluorescence intensity decrease is observed upon formation of the complexes $H_4P^{2+} \cdot 2I^-$ upon increase of KI concentration in the solution (Figure 6). At 0.01 M concentration of KI the integral fluorescence intensity is reduced for more than 300 times. Upon this fluorescence quenching the shape of the fluorescence spectrum remains the same.

The fluorescence excitation spectra have been measured in the Q -band region for the salts free sample and for the solutions with $3 \cdot 10^{-4}$ and $1 \cdot 10^{-3}$ M concentration of KI. All three spectra are identical (Figure 7A) and the maxima positions in the fluorescence excitation spectra were found at ~ 548 and $\sim 592 \text{ nm}$, as it was found for the absorption spectrum of uncomplexed H_4P^{2+} molecules (Figure 7B).

This finding indicates that fluorescence belongs to the uncomplexed diprotonated H_4P^{2+} form, and the fluorescence of complexes $H_4P^{2+} \cdot 2I^-$ is not observed. In other words, the complexed porphyrin $H_4P^{2+} \cdot 2I^-$ is nonfluorescent. If we neglect the contribution into the emission from the minor concentration of uncomplexed diprotonated form H_4P^{2+} in 0.01 M KI solution,^[31] the upper limit of the fluorescence quantum yield Φ_f for $H_4P^{2+} \cdot 2I^-$ complex can be estimated as low as $3.6 \cdot 10^{-4}$.

The fluorescence time decay ($\lambda_{exc} = 410 \text{ nm}$,

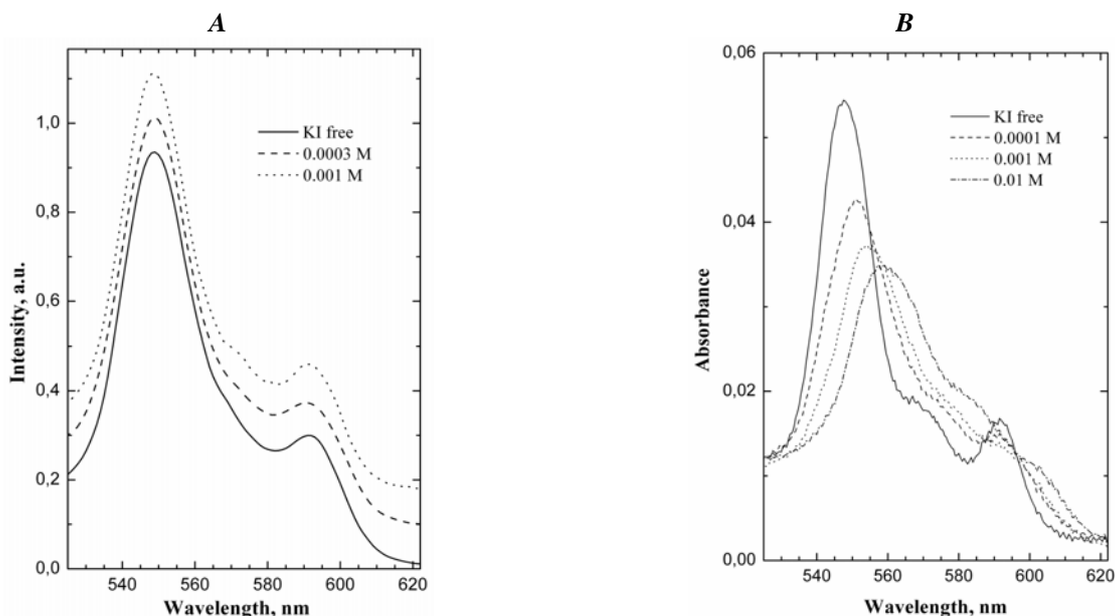


Figure 7. A - Excitation fluorescence spectra of H_4P^{2+} as a function of KI concentration in acetonitrile; $\lambda_{obs} = 654$ nm. The spectra are shifted up for 0.1 arbitrary units each for clarity sake; B - Electronic absorption spectra of H_4P^{2+} as a function of KI concentration in acetonitrile; porphyrin concentration is $1.9 \cdot 10^{-6}$ M.

