Efficient Artificial Light-Harvesting System Based on Supramolecular Peptide Nanotubes in Water

Qiao Song, Sofia Goia, Jie Yang, Stephen C. L. Hall, Michael Staniforth, Vasilios G. Stavros, and Sébastien Perrier*

ABSTRACT: Artificial light-harvesting systems in aqueous media which mimic nature are of significant importance; however, they are often restrained by the solubility and the undesired aggregation-caused quenching effect of the hydrophobic chromophores. Here, we report a generalized strategy toward the construction of efficient artificial light-harvesting systems based on supramolecular peptide nanotubes in water. By molecularly aligning the hydrophobic chromophores along the nanotubes in a slipped manner, an artificial light-harvesting system with a two-step sequential Förster resonance energy transfer process is successfully fabricated, showing an energy transfer efficiency up to 95% and a remarkably high fluorescence quantum yield of 30%, along with high stability. Furthermore, the spectral emission could be continuously tuned from blue through green to orange, as well as outputted as a white light continuum with a fluorescence quantum yield of 29.9%. Our findings provide a versatile approach of designing efficient artificial light-harvesting systems and constructing highly emissive organic materials in aqueous media.

INTRODUCTION

In nature, plants and photosynthetic bacteria are able to capture, transfer, and store solar energy utilizing their highly efficient light-harvesting systems. Within these light-harvesting systems, chromophores are aligned into arrays by the surrounding proteins, which are capable of absorbing and capturing, transferring, and storing solar energy through their highly efficient light-harvesting systems. In nature, plants and photosynthetic bacteria are able to capture, transfer, and store solar energy utilizing their highly efficient light-harvesting systems. Within these light-harvesting systems, chromophores are aligned into arrays by the surrounding proteins, which are capable of absorbing and capturing, transferring, and storing solar energy utilizing their highly efficient light-harvesting systems. Within these light-harvesting systems, chromophores are aligned into arrays by the surrounding proteins, which are capable of absorbing and capturing, transferring, and storing solar energy through the Förster resonance energy transfer (FRET) processes from donor chromophores to acceptor chromophores, including self-assembled structures such as dendrimers, supramolecular polymers, gels, micelles, vesicles, and bioinspired structures. However, limited by the solubility and the undesired aggregation-caused quenching (ACQ) effect of the hydrophobic chromophores in water, light-harvesting systems built in aqueous environments often show unsatisfying performance with considerably low fluorescence quantum yields. Within this limitation, while the utilization of fluorophores with aggregation induced emission (AIE) properties has been shown to be a powerful tool for constructing light-harvesting systems with relatively high quantum yields, a generalized strategy is essential to tackle this issue. Moreover, artificial light-harvesting systems with high stability are of significant importance for practical applications, which is rarely emphasized in supramolecular systems.

Assemblies of cyclic peptide–polymer conjugates are a relatively new type of supramolecular polymers. The alternating D- and L-amino acid configuration of the cyclic octapeptides enables strong multiple hydrogen bonding interactions between the flat rings of the cyclic peptides, driving the formation of nanotubular assemblies. Conjugating hydrophilic polymers onto cyclic peptides prevents their lateral aggregation and improves their solubility and stability in aqueous environments, forming self-assembled polymeric nanotubes with well-defined structure. Inspired by the highly ordered structures guided by cyclic peptides, we envisioned that by attaching the hydrophobic chromophores onto the cyclic peptide–polymer conjugates, they could be molecularly aligned along the supramolecular polymeric nanotubes in a slipped way, enabling us to design efficient artificial light-harvesting systems while avoiding the undesired ACQ effect of the chromophores in water. Moreover, due to the strong binding affinity between the cyclic peptides, the artificial light-harvesting system is expected to exhibit very good stability.

