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Self-complementary double-stranded porphyrin arrays assembled from an alternating pyridyl-porphyrin sequence†

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Oligomeric porphyrin arrays with an alternating pyridyl-porphyrin sequence were synthesized to explore double-strand formation through self-complementary pyridyl-to-zinc axial coordination bonds. Competitive titration experiments revealed the thermodynamic aspects involved in the zipper effect within double-strand formation. Multiple axial coordination bonds defined the stacked conformation, despite a marginal contribution to the stability of the double-strands. Thus, the zipper cooperativity was the dominant factor for the remarkable stability. Moreover, the dimeric and trimeric porphyrin arrays were independently assembled into double-strands by self-sorting from a binary mixture. Double-strand formation engineered discretely stacked π -systems. Successive slipped-cofacial stacks of the porphyrin rings progressively extended the π -system via exciton coupling over the double-strand while keeping a relatively high fluorescence quantum yield.

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Introduction

Naturally occurring double-stranded polymers, such as DNA and proteins, display an exquisite molecular organization in biological systems. Double-stranded DNA is currently emerging as a passive scaffold and provides state-of-the-art bottom-up nanotechnologies with a molecular scale precision, as exemplified by DNA-origami.1 This shape-programmable nanotechnology is enabled by both the sequence-specific double-strand formation and shape-persistent double-stranded building units. Therefore, the development of an intelligent doublestrand as a building material for sophisticated hierarchical architectures in which functionalities could be integrated should be of significant interest. The present study examines novel double-strand-forming oligomeric porphyrin arrays which may, in a straightforward manner, be used to incorporate photoelectronic functions into structures for artificial photosynthesis.

The high fidelity of Watson–Crick base pairing *via* complementary hydrogen bonds plays a crucial role in the formation of double-stranded DNA.² In a similar way, artificial double-strands require the molecular design of complementary pairs

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through either multiple hydrogen bonds³ or charge-transfer interactions.⁴ While the majority of research in the field has been devoted to the formation of helical structures,⁵ almost no synthetic effort has been aimed at the applications of these structures in nanomaterial science, though they represent a promising new paradigm in bottom-up molecular assembly. The molecular design of supramolecular building materials could include not only sequence selectivity but also the shape-persistence of the functional double-stranded structures.

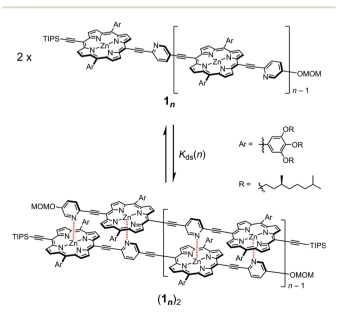
For decades, porphyrin frameworks have been part of technological advancements at the forefront of material studies because of their large π -systems and outstanding molar extinction coefficients. Their structural relevance in natural photosynthetic systems has been the source of considerable interest and has driven research in supramolecular multiporphyrin architectures.^{6,7} The rigid porphyrin plane offers an ideal platform for versatile multiporphyrin architectures which can be built using both covalent and supramolecular approaches.8,9 Biomimetic supramolecular porphyrin architectures with slipped-cofacially stacked conformations have been a useful motif for the design of materials with excellent photoelectronic functionalities, as described by Kobuke and coworkers. 10,11 A coordination-directed approach is particularly effective in the assembly of ladder complexes that are composed of fully π -conjugated multiporphyrin arrays via the use of bidentate ligands, as demonstrated by Anderson and collaborators.12 A double-strand is an intriguing motif to use to create novel artificial photosynthetic systems and engineer discrete stacked π -systems.

In our previous report, the formation of self-coordinated zinc(2-pyridylethynyl)porphyrin dimers was controlled by the choice of nucleation conditions, where the formation of the initial coordination bond governed how the second coordination bond would form.¹³ In a non-coordinating solvent, the formation of the second intra-dimer coordination bond was more energetically favorable than the initial binding was, which led to a self-complementary pattern of multiple coordination bonds. According to the same principle, we envisioned that oligomeric zinc(2-pyridylethynyl)porphyrin arrays $\mathbf{1}_n$ with an alternating pyridyl-porphyrin sequence could be assembled into a double-strand $(1_n)_2$ (Scheme 1). The formation of the initial interstrand coordination bond would induce the spontaneous formation of the second and the subsequent coordination bonds in a self-complementary fashion, because of the increasingly favorable thermodynamics of binding. This is the so-called zipper effect. Here, we present the molecular organization of novel double-stranded porphyrin arrays based on self-complementary ligand-metalloporphyrin sequence, which provides successively stacked porphyrin arrays. The photophysical properties of these systems are studied.