$\lambda_{em} = 595$ nm) has monoexponential character for all the solutions studied. The fluorescence lifetime τ_{fl} progressively decreases in going from the salt free solution to solutions containing KI (Figure 8). For the initial solution of H_4P^{2+} (without KI) the fluorescence lifetime τ_{fl} is found to be 5.86 ± 0.05 ns, and τ_{fl} value is reduced down to 3.80 ± 0.05 ns at 0.002 M concentration of KI.

Static and Dynamic Fluorescence Quenching

As it was shown above, the fluorescence of diprotonated form is efficiently quenched by iodide ions. The Stern-Volmer plot (Figure 9) deviates from the

“classical” linear dependence. The concavity of Stern-Volmer plot with regard to the ordinate axis is an indication that both static and dynamic diffusion-controlled quenching take place simultaneously.^[46] Concaving up is because the quenching in such a case is the second order function of iodide ion concentration

$$I_0/I = 1 + (K_{SV}^c + K_{SV}^d)[I] + K_{SV}^c K_{SV}^d [I]^2, \quad (4)$$

where K_{SV}^c is the Stern-Volmer constant for the static fluorescence quenching, K_{SV}^d is the Stern-Volmer constant for the dynamic fluorescence quenching and $[I]$ is concentration iodide ion. The calculated Stern-Volmer constant K_{SV}^c of the static fluorescence quenching due to

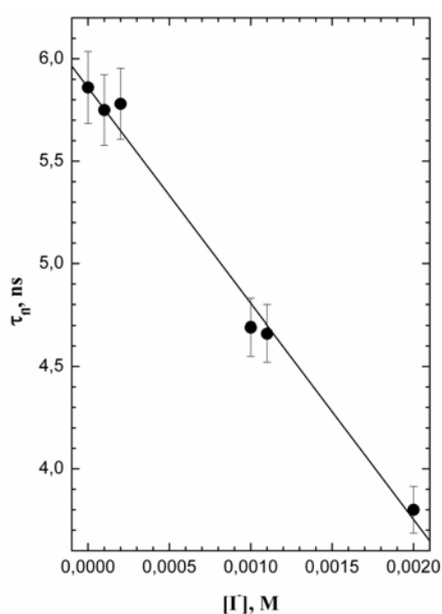


Figure 8. Fluorescence lifetime τ_{fl} as a function of KI concentration in acetonitrile; $\lambda_{exc} = 410$ nm, $\lambda_{obs} = 595$ nm.

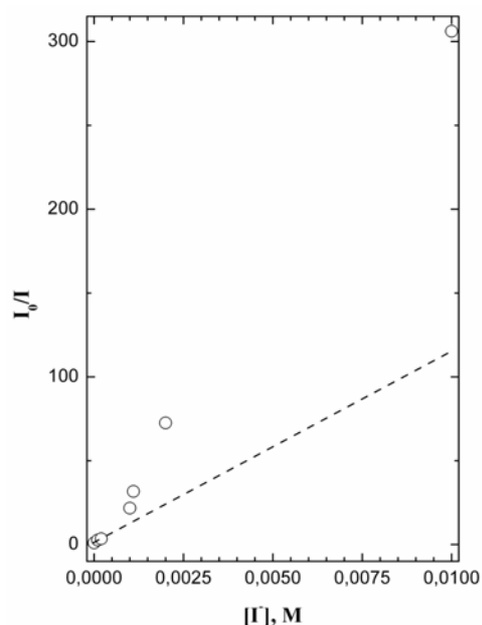


Figure 9. The Stern-Volmer plot for the fluorescence quenching of H_4P^{2+} with iodide ions in acetonitrile.