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In this work, we report an efficient artificial light-harvesting system with a two-step sequential FRET process constructed by molecularly aligning the hydrophobic chromophores along the self-assembled peptide nanotubes in water. As shown in Scheme 1, three fluorophore-cyclic peptide-polymer conjugates, pyrene-cyclic peptide-poly(ethylene glycol) (PYR-CP-PEG), naphthalene monomide-cyclic peptide-poly(ethylene glycol) (NTI-CP-PEG), and cyanine3-cyclic peptide-poly(ethylene glycol) (Cy3-CP-PEG), are designed and synthesized. An efficient FRET process takes place from PYR-CP-PEG to NTI-CP-PEG when coassembling both into supramolecular peptide nanotubes in water, which meets the requirement of promising candidates for light-harvesting systems. A two-step sequential FRET process could be further realized by incorporating a third conjugate, Cy3-CP-PEG. In this regard, light energy is transferred from PYR-CP-PEG to NTI-CP-PEG, and subsequently transferred to Cy3-CP-PEG. More importantly, due to the slipped stacking arrangement of the fluorophores along the supramolecular peptide nanotubes, the typical ACQ effect is largely inhibited with fluorescence quantum yields measured to be over 30%. Moreover, the emission color could be continuously tuned from blue to green and finally to orange simply by adjusting the ratio of the three conjugates. Particularly, a pure white-light emission of (0.32, 0.34) in CIE 1931 coordinates is obtained at the ratio of the three conjugates. Particularly, a pure white-light emission of (0.32, 0.34) in CIE 1931 coordinates is obtained at the ratio of the three conjugates. Furthermore, similar structures were also observed by SANS and TEM for the assembled polymeric nanotube structures. Furthermore, similar structures were also observed by SANS and TEM for the assembled polymeric nanotube structures. Subtle structural differences were observed between nanotubes formed by each conjugate, hypothesized to be induced by the physicochemical properties of different fluorophores (SI Table S1). These conjugates could also be visualized by TEM (Figures 1B,C and S11), showing 1D structures for the three conjugates in aqueous media. The diameter of the nanotubes was narrowly distributed around 8 nm, while the length varied within 100 nm, which is consistent with the proposed self-assembled polymeric nanotube structures. Furthermore, similar structures were also observed by SANS and TEM for the mixtures of these three conjugates (Figures 1D and S12).

**RESULTS AND DISCUSSION**

**Design, Synthesis, and Self-Assembly of Fluorophore-Cyclic Peptide-Polymer Conjugates.** The three fluorophore-cyclic peptide-polymer conjugates, PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG, were synthesized by attaching the corresponding fluorophores via either NHS coupling chemistry or HATU coupling chemistry, followed by conjugating the hydrophilic polymer PEG ($M_n = 5000$ g mol$^{-1}$) via strained alkyne/azide group ligation. PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG were subsequently characterized by ESI-MS, HPLC, and GPC (Figures S1–S7 of the Supporting Information, SI). Two fluorophore-polymer conjugates in the absence of cyclic peptides, PYR-PEG and NTI-PEG, were also synthesized as control compounds (Figures S8–S10).

Multiple hydrogen bonding interactions between the cyclic peptides are the driving forces of forming self-assembled polymeric nanotubes. The self-assembling behavior of conjugates PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG in aqueous solution was investigated by small angle neutron scattering (SANS) and transmission electron microscopy (TEM). Figure 1A shows the reduced, corrected scattering data for PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG conjugates in water. Using SASfit software, the data could be fitted with a cylindrical micelle model, suggesting the self-assemblies forming polymer nanotubes in aqueous solutions. Subtle structural differences were observed between nanotubes formed by each conjugate, hypothesized to be induced by the physicochemical properties of different fluorophores (SI Table S1). These conjugates could also be visualized by TEM (Figures 1B,C and S11), showing 1D structures for the three conjugates in aqueous media. The diameter of the nanotubes was narrowly distributed around 8 nm, while the length varied within 100 nm, which is consistent with the proposed self-assembled polymeric nanotube structures. Furthermore, similar structures were also observed by SANS and TEM for the mixtures of these three conjugates (Figures 1D and S12).