Results and discussion

Synthesis

The synthetic route to the monomeric zinc(2-pyridylethynyl) porphyrin has already been established by our group. ¹³ According to the reported procedure, we prepared the monomeric $\mathbf{1}_1$ as the precursor for the oligomeric $\mathbf{1}_n$. A systematic series of new oligomers $\mathbf{1}_n$ (n=2-3) were then prepared from $\mathbf{1}_1$ via repetitive Sonogashira–Hagihara coupling reactions (Scheme 2). ¹⁴ The details of the synthetic procedures are described in the ESI, † together with the essential thermodynamic and photophysical properties of $(\mathbf{1}_1)_2$ and $(\mathbf{1}_2)_2$. The following sections mainly demonstrate the results



Scheme 1 Formation of the double-strands $(1_n)_2$ (n = 1-3).

Scheme 2 Synthesis of the porphyrin arrays $\mathbf{1}_n$ (n = 2-3).

representative of $(1_3)_2$, which is an assembly composed of six porphyrin rings and six pyridyl groups.

Structural elucidation of double-strands

double-strand (13)2 was observed using MALDI-TOF MS measurements (Fig. 1A). $(1_3)_2$ spontaneously formed in the lone stationary state in toluene, as identified by NMR. Only three sets of protons in the pyridyl and porphyrin-β positions showed unambiguous diagonal correlations via the nuclear Overhauser effect (NOE) (Fig. 1B), which is indicative of two trimeric porphyrin arrays that are cofacially assembled in an antiparallel arrangement (Fig. 1A). NOE correlation with the protons of the MOM group determined one of three porphyrin rings. Subsequently, TOCSY correlations identified two sets of signals corresponding to the 2-, 3- and 6-pyridyl and 2"-, 3"- and 6"-pyridyl protons. An alternative assignment for these two sets was also possible for these pyridyl-porphyrin pairs. However, these pyridyl protons individually showed NOE correlations with the porphyrin rings, which indicated the pairing of the porphyrin rings with the complementary pyridyl groups. This finding is consistent with a symmetrically assembled structure, as indicated by a lack of multiplied signals for the porphyrin array. All of the aromatic resonances of the pyridyl groups were found in the non-aromatic region (6.34-2.56 ppm), which suggested that the axially coordinated pyridyl groups were strongly shielded in the vicinity of the porphyrin ring. The assignment is consistent with all of the observed NMR resonances.

In pyridine- d_5 , which can act as a competitive coordinating ligand, the non-shielded resonances of the pyridyl protons of the disassembled species were observed in the aromatic region with the disappearance of the upfield-signals observed for the double-strand. The comparison of the spectra in the two solvents suggests the assembly of double-strand $(\mathbf{1}_3)_2$ through pyridyl-to-zinc coordination bonds. All of the NMR data firmly established that the structural picture of the discrete double-strand $(\mathbf{1}_3)_2$ assembled from an alternating pyridyl-porphyrin sequence via self-complementary coordination bonds.

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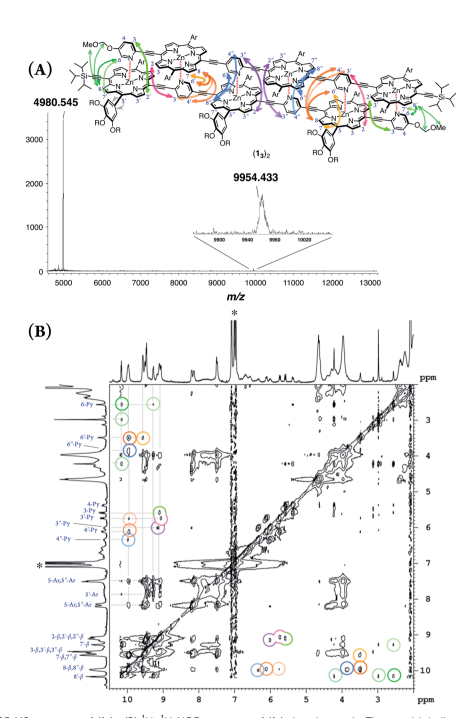