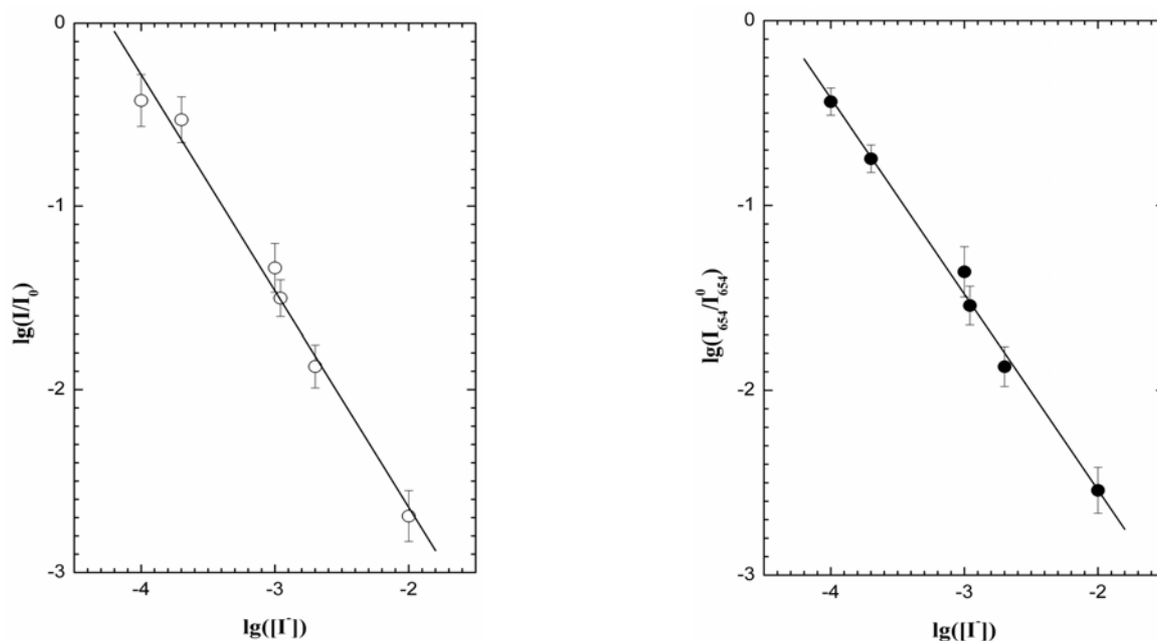


Figure 10. Double logarithmic plot for: A - the integral fluorescence intensity ratio I/I_0 , B - the peak intensity ratio I_{654}^0/I_{654} as a function of KI concentration.

complex formation with iodide-ions is 7860 M^{-1} . Fluorescence of diprotonated form of porphyrin in the complex with iodide-ions is quenched by mechanism of the internal heavy atom effect.^[4] The above data on the electronic absorption spectra and fluorescence intensity as the functions of the concentration of iodide ions in the solution indicate that the fluorescence quenching is practically parallel to the formation of the nonfluorescent $\text{H}_4\text{P}^{2+} \cdot 2\text{I}^-$ complexes, *i.e.* static quenching due to the complexation with halide ions is a dominating channel of the fluorescence quenching of the H_4P^{2+} molecule.

At the same time the shortening of the fluorescence lifetime of “halide ion free” diprotonated H_4P^{2+} form cannot be explained with the static quenching due to internal heavy atom effect. The dependence of the fluorescence lifetime on the concentration of iodide ions I^- in the solution is linear one (Figure 8). This fact indicates that the quenching can be described with Stern-Volmer approach.^[46] Therefore, one can state that the dynamic diffusion-controlled fluorescence quenching due to external heavy atom effect coexists with the static quenching due to complex formation. The Stern-Volmer constant K_{SV}^{d} of the dynamic fluorescence quenching by iodide-ions 277 M^{-1} has been calculated with an equation

$$\tau_{\text{fl}}^0/\tau_{\text{fl}} = 1 + K_{\text{SV}}^{\text{d}} [\text{I}^-], \quad (5)$$

where τ_{fl}^0 and τ_{fl} are the fluorescence lifetimes measured in the absence of quencher and with halide salts added in the solution, respectively. The calculated K_{SV}^{d} value for the dynamic quenching is almost 30 times smaller than that for the static quenching. This finding proves once more that the formation of the nonfluorescent complexes with halide ions is the dominating channel of the H_4P^{2+} porphyrin fluorescence quenching by halide ions.