**Proof and Evaluation of the Artificial Light-Harvesting System with a Two-Step Sequential FRET Process.** Two factors are required for an efficient FRET system: (1) the emission spectrum of the donor fluorophore must overlay well with the absorption spectrum of the acceptor fluorophore; and (2) the donor and acceptor should be close to each other (within $\sim 10$ nm). We initially investigated the FRET...
process between PYR-CP-PEG and NTI-CP-PEG by measuring the absorption and emission spectra of PYR-CP-PEG and NTI-CP-PEG as well as the two control compounds PYR-PEG and NTI-PEG. Interestingly, the emission spectrum of PYR-CP-PEG showed minimal overlap with the absorption spectrum of NTI-PEG (Figure S13), in contrast to the emission spectrum of PYR-CP-PEG which overlapped well with the absorption spectrum of NTI-CP-PEG (Figure 2A). This could be ascribed to the formation of pyrene excimers caused by the stacking of the cyclic peptides, as indicated by the peak in the emission spectrum at 460 nm. As a result, an efficient energy transfer process is expected to occur when coassembling PYR-CP-PEG and NTI-CP-PEG conjugates. Indeed, as demonstrated in Figure 2B, with the increase of NTI-CP-PEG/PYR-CP-PEG ratio, PYR excimer emission intensity at 460 nm decreased gradually, and the emission intensity of the NTI fluorophore at 520 nm increased when excited at 335 nm. Fluorescence decay experiments were performed to further confirm the energy transfer process. As shown in Figure 2C, the fluorescence decay of PYR-CP-PEG was significantly faster in the presence of NTI-CP-PEG. The decay curves were fitted using a three-exponential decay model (Figure S14, Table S2), showing the fluorescence lifetimes of \( \tau_1 = 3.4 \text{ ns} \), \( \tau_2 = 20.9 \text{ ns} \), and \( \tau_3 = 67.2 \text{ ns} \) for PYR-CP-PEG. The fluorescence lifetimes decreased to \( \tau_1 = 2.7 \text{ ns} \), \( \tau_2 = 16.7 \text{ ns} \), and \( \tau_3 = 59.4 \text{ ns} \) when mixing with 2% NTI-CP-PEG, and further decreased to \( \tau_1 = 1.5 \text{ ns} \), \( \tau_2 = 9.8 \text{ ns} \), and \( \tau_3 = 40.9 \text{ ns} \) by coassembling 8% NTI-CP-PEG. This clearly indicates that the energy can be transferred from PYR-CP-PEG to NTI-CP-PEG when coassembling both into supramolecular polymeric nanotubes. To prove the importance of forming self-assembled polymeric nanotubes, control experiments were conducted. As shown in Figure S15, no FRET was observed when mixing either PYR-PEG/NTI-CP-PEG or PYR-CP-PEG/NTI-PEG in aqueous solutions, suggesting that the closely packed structure formed by coassembling both fluorophores into polymeric nanotubes is essential for FRET.

To quantitatively evaluate its performance as an artificial light-harvesting system, energy transfer efficiency (\( \Phi_{ET} \)) and antenna effect (AE) were measured and calculated at different PYR-CP-PEG/NTI-CP-PEG ratios (Figure 2D, Table S3). \( \Phi_{ET} \) represents the fraction of absorbed energy by the donor that is transferred to the acceptor, while AE describes how much brighter the acceptor emits by exciting the donor instead of directly exciting the acceptor. Overall, the \( \Phi_{ET} \) values increased with the increase of NTI-CP-PEG ratio and eventually reached up to 90%. The AE values, however, decreased with the increase of NTI-CP-PEG ratio. Particularly, we found that the AE value could be maintained above 10 while the \( \Phi_{ET} \) reached 90%. For example, at a PYR-CP-PEG/NTI-CP-PEG ratio of 100/4, \( \Phi_{ET} \) is calculated to be 72.6% with an AE value of 15.8, at a PYR-CP-PEG/NTI-CP-PEG ratio of 100/8, \( \Phi_{ET} \) is 88.1% with an AE value of 10.4. More importantly, PYR-CP-PEG exhibited a fluorescence quantum yield as high as 39.7 ± 3.7%, while the quantum yield of PYR-CP-PEG/NTI-CP-PEG = 100/8 was measured to be 34.1 ± 1.5%. To the best of our knowledge, this represents one of the highest values reported for aqueous artificial light-harvesting systems,

An ideal artificial light-harvesting system should be robust in various environmental conditions. Due to the high fidelity of the multiple hydrogen bonding interactions between the cyclic peptides, this supramolecular peptide nanotube-based light-harvesting system exhibits impressive stability at a wide range of concentrations and temperatures. Herein, AE and \( \Phi_{ET} \) were used to evaluate the performance of the light-harvesting system (PYR-CP-PEG/NTI-CP-PEG = 100/8) while varying the concentration or temperature. When decreasing the concentration or temperature, the AE value of 15.8; at a PYR-CP-PEG/NTI-CP-PEG ratio of 100/4, \( \Phi_{ET} \) is 72.6% with an AE value of 15.8, at a PYR-CP-PEG/NTI-CP-PEG ratio of 100/8, \( \Phi_{ET} \) is 88.1% with an AE value of 10.4. More importantly, PYR-CP-PEG exhibited a fluorescence quantum yield as high as 39.7 ± 3.7%, while the quantum yield of PYR-CP-PEG/NTI-CP-PEG = 100/8 was measured to be 34.1 ± 1.5%. To the best of our knowledge, this represents one of the highest values reported for aqueous artificial light-harvesting systems,

