

Construction of Segregated Arrays of Multiple Donor and Acceptor Units Using a Dendritic Scaffold: Remarkable Dendrimer Effects on Photoinduced Charge Separation

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Abstract: Dendritic molecules appended with multiple zinc porphyrin units (DP_m, m [number of zinc porphyrin units] = 6, 12, and 24) trap bipyridine compounds carrying multiple fullerene units (Py₂F_n, n [number of C₆₀ units] = 1−3), affording coordination complexes DP_m⊃Py₂F_n having a photoactive layer consisting of spatially segregated donor and acceptor arrays on their surface. Complexes DP_m⊃Py₂F_n are stable enough (K [average binding affinity] = 1.1 × 10⁶−4.4 × 10⁶ M⁻¹ in CHCl₃ at 25 °C) to be isolated by gel permeation chromatography. UHV−STM microscopy enables clear visualization of a petal-like structure of DP₁₂⊃Py₂F₃. Photoexcitation of the zinc porphyrin units in DP_m⊃Py₂F_n results in a zinc porphyrin-to-fullerene electron transfer to generate a charge separation. The charge-separation rate constant (K_{CS}) in CH₂Cl₂ at 20 °C increases from 0.26 × 10¹⁰ to 2.3 × 10¹⁰ s⁻¹ upon increment of E_{CS} 0 m and E_{CS} 1. Consequently, DP₂₄⊃Py₂F₃ furnishes the largest ratio of E_{CS} 1 months and E_{CS} 2 months and E_{CS} 3 furnishes the largest ratio of E_{CS} 1 months and E_{CS} 3 months and E_{CS} 4 months are constant (E_{CS} 6 months and E_{CS} 6 months are constant (E_{CS} 1 months and E_{CS} 2 months are constant (E_{CS} 3 months and E_{CS} 4 months are constant (E_{CS} 3 months and E_{CS} 4 months are constant (E_{CS} 4 months are constant (E_{CS} 4 months and E_{CS} 6 months are constant (E_{CS} 6 months and E_{CS} 6 months are constant (E_{CS} 1 months and E_{CS} 2 months are constant (E_{CS} 3 months are constant (E_{CS} 4 months are constant (E_{CS} 6 months are constant ($E_$

Introduction

In biological photosynthesis, photoinduced electron transfer (PET) is one of the most essential events for conversion of solar energy into chemical energy. Molecular design of artificial photosynthetic systems involving restricted spatial arrangements of covalently¹ and noncovalently² linked electron donor (D) and acceptor (A) units has long been a central interest and now attracts even greater attention for the development of optoelectronic devices such as solar cells.³ For the fabrication of those materials, charge-transfer complexations between D and A units, leading to rapid quenching of charge-carrier transports, must be avoided, while D and A units are required to assemble individually to form their segregated arrays.⁴ Having this context

in mind, we were motivated to achieve at the molecular level such segregated arrays of multiple D and A units. Although a variety of D/A systems having multiple D or A units have been reported,⁵ those containing large numbers of both D and A units are unprecedented. In the present paper, we report novel photofunctional dendrimers consisting of spatially segregated arrays of multiple D and A units on their surface.

Our molecular design strategy (Scheme 1) made use of petallike dendritic structures as scaffolds⁶ to realize a wheel-like or spherical arrangement of multiple zinc porphyrin (P) units (DP_m; m = 6, 12, and 24),⁷ which are capable of ligating multiple molecules of bipyridine compounds having 1–3 fullerene (F) units (Py₂F_n; n = 1-3). Since the zinc porphyrin moieties in DP_m are located in the outermost dendritic layer, and the