Determination of the Iodide Ion Concentration

The concentration dependencies of both the integral fluorescence intensity and peak fluorescence intensity at $\lambda = 654 \text{ nm}$ are shown in Figure 10. The double logarithmic

plot is an analogue of the Scatchard plot, which is used widely in the fluorometric analysis of the ligand binding at the different hosts.^[47,48] The both plots demonstrate the linear character and can be proposed as the calibration for the iodide ion concentration measurements in the solution.^[32,49] In order to measure the iodide ions concentration the integral fluorescence intensity I_0 for the reference sample and intensity I for the studied sample with unknown iodide ion concentration should be measured first. Then the I_0/I ratio is calculated. The iodide ion concentration (in M) is available from the calibration formula (6)

$$[\text{I}^-] = 10^{(0.84 \lg(I_0/I) - 4.24)} \quad (6)$$

In a similar manner, the halide ion concentration can be determined with the ratio I_{654}^0/I_{654} of the peak fluorescence intensities at 654 nm for the studied and reference samples. With this approach the iodide ion concentration (in M) in the solution is given from the calibration formula (7)

$$[\text{I}^-] = 10^{(0.94 \lg(I_{654}^0/I_{654}) - 4.40)} \quad (7)$$

Moreover, the iodide ion concentration can also be determined with the measured concentration dependence of porphyrin fluorescence lifetime τ_{fl} (see Figure 8). The measurements of the fluorescence time decay can be achieved at any given wavelength in the range from 560 to 720 nm. The concentration of iodide ions (in M) is given as

$$[\text{I}^-] = 0.0055 - 0.00095\tau_{\text{fl}}, \quad (8)$$

where τ_{fl} is the fluorescence lifetime in nanoseconds.

Conclusions

The philosophy of measurements described above underlies the submitted Russian Federation patent pending.^[49] The proposed methods allow the determination of the iodide ions in the concentration range from

$3.0 \cdot 10^{-2} \text{ M}$ to $3.0 \cdot 10^{-5} \text{ M}$. As far as we know, the luminescent methods for the iodide ion concentration measurements described to date provide the sensitivity not higher than $1.5 \cdot 10^{-4} \text{ M}$.^[50] Therefore, the developed approach seems to be promising way to increase the sensitivity of the luminescent methods of the halide ion determination in the solutions.

The additional advantage of the proposed method is the possibility to carry out the measurements in the air-equilibrated solutions. The upper limit of the Stern-Volmer constant K_{SV} for the fluorescence quenching by molecular oxygen do not exceed 100 M^{-1} ($K_{SV} = \tau_0^n k_q$, where k_q is a bimolecular rate constant of the fluorescence quenching by molecular oxygen). This value is almost two orders of magnitude smaller than Stern-Volmer constant K_{SV}^d dominating in the fluorescence quenching by iodide ions. Therefore, the interfering influence of fluorescence quenching by molecular oxygen can be neglected.

The presented results demonstrate that diprotonated form of 3,7,13,17-tetramethyl-2,8,12,18-tetrabutylporphyrin can be used as a basic compound for the design of the luminescent molecular sensor for the measurements of the halogenide ion concentration in the solution. The diprotonated porphyrin binds the halide ions with equilibrium constant of order 10^3 - 10^5 M^{-1} . The fluorescence of porphyrin molecule was shown is quenched by halide ions due to both external and internal heavy atom effect. The methods for measurements of the iodide ion concentration using the fluorescence quenching of diprotonated porphyrin by halide ions have been proposed. These methods allow the ions to be detected in the air-equilibrated solutions at room temperatures in the wide concentration range.

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