Fig. 1 (A) MALDI-TOF MS spectrum of $(\mathbf{1}_3)_2$. (B) $^1\text{H}-^1\text{H}$ NOE spectrum of $(\mathbf{1}_3)_2$ in toluene- d_8 . The asterisk indicates residual toluene. One alternative assignment is shown.

Thermodynamic behaviors

Double-strand $(\mathbf{1}_3)_2$ was sufficiently durable to obey Beer's law over a wide concentration range $(\mathbf{10}^{-7}$ to $\mathbf{10}^{-4}$ M) in toluene. In contrast, the spectral shape of the electronic absorption spectra of $\mathbf{1}_1$ depended on the concentration, which suggested a small association constant for the formation of $(\mathbf{1}_1)_2$ ($K_{ds}(\mathbf{1}) = 1.3 \pm 0.2 \times 10^4$ M $^{-1}$). The association constant of $(\mathbf{1}_3)_2$, $K_{ds}(3)$, was found to be too high to directly evaluate the thermodynamic stability of double-strand $(\mathbf{1}_3)_2$. Over the course of competitive

titration experiments with pyridine, the spectral changes showed several pseudo-isosbestic points, which suggested that the equilibria involved essentially two stationary states, *i.e.*, the double-strand and the unzipped single-strand (Fig. 2). Competitive titration experiments allowed us to analyze the thermodynamic stability of the double-strands according to the thermodynamic cycle (Scheme 3). We analyzed the unzipping equilibria by employing a tentative one-step unzipping model, which is useful for the description of the simplified overall equilibria. Nonlinear least-square fittings gave reliable binding

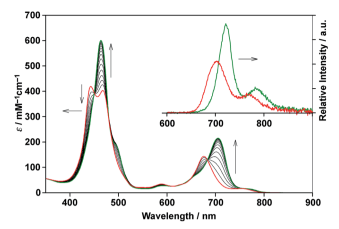


Fig. 2 Spectrometric titration of $(\mathbf{1_3})_2$ $([\mathbf{1_3}]_0 = 2.9 \times 10^{-6}$ M) with pyridine (up to 480 equiv., red to green) at 25 °C in toluene. The inset shows the fluorescence spectra of $(\mathbf{1_3})_2$ and $\mathbf{1_3}$ in the presence of excess pyridine $(10^4$ equiv.) in toluene $(\lambda_{ex} = 450$ nm, a pseudo-isosbestic point).

properties for the overall unzipping equilibria.¹⁴ The overall unzipping constant ($K_{\rm uz}$) is then described as follows:

$$K_{\rm uz}(n) = \frac{[1_n L_n]^2}{[(1_n)_2][L]^{2n}}$$
 (1)

Assuming that the microscopic binding constant (K_{μ}) is identical for each ligand-to-zinc axial coordination bond of the single-strand, the microscopic binding constant can be approximated by eqn (2).

$$K_{\mu} = \frac{\left[1_{n} L_{n}\right]^{1/n}}{\left[1_{n}\right]^{1/n} [L]} = \frac{\left[2_{\text{Si}} L\right]}{\left[2_{\text{Si}}\right] [L]}$$
(2)

In practice, the titration of model zinc porphyrin $2_{\rm Si}$ with pyridine to yield the axially coordinated zinc porphyrin gave the experimental K_{μ} values ($K_{\mu} = (3.2 \pm 0.1) \times 10^4 \ {\rm M}^{-1}$) (Scheme 3B). The values, in turn, gave the binding constant for the double-strand formation ($K_{\rm ds}(n)$) according to eqn (3).