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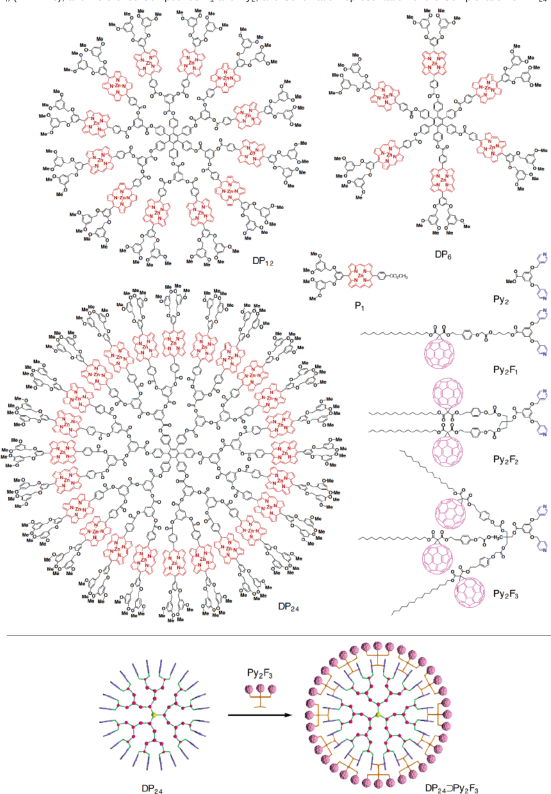
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Scheme 1. Molecular Structures of Zinc Complexes of Multiporphyrin Dendrimers DP_m (m=6, 12, and 24), Fullerene-Appended Bipyridine Ligands Py_2F_n (n=1-3), and Reference Compounds P_1 and Py_2 , and Schematic Representation of the Complexation of DP_{24} with Py_2F_3



bipyridine and fullerene units in Py_2F_n are apart by roughly 2-3 nm from one another, the coordination of Py_2F_n to DP_m is expected to form a photoactive layer consisting of spatially segregated arrays of multiple donor (zinc porphyrin) and acceptor (fullerene) units on the dendrimer surface. Molecular design of DP_m was inspired by the unique structures of light-harvesting antenna complexes such as LH1 and LH2 in purple

bacteria,⁸ where multiple bacteriochlorophyll units are spatially arranged like a wheel and ensure efficient harvesting of dilute photons. The excitation energy captured by one of the bacteriochlorophyll unit migrates, without dissipation, to neighboring chlorophyll units in the same and connecting wheel-like chromophore arrays and is subsequently funneled to the photosynthetic reaction center, thereby triggering an electron

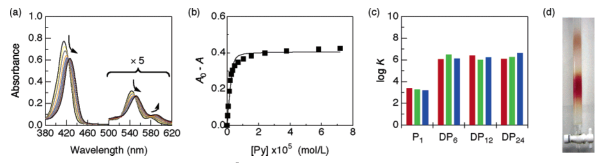


Figure 1. (a) Absorption spectral change of DP_{24} (1.5 × 10^{-7} M) upon titration with Py_2F_3 ($[Py_2F_3]/[DP_{24}] = 0$, 1.1, 2.3, 4.6, 6.8, 9.1, 11, 16, 23, 34, 52, 80, 126, 194, and 240) in CHCl₃ at 25 °C. (b) Change in absorbance ($A_0 - A$) of DP_{24} , monitored at 415.0 nm, as a function of [Py] (=2 × $[Py_2F_3]$), and its fitting profile. (c) Binding affinities ($\log K$) of the pyridine units in Py_2F_1 (red), Py_2F_2 (green), and Py_2F_3 (blue) toward the zinc porphyrin units in P_1 , DP_6 , DP_{12} , and DP_{24} in CHCl₃ at 25 °C. (d) A snapshot of gel permeation chromatography (GPC) of a mixture of DP_{24} and Py_2F_3 ($[Py_2F_3]/[DP_{24}] = 25$) with CHCl₃ as an eluent.

