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Mechanism for Copper(II)-Mediated Disaggregation of a Porphyrin J-Aggregate

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ABSTRACT: J-aggregates of anionic meso-tetakis(4-sulfonatophenyl)porphyrin form at intermediate pH (2.3–3.1) in the presence of NiSO₄ or ZnSO₄ (ionic strength, I.S. = 3.2 M). These aggregates convert to monomeric porphyrin units via metallation with copper(II) ions. The kinetics for the disassembly process, as monitored by UV/vis spectroscopy, exhibits zeroth-order behavior. The observed zeroth-order rate constants show a two-term dependence on copper(II) ion concentrations: linear and second order. Also observed is an inverse dependence on hydrogen ion concentration. Activation parameters have been determined for the disassembly process leading to AH° = (+163 ± 15) kJ·mol⁻¹ and AS° = (+136 ± 11) J·K⁻¹. A mechanism is proposed in which copper(II) cation is in pre-equilibrium with a reactive site at the rim of the J-aggregate. An intermediate copper species is thus formed that eventually leads to the final metallated porphyrin either through an assisted attack of a second metal ion or through a direct insertion of the metal cation into the macrocycle core.

INTRODUCTION

Kinetics and mechanisms for metal ion coordination and the removal from a number of porphyrins in aqueous and nonaqueous media have been extensively investigated. The reactivity in a variety of confined microenvironments, such as micelles and microemulsions, or even nucleic acids or proteins has been explored. On the contrary, the reactivity of porphyrin homoaggregates with metal ions has been the subject of very few studies.

Many water-soluble porphyrins have been shown to aggregate under appropriate conditions of concentration, pH, and ionic strength. The study of such species is facilitated by the impact of aggregation on their basic physical–chemical properties. For example, porphyrin electronic absorption is affected by aggregation leading to blue-shifted or red-shifted bands depending on whether the chromophores interact in a face-to-face (H-dimers) or in a side-by-side geometry (J-dimer), respectively. Extended J-aggregates of porphyrins are of particular interest because of their intriguing optical and electronic properties, such as line narrowing of the absorption bands, resonance light scattering (RLS), enhanced Raman scattering, nonlinear optical effects, and photoc conductivity. Water soluble anionic meso-tetakis(4-sulfonatophenyl)porphyrin (TPPS₄) in its diacid form is able to self-assemble through an extensive network of hydrogen bonding, electrostatic, and hydrophobic interactions. The kinetics of growth of these supramolecular systems depends on the concentration of the porphyrin and medium properties such as pH, ionic strength, the presence of some cationic species, and temperature. Their kinetics of formation has been described using models that begin with a nucleation step followed by autocatalytic growth of the assembly. Despite many reports dealing with the kinetics of assembly formation, relatively few studies have focused on the disassembly process leading to blue-shifted or red-shifted bands, resonance light scattering (RLS), enhanced Raman
Quite recently, we have shown that J-aggregates of TPPS₄ can be restored to the monomeric form through coordination of copper(II) ions into the porphyrin. The insertion of this metal ion into the porphyrin core leads to disruption of the nanoassemblies because it prevents any of the possible stabilizing interactions between adjacent porphyrins.

Reactions of metal ions with porphyrins have been investigated largely under pseudo-first-order conditions (i.e., a large excess of metal ions with respect to porphyrin), and the derived rate laws are in most cases first order both in porphyrin and in metal ion concentration. Furthermore, the rate law may contain a term or terms depending on the specific counter-anion and proton concentration. As previously shown, TPPS₄ J-aggregates react with copper(II) in a manner strongly dependent on the details of the mixing protocol of the reagents. The combined experimental evidence supports an attack by the metal ion at the external rims of the supramolecular assemblies. Because of the importance of disaggregation to various applications, we decided to extend our earlier study on the factors affecting the rate of this process. To permit a wide range of conditions as possible, we chose to work under relatively mild acid conditions (pH 2.3−3.1) and to promote aggregation through the addition of metal salts (ionic strength, I.S. = 3.2 M). To the best of our knowledge, this is the first detailed study on the mechanism of metal ion insertion into an already highly aggregated porphyrin system.

■ RESULTS AND DISCUSSION

The formation of J-aggregates from the porphyrin diacid (H₄TPPS₄) can be triggered by using low pH conditions or, at intermediate pH, by providing a high ionic strength environment. In a micromolar concentration range at pH values between 2.3 and 3.1, the TPPS₄ porphyrin is in its non-J-aggregated diacid form (pKₐ = 4.9). An increase in the ionic strength of the aqueous solution leads to the formation of J-aggregates arising from the screening of the electrostatic repulsions between interacting porphyrin units. In our experiments, Ni(II) or Zn(II) sulfate was used to provide the desired fixed ionic strength (I.S. = 3.2 M). These two d-block metal ions were chosen based on their similarity to copper(II) except for their limited reactivity under acidic conditions to form the corresponding metalloporphyrin complexes. Zinc(II) rapidly inserts into the porphyrin, but the corresponding metal complex is labile in our pH range, leading to rapid demetallation, or exchange with copper(II) ions. Ni(II) is significantly less reactive than either zinc(II) or copper(II). It was chosen for the experiments reported in the main text; results for the more labile zinc(II), which are used for comparison and follow similar trends, are collected in Supporting Information.