Figure 2. FRET process between PYR-CP-PEG and NTI-CP-PEG. (A) Normalized fluorescence spectra (dashed curves) of PYR-CP-PEG (blue trace, \( \lambda_{ex} = 335 \text{ nm} \)) and NTI-CP-PEG (green trace, \( \lambda_{ex} = 460 \text{ nm} \)), and their normalized absorbance spectra (solid curves). (B) Fluorescence spectra of PYR-CP-PEG in water with different concentrations of NTI-CP-PEG (\( \lambda_{ex} = 335 \text{ nm} \)). (C) Fluorescence decay profiles of PYR-CP-PEG and PYR-CP-PEG/NTI-CP-PEG. (D) AE and \( \Phi_{ET} \) at different NTI-CP-PEG/PYR-CP-PEG ratios.
AE and $\Phi_{ET}$ were both decreasing slowly with the increase of temperature. Nevertheless, even at 80 °C, only less than 10% decrease of AE and $\Phi_{ET}$ (4.7% and 7.1%, respectively) was observed. Hence, the stability of this supramolecular peptide nanotube-based system demonstrates its great potential for practical applications as an artificial light-harvesting system.

Nature uses multistep sequential energy transfer rather than only one-step energy transfer to make better utilization of light in a wide wavelength range. We explored the possibility of fabricating multistep sequential energy transfer system based on self-assembled polymeric nanotubes. Cy3-CP-PEG conjugate was rationally chosen as the second acceptor to harvest light emitted from NTI-CP-PEG, as the absorption spectrum of Cy3-CP-PEG overlaps well with the emission spectrum of NTI-CP-PEG (Figure 3A). In this respect, it is anticipated that

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\text{Fluorescence spectra of } \text{PYR-CP-PEG/NTI-CP-PEG} = 100/8 \text{ in presence of Cy3-CP-PEG, as the absorption spectrum of Cy3-CP-PEG overlaps well with the emission spectrum of NTI-CP-PEG.}
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Figure 3. Two-step sequential FRET process between PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG. (A) Normalized fluorescence spectra (dashed curves) of NTI-CP-PEG (green trace, $\lambda_{ex}$ = 460 nm) and Cy3-CP-PEG (orange trace, $\lambda_{ex}$ = 476 nm), and their normalized absorbance spectra (solid curves). (B) Fluorescence spectra of NTI-CP-PEG in water with the addition of different molar ratios of Cy3-CP-PEG ($\lambda_{ex}$ = 460 nm). (C) Fluorescence decay profiles of NTI-CP-PEG and NTI-CP-PEG/Cy3-CP-PEG. (D) Fluorescence spectra of PYR-CP-PEG/NTI-CP-PEG = 100/8 in water with the addition of different molar ratios of Cy3-CP-PEG ($\lambda_{ex}$ = 335 nm).