transfer. We and other groups have reported that some dendritic multiporphyrin arrays including DP_m , upon photoexcitation, display highly efficient energy migration characteristics analogous to those of the biological light-harvesting systems. ^{6a,9} Hence, we decided to make use of zinc complexes of multiporphyrin dendrimers DP_m with a certain structural rigidity for an attempt to construct a concentric double layer of spatially segregated arrays composed of multiple D and A units (Scheme 1). As an electron acceptor for DP_m , we chose a fullerene such as C₆₀ because of its small reorganization energy and excellent electron-accepting properties.¹⁰ Thus, we synthesized Py₂F_n having 1-3 fullerene units (n = 1-3), where the bipyridine (Py₂) unit was expected to coordinate strongly in a bidentate fashion to two neighboring zinc porphyrin units in DP_m , thereby allowing the formation of a fullerene array outside of the zinc porphyrin array on the dendrimer surface $(DP_m \supset Py_2F_n)$. Here, one can change the packing densities of these D and A units in the photoactive layer by varying the m and n values in the DP_m and Py_2F_n components, respectively, and might therefore be able to modulate the photoinduced charge-separation event.

Results and Discussion

Zinc complexes of multiporphyrin dendrimers DP_m (m = 6, 12, and 24) were synthesized according to a method analogous to that reported previously for chiroptical sensing of chiral bipyridine compounds.⁷ On the other hand, fullerene-appended bipyridine ligands Py_2F_n (n = 1-3) were synthesized by coupling of a fullerene-containing carboxylic acid precursor with

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bipyridine-terminated alcohols bearing 1–3 hydroxyl groups. ¹¹ In CHCl₃, DP₆ showed an absorption spectral profile typical of 5,15-diarylporphyrin zinc complexes, having a Soret absorption band centered at 414 nm and Q-bands at 542 and 578 nm. Compared with DP₆, DP₂₄ showed a broad Soret absorption band centered at 415 nm, while DP₁₂ showed a blue-shifted shoulder at 398 nm along with a major Soret band at 411 nm. The latter observation suggests that DP₁₂ may adopt a planar geometry with a H-aggregate-type arrangement of the zinc porphyrin units along its periphery. ⁷ On the other hand, Py₂F₁–Py₂F₃ all displayed an electronic absorption band centered at 326 nm. ¹¹ Upon excitation at 326 nm, all Py₂F_n fluoresced at 694 nm from their fullerene units. ^{10,11}

As expected, DP_m (m = 6, 12, and 24) bound Py_2F_n (n =1−3) strongly to form stable $DP_m \supset Py_2F_n$. For example, upon titration with Py₂F₃ in CHCl₃ at 25 °C, DP₂₄ (1.5 \times 10⁻⁷ M) displayed a large spectral change in the Soret and Q-bands (Figure 1a), characteristic of the axial coordination of zinc porphyrins, with a clear saturation profile at a molar ratio [Py₂F₃]/[DP₂₄] exceeding 12 (Figure 1b). This spectral change profile did not give distinct isosbestic points possibly due to a large effect of the multivalency of the complexation between DP_{24} and Py_2F_3 . However, the average binding affinity (K), as estimated by simply assuming a one-to-one coordination between the individual zinc porphyrin and pyridine units, was $1.2 \times 10^6 \,\mathrm{M}^{-1}$. This value is more than 2 orders of magnitude greater than association constants reported for monodentate coordination between zinc porphyrins and pyridine derivatives.¹² Other combinations of DP_m and Py_2F_n for the titration all showed analogous spectral change profiles with a marked saturation tendency at a mole ratio $[Py_2F_n]/[DP_m]$ close to m/2.¹¹ Figure 1c shows average binding affinities K between the zinc porphyrin and pyridine units, which are almost comparable to one another in a range $1.1 \times 10^6 - 4.4 \times 10^6 \,\mathrm{M}^{-1}$ irrespective of the m and n values in DP_m and Py_2F_n , respectively. We also found that resulting complexes $DP_m \supset Py_2F_n$ are all stable under conditions for gel permeation chromatography (GPC). For example, a CHCl3 solution (0.5 mL) of a mixture of DP24 and Py_2F_3 ([DP₂₄] = 1.6 × 10⁻⁵ M, [Py₂F₃]/[DP₂₄] = 25) was loaded onto a Bio-beads S-X1 GPC column and then eluted with CHCl₃. As shown in Figure 1d, the chromatographic profile

(11) See Supporting Information.