**Figure 1.** (A) UV/vis spectral changes for the formation of J-aggregates of the porphyrin TPPS₄, according to a porphyrin last mixing protocol, spectra have been collected with a scanning time of 10 s ([TPPS₄] = 3 μM, [H₂SO₄] = 0.05 M, I.S. = 3.2 M, T = 298 K). The inset shows the corresponding kinetic profile at 490 nm. (B) UV/vis spectral changes for the disassembly process of equilibrated TPPS₄ J-aggregates upon addition of CuSO₄, spectra have been collected with a scanning time of 10 s ([TPPS₄] = 3 μM, [H₂SO₄] = 0.05 M, 
[CuSO₄] = 0.3 M, I.S. = 3.2 M (NiSO₄), T = 298 K). The inset shows the corresponding kinetic profile at 490 nm. The notch detectable at 486 nm is an experimental artifact because of the deuterium emission spectra from the instrument lamp and an imperfect baseline correction.

The presence of NiSO₄ Upon mixing the reagents, the formation of the nanoassemblies can be monitored by the rapid appearance and increase in intensity of a band at 490 nm, that is, the spectroscopic signature of J-aggregated TPPS₄. The kinetic traces for this process follow a stretched exponential behavior that is characteristic for the "porphyrin last" mixing protocol described in previous investigations. When copper(II) sulfate is added to an equilibrated solution of J-aggregates, the J-band at 490 nm decreases linearly with time, that is, zero-order kinetics are observed (Figure 1B). When ZnSO₄ is used...
as a supporting electrolyte, a two-component B-band region grows in 412 and 405 nm because of the formation of the final reaction product, CuTPPS₄ (data not shown). These spectral features have been identified as arising from the monomer/dimer equilibrium of CuTPPS₄ previously studied with very fast relaxation methods. In the presence of NiSO₄ as a supporting electrolyte, as is the case of the reported spectra, the \(^{3}\text{Ag}_e \rightarrow \ ^{3}\text{T}_{1g} (P)\) transition around 400 nm due to \([\text{Ni}(\text{H}_2\text{O})_6]^{2+}\) masks these spectral changes.

To investigate the dependence of the rate for the disassembly process on experimental parameters, we prepared samples of J-aggregates (according to the above protocol) and induced their deaggregation by adding CuSO₄. From the slope of the zeroth-order kinetic traces, we calculated the values for the rate constants, \(k_0\) (Figure 2 and Figure S12). These data can be conveniently fitted by a second-order polynomial function of the form:

\[
k_K = (7.96 \pm 2.52) \times 10^{-11} \text{M}^{-1}\text{s}^{-1} + (1.29 \pm 0.15) \times 10^{-9} \text{M}^{-1}\text{s}^{-1} + (2.96 \pm 0.02) \times 10^{-9} \text{M}^{-1}\text{s}^{-1} \quad [\text{Cu}^{2+}]^2\]

at \(pH = 2.56, R^2 = 0.99939\). This behavior is different from that previously reported for this system for \([\text{Cu}^{2+}] < 0.3 \text{ M}\), suggesting different reaction pathways leading to the final reaction products in different regimes of metal ion concentration. At fixed \([\text{Cu}^{2+}]\), the rate constant shows a linear dependence on the inverse hydrogen ion concentration (Figure 3 and Figure S13), giving a slope of \((6.83 \pm 0.15) \times 10^{-12} \text{M}^2\text{s}^{-1}\). Such a dependence has been observed previously for the reaction with copper(II) of the water-soluble 2,4-disulfonated deuteroporphyrin dimethyl ester in acidic and neutral solutions. We note that the range of \(pH\) accessible in the experiments is limited by the acid stability of the J-aggregates at \(pH < 2.3\).

The rate law for the disassembly process is derived from the mechanism shown in Scheme 2 and is zero order because it is independent on the aggregate or the porphyrin concentration

\[
\text{Rate} = -d[\text{Agg}]/dt = k_0 \cdot k \cdot ([\text{Cu}^{2+}] / [H^+]) + k_2 \cdot K \cdot ([\text{Cu}^{2+}]^2 / [H^+]) = k_0
\]

where the meaning of \(k_0\), \(k_1\), \(k_2\), and \(K\) is discussed below.