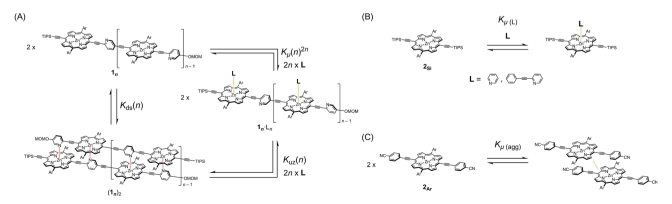
$$K_{\rm ds}(n) = \frac{\left[(1_n)_2\right]}{\left[1_n\right]^2} = \frac{K_{\mu}^{2n}}{K_{\rm uz}(n)}$$
 (3)

These estimated thermodynamic parameters are useful to the discussion on the durability of double-strand $(\mathbf{1}_n)_2$ (Table 1). The values of $K_{\rm ds}(n)$, $K_{\rm ds}(2)=(2.5\pm0.3)\times10^9~{\rm M}^{-1}$ and $K_{\rm ds}(3)=(6.5\pm1.2)\times10^{11}~{\rm M}^{-1}$, were remarkable, considering the small magnitude of the microscopic binding constant as described below.

The zipper effect was quantified by the synergetic free energy change ($\Delta\Delta G(n)$), the excess energy beyond the sum of the independent free energy changes induced by pyridyl-to-zinc axial coordinating and π -stacked microscopic binding.¹⁵

$$\Delta \Delta G(n) = \Delta G_{\rm ds}(n) - 2n\Delta G_{\mu} \tag{4}$$

The $\Delta\Delta G(n)$ values substantially dominated the stability of the double-strands, with multiple axial coordination bonds defining the discrete conformation of the double-strand despite their minimal thermodynamic contribution. To quantify a reliable microscopic binding constant, we employed model compounds. The binding constant for the axial coordination of 2-(phenylethynyl)pyridine to 2_{si} as the model for each ligand-tozinc axial coordination bond is $K_{\mu(L)} = 7.8 \pm 1.1 \text{ M}^{-1}$ (Scheme 3B) and that for the self-aggregation of 2_{Ar} as the model for π stacked interactions is $K_{\mu(\text{agg})} = 11 \pm 1 \text{ M}^{-1}$ (Scheme 3C). The microscopic binding constant for single coordination together with π -stacked interactions was, then, determined to be 9.4 \pm 1 M^{-1} ($K_{\mu}=K_{\mu(L)}^{1/2}K_{\mu(agg)}^{1/2}$, $\Delta G_{\mu}^{\circ}=-5.5\,\pm\,0.3\,$ kJ $\,\mathrm{mol}^{-1}$). The dramatic change from $\Delta\Delta G^{\circ}(1)$ to $\Delta\Delta G^{\circ}(2)$ suggested that the zipper effect, as a consequence of chelate cooperativity including interactions such as π -stacking, van der Waals interactions, and desolvation entropy, became greater with the addition of the repeating units. In contrast, the $\Delta\Delta G^{\circ}(3)$ value was similar to that of $\Delta\Delta G^{\circ}(2)$, which indicated that the compensatory effects gave rise to structural strain, such as the distortion of the porphyrin planes, and a loss of structural entropy. The analyses elucidated that the increasing number of repeat units increased the durability of $(\mathbf{1}_n)_2$, although the cooperativity per interaction decreased. The nucleation step of



Scheme 3 (A) Generic closed thermodynamic cycle (n = 2-3). (B) Microscopic binding equilibrium of model porphyrin $\mathbf{2}_{Si}$ with axial ligands; L = pyridine and 2-(phenylethynyl)pyridine. (C) Aggregation equilibrium of model porphyrin $\mathbf{2}_{Ar}$.