NTI-CP-PEG could act as a bridge to harvest light emitted by PYR-CP-PEG, and subsequently transfer the energy to Cy3-CP-PEG. First, the FRET efficiency between NTI-CP-PEG and Cy3-CP-PEG was studied. As evidently illustrated by Figure 3B, with the increase of Cy3-CP-PEG concentration, NTI emission intensity at 550 nm decreased gradually, and the emission intensity of Cy3 at 570 nm increased significantly when excited at 460 nm. Fluorescence decay experiments were also conducted to confirm the FRET process between NTI-CP-PEG and Cy3-CP-PEG, which shows a faster fluorescence decay of NTI-CP-PEG in the presence of Cy3-CP-PEG (Figures 3C and S18, and Table S4). Second, to facilitate the proposed sequential energy transfer process, a three-component system was constructed by coassembling PYR-CP-PEG, NTI-CP-PEG and Cy3-CP-PEG. As shown in Figure 3D, when Cy3-CP-PEG was added into PYR-CP-PEG/NTI-CP-PEG, the emission band ascribed to NTI decreased and a new emission band at 570 nm belonging to Cy3-CP-PEG appeared simultaneously when excited at 335 nm. These data indicate the occurrence of a two-step sequential FRET process, which is further supported by fluorescence decay experiments (Figure S19, Table S5). It should be noted that the energy might be transferred directly from PYR-CP-PEG to Cy3-CP-PEG; however, we anticipate this to be minor owing to the poor overlap between the emission spectrum of PYR-CP-PEG and the absorption spectrum of Cy3-CP-PEG (Figure S20). Thus, in the three-component system, a weak one-step energy transfer from PYR-CP-PEG to Cy3-CP-PEG is possible, though, when excited at 335 nm, a two-step sequential FRET process is anticipated to be preferred. Similar to the two-component system composed of PYR-CP-PEG and NTI-CP-PEG, increasing the relative concentration of the third component, Cy3-CP-PEG, caused a significant increase in $\Phi_{ET}$, reaching 95.0% at a low donor/acceptor ratio of 100/8/8 (Figure S21, Table S6). The fluorescence quantum yield was determined to be 29.9 ± 0.7% at a ratio of 100/8/2, further emphasizing the high energy transfer efficiency between the three components. The above results confirm that the supramolecular polymeric nanotubes fabricated by PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG could function as a promising artificial light-harvesting system with high energy transfer efficiency and high fluorescence quantum yield.

To further understand the dynamical processes involved in the energy transfer between the fluorophores, transient electronic absorption spectroscopy (TEAS) measurements were performed on PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG conjugates. Transient absorption spectra (TAS) were collated following photoexcitation at 335 nm of the aqueous solutions containing conjugates of different ratios; the TAS are presented as heat maps in Figure 4A–C. Briefly, three features could be observed: an initial ground state bleach centered at 350 nm, a narrow excited state absorption centered at 375 nm, and a broader excited state absorption centered at 490 nm. These features started decaying at pump–probe time delays ($\Delta t$) > 1 ps, with their decay persisting to the maximum experimental $\Delta t$ = 3 ns of the instrumental setup (see Figure S22 for different time delays chosen to highlight these features). To quantitatively analyze the dynamical processes observed, a global sequential fit (A → B → C→ ···) requiring up to five time constants was employed to extract the kinetics in the TAS data. The first two time constants for PYR-CP-PEG and PYR-CP-PEG/NTI-CP-PEG, and the first time constant for PYR-CP-PEG/NTI-CP-PEG/Cy3-CP-PEG were required to account for the instrument response (these likely include contributions from the solvent and the glass of the cuvette, thus the need in some circumstances for multiple components in the fit). The remaining three time constants (for each system) were very similar, implicating three transient species in the overall kinetic scheme with the following (approximate) time constants: 3 ps, 150 ps, and >3 ns. The fitted data is presented as evolution-associated difference spectra (EADS) in Figure 4D–F. The addition of NTI-CP-PEG as the first FRET acceptor was evident in the TAS as the intensity of the broad excited state absorption centered at 490 nm faded and narrowed down (Figure 4B). Moreover, as evidenced in Figure 4E, the small decrease in intensity centered at 550 nm was also indicative of the energy transfer from PYR-CP-PEG to NTI-CP-PEG, which coincides with the stimulated emission observed in the TAS of NTI-CP-PEG on its own (Figure S23a,e); the absence of a stimulated emission in Figure 4E could be due to masking from an excited state absorption. The addition of Cy3-CP-PEG as the second acceptor was...
evidenced in Figure 4C,F through the formation of an intense stimulated emission feature centered at 560 nm (Figure S23b,f), confirming the energy being further transferred to Cy3-CP-PEG as the second acceptor is only negligibly excited when analyzed on its own at 335 nm (Figure S24b,e). This stimulated emission feature appears to grow in with increasing delay times, which could be indicative of a two-step sequential energy transfer process. In the meantime, the presence of the stimulated emission even at the earliest times might be explained by a direct energy transfer from PYR-CP-PEG to Cy3-CP-PEG.