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Table 1. Average Numbers of Py₂F_n Bound to Single DP_m Molecule in DP_m⊃Py₂F_n, upon Mixing DP_m with Py₂F_n ([Py₂F_n]/ [DP_m] = m) in CHCl₃ at 25 °C, Followed by GPC Separation with CHCl₃ as an Eluent (Figure 1d)¹¹

	Py_2F_1	Py_2F_2	Py_2F_3
DP ₆	2.9	2.6	2.6
DP_{12}	5.3	5.5	5.3
DP_{24}	9.0	9.6	9.5

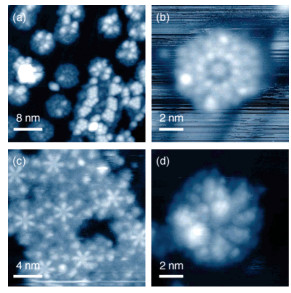


Figure 2. UHV-STM micrographs of (a, b) DP_{12} in the presence of 6 equiv of Py_2F_3 , (c) DP_6 in the presence of 3 equiv of Py_2F_1 , and (d) DP_{24} in the presence of 12 equiv of Py_2F_1 , on a Au(111) surface at a liquid nitrogen temperature. Conditions: (a) I = 1 pA, Vs = +3 V; (b) I = 1 pA, Vs = +2 V; (c) I = 2 pA, Vs = +4.5 V; (d) I = 1 pA, Vs = +2 V.

displayed two colored fractions. By means of absorption spectroscopy, the first fraction was found to contain both DP_{24} and Py_2F_3 at a molar ratio $[Py_2F_3]/[DP_{24}]$ of 9.5 (Table 1), whereas the second fraction included Py_2F_3 alone. ¹¹ The observed ratio of $[Py_2F_3]/[DP_{24}]$ is fairly close to an expected ratio of 12 for the bidentate ligation of Py_2F_3 to the zinc porphyrin units in DP_{24} . As summarized in Table 1, we likewise isolated by $GPC\ DP_m \supset Py_2F_n$ formed from the other combinations of DP_m and Py_2F_n and confirmed from their compositions that almost all the zinc porphyrin units in DP_m participate in the bidentate ligation with Py_2F_n . In contrast, P_1 , a nondendritic zinc porphyrin reference (Scheme 1), incapable of bidentate ligation with Py_2F_n , exhibited much smaller binding affinities ($\sim 2.0 \times 10^3\ M^{-1}$) toward $Py_2F_1 - Py_2F_3$ (Figure 1c).

We succeeded in visualizing some of the coordination complexes between DP_m and Py_2F_n by scanning tunneling microscopy under ultrahigh vacuum conditions (UHV-STM). For example, when a CHCl₃ solution of a mixture of DP_{12} and Py_2F_3 ([DP_{12}] = 2.0 × 10^{-6} , [Py_2F_3]/[DP_{12}] = 8.2) was deposited on a Au(111) surface by pulse injection, 13 UHV-STM at a liquid nitrogen temperature displayed petal-like patterns with a uniform diameter of 7 nm, assignable to DP_{12} adopting a planer conformation on the substrate surface (Figures 2a and 2b). UHV-STM also exhibited many bright spots at the periphery of DP_{12} molecules, which are most likely fullerene clusters of Py_2F_3 . While $DP_6 \supset Py_2F_1$ showed a petal-like pattern