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Table 1 Thermodynamic parameters of 1_n at 25 °C in toluene

	n	$K_{\rm uz}(n)^a/{ m M}^{1-2n}$	$K_{\mathrm{ds}}(n)/\mathrm{M}^{-1}\left(\Delta G_{\mathrm{ds}}(n)/\mathrm{kJ}\ \mathrm{mol}^{-1}\right)$	$\Delta\Delta G^{\circ}(n)^d/\text{kJ mol}^{-1}\left(\text{EM}^e/\text{M}\right)$
1 ₁ 1 ₂	1 2	$-$ (4.1 \pm 0.2) \times 10 ⁸	$(1.3 \pm 0.2) \times 10^4 (-23 \pm 1)^b (2.5 \pm 0.3) \times 10^9 (-53 \pm 1)^c$	$-12 \pm 0.4 (147 \pm 9) -31 \pm 0.2 (68 \pm 8)$
1 ₃	3	$(6.5\pm 1.2)\times 10^{15}$	$(1.7\pm0.3)\times10^{11}(-64\pm1)^c$	$-31\pm0.2~(12\pm1)$

^a Estimated from the competitive titration experiments with pyridine. ^b Directly estimated from variations in concentration. ^c Estimated using eqn (3), wherein $K_{\mu} = (3.2 \pm 0.1) \times 10^4 \text{ M}^{-1}$. ^d Estimated by employing $\Delta G_{\mu}^{\circ} = -5.5 \pm 0.3 \text{ kJ mol}^{-1}$, wherein $K_{\mu} = K_{\mu(L)}^{1/2} K_{\mu(agg)}^{1/2} = 9.4 \pm 1 \text{ M}^{-1}$. Effective molarity (EM) was evaluated from eqn (6).

multiple coordination bonds is dominant when $\mathbf{1}_n$ self-assembles, and synergetic effects govern the durability of $(\mathbf{1}_n)_2$ for $n \leq$ 3. The self-complementary coordination bonds were significantly stabilized by the zipper effect, thereby thermodynamically funnelling the self-assembled structures into the most stable form without kinetic entrapment by metastable states. This allowed for the realization of the specific double-strand formation.

The situations are alternatively examined in terms of "effective molarity (EM)", an empirical parameter used to describe the effectiveness of intramolecular interactions, to index the upper limit of concentration for the double-strand formation. The EM value is defined in eqn (5) and (6).16,17

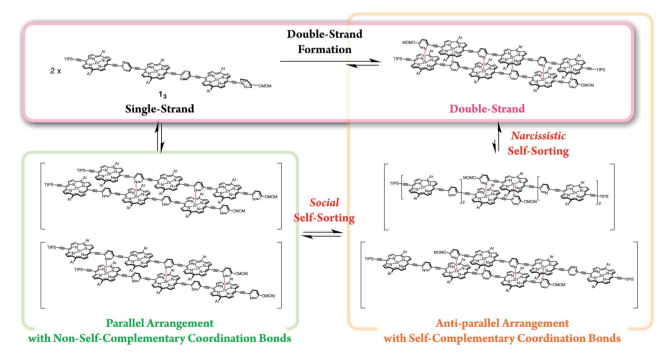
$$EM = (K_{ds}(n)/K_{\mu}^{2n})^{1/(2n-1)}$$
 (5)

$$= \exp\{-\Delta \Delta G(n)/(2n-1)RT\} \tag{6}$$

The EM values of $(\mathbf{1}_1)_2$ and $(\mathbf{1}_2)_2$ (Table 1) were in the high end range of the typical values for supramolecular systems,17 inferring that the model systems were, in the most precise sense, imperfect for the formation of discrete double-strands. However, the EM values predicted a significant selectivity of the double-strand formation even at very concentrated conditions.

Self-sorting behaviors

The exclusive formation of the self-complementary doublestrand $(\mathbf{1}_n)_2$ intrinsically involves self-sorting behaviors due to the possibility of several binding patterns for the self-association of $\mathbf{1}_n$ (Scheme 4). Briefly, social self-sorting favors selfcomplementary binding patterns, whereas narcissistic self-sorting distinguishes differences in the numbers of the repeating units of the self-complementary binding patterns. 12b,18 Based on the lack of hysteresis, the self-sorting assembly of the doublestrands occurred during the heating/cooling processes (25-70 °C) of a binary mixture of $(\mathbf{1}_2)_2$ and $(\mathbf{1}_3)_2$. The electronic absorption spectra appeared as a superimposition of the spectra of $(\mathbf{1}_2)_2$ and $(\mathbf{1}_3)_2$ at 25 °C before and after heating, although thermally dissociated 12 and 13 mutually interacted to a degree at 70 °C. The mutually orthogonal self-assembly from a thermally unzipped mixture perfectly restored the initial



Scheme 4 Double-strand formation of 13 self-sorted from possible self-assembled patterns.