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**Figure 2.** Plot of the observed zeroth-order rate constant \(k_0\) vs \([\text{Cu}^{2+}]\) at constant \(pH\) (\([\text{TPPS}_4] = 3 \mu\text{M}, [\text{H}_2\text{SO}_4] = 0.0016 \text{M}, \text{I.S.} = 3.2 \text{M (NiSO}_4)\), \(T = 298 \text{ K})

**Figure 3.** Plot of the observed zeroth-order rate constant \(k_0\) vs \(1/[H^+]\) (\([\text{TPPS}_4] = 3 \mu\text{M}, [\text{Cu}^{2+}] = 0.4 \text{ M, I.S.} = 3.2 \text{ M (NiSO}_4)\), \(T = 298 \text{ K})

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**Scheme 2.** Mechanism Proposed for the Disassembly Process of TPPS₄ J-Aggregates by Copper(II) Metal Ions

\[
\begin{align*}
\text{Agg} & \text{EH}_4 + \text{Cu}^{2+} \rightarrow \text{Agg} \text{EH}_4 \text{Cu}^{+} + \text{H}^+ \\
\text{Agg} \text{EH}_4 \text{Cu}^{+} + \text{Cu}^{2+} & \rightarrow \text{CuP} + 3\text{H}^+ + \text{Cu}^{2+} + \text{Agg} \\
\text{Agg} \text{EH}_4 \text{Cu}^{+} & \rightarrow \text{CuP} + 3\text{H}^+ + \text{Agg}
\end{align*}
\]

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**Scheme 3.** Formation of the Intermediate Species by Interaction of Cu²⁺ with Only One of the Nitrogen Atoms of the Protonated Core of a Reactive TPPS₄ Unit at the Rim of the J-Aggregates and Subsequent Insertion of the Metal Ion and Release of the Final CuTPPS₄ Metal Complex and Solvent Molecules

The reactivity of porphyrins within the extended assembly “interior porphyrins” is considered to be negligible because the porphyrin core is completely hindered by the porphyrin–porphyrin interactions stabilizing the network. Any involvement of these porphyrins would lead to a fragmentation of the aggregates, with a time-dependent formation of new reactive ends and acceleration of the rates. The linear kinetic traces and RLS experiments have ruled out such a possibility. The “porphyrin end” is the preferred site of attack for a copper(II) ion leading to the formation of an intermediate species (Agg \(\text{EH}_4 \text{Cu}^{+}\)), in which the metal interacts with the porphyrin core with a concomitant loss of a proton (equilibrium constant \(K\)). Because the porphyrin reactive end is part of the aggregate, only one of the two exposed nitrogen atoms of the porphyrin core is coordinating the copper ion, acting as a monodentate.
ligand (AggH4ECu+, Scheme 3). As further support of this point, recall that in reactions involving the metallation of monomeric porphyrins in aqueous solution, a complex dependence of rate on the proton concentration is commonly observed because of the two protonation equilibria involving the macrocycle core. In the present case, the observation of a simple inverse dependence suggests the loss of a single proton. In a successive step, the intermediate species, still embedded in the porphyrin nanoassembly, can undergo a metal ion insertion into the macrocycle core, with the formation of the final square planar copper(II) porphyrin complex, CuP. The release of proton ions in this mechanism is not expected to alter the pH of the bulk solution, because of the much larger concentration of H2SO4 (millimolar range) with respect to the porphyrin (micromolar range). Two distinct reaction pathways are proposed: (i) a metal assisted path via k2 and (ii) a direct insertion of copper(II) via k1. This latter type of mechanism is observed for many porphyrins,51 but the participation of a second metal ion (equal or different with respect to the inserted one) has been reported for meso-tetra(pyridyl)-porphyrins with di- and trivalent metal ions in aqueous solution.52 In both cases, the aggregate, containing n−1 units, is eventually released (Agg) and a new porphyrin reactive end will be available again until complete disruption of the nanoassembly (right side of Scheme 3). The time independence of the “porphyrin ends” concentration along the process (i.e., the concentration of reactive porphyrin remains small and is continuously replenished during the process) is in line with the observed zero-order kinetics, k0, and the overall rate law.