Emission Color Tuning Utilizing the Three-Component System. The highly efficient FRET between PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG enables us to straightforwardly tune the emission color. To determine the color range that could be obtained from this three-component system, two solutions with relatively high acceptor/donor ratios (NTI-CP-PEG/PYR-CP-PEG = 1/1, Cy3-CP-PEG/NTI-CP-PEG/PYR-CP-PEG = 1/1/1) were prepared, guaranteeing almost 100% FRET efficiency. Figure 5A shows the emission spectra of PYR-CP-PEG, NTI-CP-PEG/PYR-CP-PEG = 1/1, and Cy3-CP-PEG/NTI-CP-PEG/PYR-CP-PEG = 1/1/1 excited at 335 nm, which are then related to specific color coordinates in the CIE diagram. As expected, PYR-CP-PEG showed blue color emission (0.154, 0.180), and the emission color turned into green (0.353, 0.593) with the addition of one equivalent NTI-CP-PEG. The emission color could be further adjusted to orange (0.570, 0.418) when adding another equivalent of Cy3-CP-PEG (Figure 5B). By adjusting different PYR-CP-PEG/NTI-CP-PEG/Cy3-CP-PEG ratios, self-assembled polymeric nanotubes with continuously tunable emission color within the triangular color gamut could be easily obtained (Figures S25, S28). The high binding fidelity of the cyclic peptides endows the supramolecular polymeric nanotubes with unusually high stability. As shown in Figure S27, there was no significant variation of emission color when changing either concentration or temperature. Moreover, the emission color remained unchanged when integrating these supramolecular polymeric nanotubes into acrylamide hydrogels (Figure S28).

Notably, a white-light emission of (0.321, 0.336) in CIE coordinates could be obtained with the ratio of PYR-CP-PEG/NTI-CP-PEG/Cy3-CP-PEG = 100/1/2 (Figure S29). Under this condition, ΦET from PYR-CP-PEG to NTI-CP-PEG was calculated to be 37.0%, while ΦET from NTI-CP-PEG to Cy3-CP-PEG was 67.9% (Figure S29). The fluorescence quantum yield was measured to be 29.9 ± 3.1%, close to the reported record value for white-light-emitting organic materials in water (38%). Moreover, the white-emitting color could be maintained at a wide range of solution concentrations (Figure S30). Considering the high stability and high fluorescence quantum yield of the supramolecular polymeric nanotubes, this three-component system is believed to be suitable for applications as white-light-emitting materials.
CONCLUSIONS

We have reported an artificial light-harvesting system in aqueous environment based on self-assembled peptide nanotubes constructed by cyclic peptide–polymer conjugates. By simply coassembling the three conjugates, PYR-CP-PEG, NTI-CP-PEG, and Cy3-CP-PEG, an efficient artificial light-harvesting system with two-step sequential FRET process was achieved. Through effective alignment of the donor and acceptor fluorophores thanks to the supramolecular peptide nanotubes, the ACQ effect which typically plagues these systems was significantly reduced, resulting in high fluorescence quantum yields (≥30%) in aqueous media. The emission color could be continuously tuned from blue to green and finally to orange by adjusting the ratio of the three conjugates. Remarkably, pure white light with a fluorescence quantum yield as high as 29.9% is also accessible. Moreover, this system shows good stability, which could preserve its performance at low concentration (<1 μM) and high temperature (80 °C). This supramolecular peptide nanotube-based system is expected to serve as an efficient artificial light-harvesting system with potential light-emitting applications. Moreover, to better mimic the natural photosynthesis process, acceptor fluorophores with photocatalytic activities could be used to conduct photochemical reactions, thus making full use of the harvested energy.

ASSOCIATED CONTENT

Supporting Information
The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/jacs.0c11060.

Materials and instruments, details about synthetic procedures and characterization, and supplemental figures (PDF)

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REFERENCES


(9) Schultz, M.; Kunz, V.; Frieschmann, P. D.; Würthen, F. A supramolecular ruthenium macrocycle with high catalytic activity for water oxidation that mechanistically mimics photosystem II. *Nat. Chem.* 2016, 8 (6), 576–583.


(49) Broussard, J. A.; Rappaz, B.; Webb, D. J.; Brown, C. M. Fluorescence resonance energy transfer microscopy as demonstrated by measuring the activation of the serine/threonine kinase Akt. Nat. Protoc. 2013, 8 (2), 265–281.