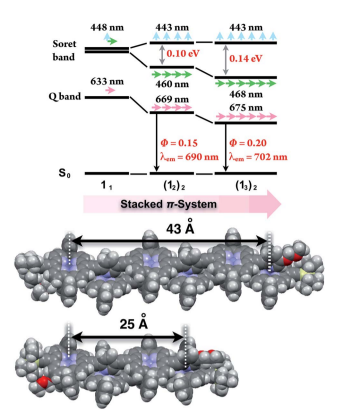


Fig. 3 Energy diagrams of the double-strands $(\mathbf{1}_n)_2$, and their geometry-optimized structures calculated using the MM + force field (HyperChem Ver. 8.0 software). The alkoxy side chains are omitted for visual clarity. The wavelengths denote the absorption maxima.

Table 2 Photophysical properties of double-strands $(\mathbf{1}_n)_2$, $\mathbf{1}_n$ accommodated with pyridine $(\mathbf{1}_n \cdot \mathrm{Py}_n)$ and 2 in toluene

	$\lambda_{\mathrm{em}}{}^{a}/\mathrm{nm}\left(\Phi\right)$	$\tau/\mathrm{ns}\left(\alpha\right)$	$k_{\rm em}^{e}/{\rm s}^{-1}$
(12)2(13)22Si·Py	690 (0.15) 702 (0.20) 635 (0.07) 650 (0.08)	$0.65 (0.51), 1.15 (0.49)^b$ $0.56 (0.32), 1.03 (0.68)^b$ 1.61^c 1.45^d	1.7×10^{8} 2.3×10^{8} 4.3×10^{7} 5.6×10^{7}

^a Emission maximum ($\lambda_{\rm em}$) and fluorescence quantum yield (Φ) obtained using an integration sphere (excitation at 452 nm for ($\mathbf{1}_2$)₂, 450 nm for ($\mathbf{1}_3$)₂, and 405 nm for $\mathbf{2}_{\rm Si}$ and $\mathbf{2}_{\rm Si}$ ·Py). ^b Fluorescence lifetime (τ) and the normalized amplitude (α) determined from the fluorescence decay profiles in the range of 623–773 nm upon excitation at 483 nm (Fig. 4). ^c Emission at 635 nm upon excitation at 405 nm. ^d Emission at 650 nm upon excitation at 405 nm. ^e Radiative rate constant defined as $k_{\rm em} = \Phi/\tau$. The single-strand $\mathbf{2}_{\rm Si}$ ·Py was observed in the presence of excess pyridine ($\mathbf{10}^4$ equiv.).

fluorescence properties, as indicated by the superimposition of $(\mathbf{1}_2)_2$ and $(\mathbf{1}_3)_2$ at 25 °C. The self-sorting capacity of double-strands $(\mathbf{1}_n)_2$ makes them promising in the construction of units that are capable of simultaneous multiple molecular assembly within bottom-up nanotechnology applications.

Engineering discrete stacked π -systems

In the last stage, our attention turned to the electronic properties of the double-strands $(\mathbf{1}_2)_2$ and $(\mathbf{1}_3)_2$. The double-strand

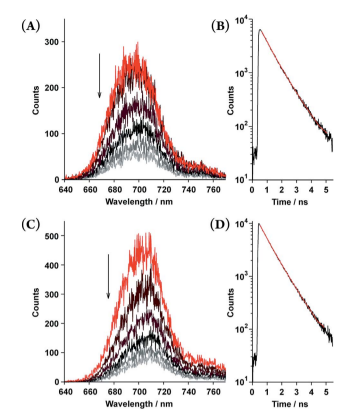


Fig. 4 Time-resolved fluorescence spectra recorded every 0.3 ns (upper to lower; 0.45–0.55, 0.75–0.85, 1.05–1.15, 1.35–1.45, 1.65–1.75, and 1.95–2.05 ns) and fluorescence decay profiles of the double-strands (1₂)₂ at [1₂] = 5.5×10^{-6} M (A and B) and (1₃)₂ at [1₃] = 6.9×10^{-6} M (C and D) in toluene. The fluorescence decay profiles (black lines) are shown with fitted curves based on the biexponential decay constants (red lines, τ shown in Table 2) in the range of 623–773 nm.