The very small term A in the fit of the rate constants in Figure 2 (A = (7.96 ± 2.52) × 10−11 M−1 s−1) can be ascribed to statistical errors or to a metal ion independent pathway, in which a very slow dissociation of monomers from the nanoassemblies is responsible for this small contribution. Indeed, under the same experimental conditions, the reactivity of the free diacid porphyrin toward copper(II) is at least an order of magnitude larger, with respect to the porphyrin embedded in the aggregates (t1/2 ≈ 80 s vs t1/2 ≈ 1500 s). Activation parameters for the disassembly process have been determined from an Eyring plot giving ΔH° = (+163 ± 15) kJ mol−1 and ΔS° = (+136 ± 11) J K−1 (Figure 4 and Table S14). The large positive value for the activation entropy may be due to the release of the metal derivative CuP from the nanoassembly to the solution and the loss of the water ligands coordinated to the copper center in the intermediate species, when the metal enters the porphyrin core (Scheme 3). For the sake of comparison, we determined the activation parameters for the copper(II) insertion into the monomeric diacid TPPS4, obtaining ΔH° = (+143 ± 6) kJ mol−1 and again a large positive ΔS° = (+194 ± 11) J K−1 (see Figure S15 and Table S15). These values are in line with literature data on the same porphyrin, even at different ionic strengths.53 When comparing the activation parameters for the copper(II) insertion in the free monomer and in the aggregate, the observed differences could be related to how the nanoassembly structure induces changes in the way the metal ion approaches the available face of the terminal porphyrin unit. Also, the sulfonate group of the porphyrin preceding the exposed terminal end could be responsible for a change in the interaction of the reactive end with the copper ion.

**CONCLUSIONS**

Given the large and burgeoning interest in J-aggregates for many potential applications, such as optoelectronic devices,18 optical waveguide,53 or nonlinear optics,22 a precise understanding of the factors governing their stability and controlling their sizes is very important. Moreover, a detailed knowledge of the kinetics of their disassembly allows controlling the release of porphyrins in a time-dependent way. This mechanism can be conveniently triggered by external stimuli, such as pH or metal ion concentration changes, making these aggregates as nanoreservoirs of porphyrin units, exploitale, for example, in photodynamic therapy or nanomedicine.54 Copper(II) has been chosen in the present study because of its lability, the relative stability of its metal complexes under acidic conditions and its limited redox behavior. However, other metal ions could be used to trigger the assembly/disassembly process of TPPS4 porphyrin, opening the way to a variety of potential stimuli controlled applications.

**EXPERIMENTAL SECTION**

The porphyrin 5,10,15,20-tetrakis(4-sulfonatophenyl)-porphyrin (TPPS4) as the tetrasodium salt was purchased from Frontier Scientific. CuSO4·SH2O, NiSO4, ZnSO4 and sulfuric acid (ACS grade) were obtained from Aldrich Co. and used as received. Milli-Q water was used for preparing all solutions. A concentrated stock solution of TPPS4 was freshly prepared, and the concentration of the stock solution was calculated using ε412 = 5.33 × 105 M−1 cm−1 at the B-band in neutral aqueous solution of low (∼0) ionic strength.

UV/vis extinction spectra were recorded on an Agilent 8453 diode array and on a Jasco V-550 spectrophotometer using 1 cm path length quartz cells. Because of the nonmolecular nature of the system under investigation, differences in the kinetic parameters are expected, when different sets of experiments are compared. The strong dependence of kinetics under initial conditions or even on the mixing protocol has already pointed out in our previous papers. In the present work, to obtain a better reproducibility of the data, when investigating a particular effect (i.e., copper(II) concentration or pH), a complete series of kinetic runs have been performed using the same batch of preformed J-aggregate. To this purpose, a large volume of solution of TPPS4 J-aggregate (10 mL) was prepared using a porphyrin last protocol by adding a known volume of concentrated TPPS4 solution (ca. 200 μM) to a solution containing NiSO4 (or ZnSO4) and H2SO4 at the required concentration, and the

![Figure 4](image-url)
solution was equilibrated for at least an hour. Kinetic runs were initiated in a plastic cell placed in the thermostated compartment of the instruments by mixing equal volumes of a solution containing CuSO₄, NiSO₄ (or ZnSO₄), and H₂SO₄ at the required concentration and the solution containing the preformed J-aggregate (in all the experiments I.S. = 3.2 M by addition of NiSO₄ or ZnSO₄).

The zero-order kinetics was analyzed by a linear fit of the extinction data collected at 490 nm after normalization to unity. The concentration of aggregated monomers at any time has been obtained by subtracting the amount of diacid or metallated porphyrin (determined using literature values of the extinction coefficients) from the initial TPP₅ concentration, obtaining zero-order kinetic constants (M⁻¹s⁻¹) as slopes.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsomega.8b02913.

Dependencies of rate constants on [TPPS₄], [Cu²⁺], and pH (I.S. with ZnSO₄), Eyring plots for the disaggregation in the presence of ZnSO₄ and for metallation of TPP₅ with copper(II), and primary kinetic data (PDF).

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Author Contributions
The manuscript was written through contributions of all authors. All authors have given approval to the final version of the manuscript.

Notes
The authors declare no competing financial interest.

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