formation of $\mathbf{1}_n$ resulted in a successively slipped-cofacial stack of porphyrin arrays. The double-strands $(\mathbf{1}_n)_2$ showed a splitting of the Soret band with a bathochromic shift of the lower band due to exciton coupling in the electronic absorption spectra. 10,19 The split width of the Soret band of (13)2 (0.14 eV) was wider than that of $(\mathbf{1}_2)_2$ (0.10 eV), where the lowest exciton band was red-shifted and the highest exciton band remained unchanged. At the same time, the Q band of $(1_3)_2$ displayed a larger bathochromic shift to 675 nm (1.83 eV) than that of $(\mathbf{1}_2)_2$ at 669 nm (1.85 eV), and the longest Zn···Zn distances were estimated to be 43 Å for $(\mathbf{1}_3)_2$ and 25 Å for $(\mathbf{1}_2)_2$ based on the geometry-optimized structures (Fig. 3). The comparison of the electronic structures of $(\mathbf{1}_3)_2$ and $(\mathbf{1}_2)_2$ unveiled that the doublestranded multiporphyrin arrays exhibited exciton coupling due to successive slipped-cofacial stacks, leading to long-range π -electronic interactions.

The emission properties are noteworthy because the double-strands $(\mathbf{1}_n)_2$ did not dissipate a photoexcited singlet. The absolute fluorescence quantum yields (Φ) of $(\mathbf{1}_2)_2$ and of $(\mathbf{1}_3)_2$ were 0.15 and 0.20, respectively (Table 2). Increasing the number of porphyrin units raised the Φ values by extending the π -systems. The relatively high fluorescence efficiency of $(\mathbf{1}_n)_2$ suggested that the double-stranded structure was effective in

circumventing nonradiative deactivation pathways, even in the near-infrared wavelength region. The successive slipped-cofacial stacks of the porphyrin planes in the double-strand serve as structural and functional mimics of bacterial light-harvesting antenna complexes which display an efficient capture of sunlight and photoexcited energy transfer.⁷

Time-resolved fluorescence spectroscopy gave further insight into the photophysical dynamics of the excited states (Fig. 4). The biexponential fluorescence decay profiles of both double-strands indicated the existence of dual fluorescent states (Table 2). Over time, double-strand $(\mathbf{1}_3)_2$ showed a small red-shift in its emission wavelength. In contrast, the spectral shift of $(\mathbf{1}_2)_2$ was smaller than that of $(\mathbf{1}_3)_2$. It is intriguing to consider that the photophysical properties are relevant to the thermodynamic aspect of the double-strands; $(\mathbf{1}_3)_2$ was much more strained than $(\mathbf{1}_2)_2$. Structural relaxation could provide an energy sink that would be capable of trapping photoexcited singlets. For instance, it is known that porphyrin planes are ruffled in the photoexcited singlet state. 20,21 This interpretation is a likely reason as to why the slow fluorescence decay manifests as a red shift.

Conclusions

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In summary, we have synthesized double-stranded porphyrin arrays that yielded successive slipped-cofacial stacks with strong exciton coupling. Titration experiments revealed the thermodynamic aspects underlying the specific double-strand formation, and are used to describe a process in which selfcomplementary coordination bonds define the discrete structure. The zipper effect dominated the stability of the doublestranded structure. The remarkable selectivity for double-strand formation may serve as a powerful building tool for sophisticated bottom-up molecular assemblies with molecular scale precision, similar to those found in DNA nanotechnology. Moreover, the double-strands extended the π -electron network without any dissipation of the fluorescence properties due to the assembly of successive slipped-cofacial stacks. The shapepersistent double-stranded porphyrin arrays provide new options for artificial photosynthetic systems based on shapeprogrammable, and bottom-up molecular architectures.

Acknowledgements

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