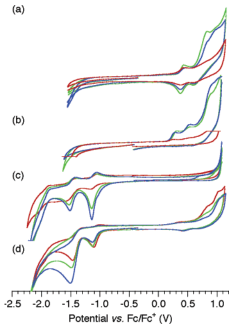


Figure 3. Cyclic voltammograms of (a) DP₆ (green), DP₁₂ (blue), and DP₂₄ (red) alone, (b) those in the presence of Py₂ ([pyridine]/[zinc porphyrin] = 1), (c) Py₂F₁ (red), Py₂F₂ (green), and Py₂F₃ (blue) alone, and (d) those in the presence of DP₁₂ ([pyridine]/[zinc porphyrin] = 1). Conditions: 0.1 M Bu₄N⁺PF₆⁻ as a supporting electrolyte, scan rate: 100 mV/s, Pt electrodes, in CH₂Cl₂ at 25 °C.

(Figure 2c), the complex likely lost most of the ligating Py_2F_1 molecules during the process for pulse injection. On the other hand, as shown in Figure 2d, $DP_{24} \supset Py_2F_1$ in UHV-STM developed a rather obscure molecular image, possibly due to a difficulty of large DP_{24} in flattening on the substrate surface.

Electrochemical properties of DP_m , Py_2F_n , and reference complexes for $DP_m \supset Py_2F_n$ were investigated by means of cyclic voltammetry (CV). In CH₂Cl₂ at 25 °C, DP₆, DP₁₂, and DP₂₄ displayed nearly identical redox properties to one another with the first oxidation potential $(E_{Ox}^{0/\cdot+})$ at around 0.40 V versus Fc/ Fc⁺ (Figure 3a). For the evaluation of the redox potentials of the pyridine-ligating zinc porphyrin units in $DP_m \supset Py_2F_n$, we utilized, in place of Py₂F_n, reference bipyridine compound Py₂ without fullerene units (Scheme 1). Upon addition of 3 equiv of Py2 to a CH2Cl2 solution of DP6, the oxidation peak, as expected, showed a cathodic shift down to $E_{\rm Ox}^{0/-+}$ of 0.28 V due to an electron donation from ligating Py2 to the zinc porphyrin units (Figure 3b). A similar electronic effect of Py₂ was observed for the oxidation potentials of DP₁₂ and DP₂₄, although the extent of such a cathodic shift was a little more explicit as the steric congestion of the dendrimer became larger: $E_{\rm Ox}^{0/\cdot+}=0.26$ (DP₁₂) and 0.23 V (DP₂₄). On the other hand, Py₂F_n showed the first reduction potential ($E_{\rm Red}^{0,-}$) at -1.08 (n=1), -1.10 (n=1), -1.1= 2), and -1.10 (n = 3) versus Fc/Fc⁺ in CH₂Cl₂ (Figure 3c), which remained virtually intact upon complexation with DP_m such as DP_{12} (Figure 3d).

Photoinduced electron transfer in $DP_m \supset Py_2F_n$ was confirmed by means of steady-state emission spectroscopy and nanosecond flash photolysis measurements. For example, excitation of a CHCl₃ solution of DP_{24} (1.5 × 10⁻⁷ M) at 550 nm resulted in a fluorescence emission from the zinc porphyrin units at 591 and 635 nm (Figure 4a). Upon titration with Py_2F_3 , the fluorescence stepwise decreased in intensity and was quenched almost completely in the final stage. Stern—Volmer constants

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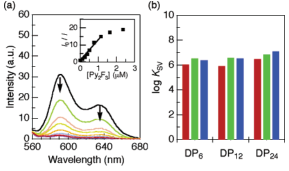


Figure 4. (a) Fluorescence spectral change and Stern–Volmer plot (inset) of DP_{24} (1.5 × 10^{-7} M) upon excitation at 550 nm in the presence of Py_2F_3 ($[Py_2F_3]/[DP_{24}] = 0$, 0.85, 1.7, 2.6, 3.4, 5.1, 7.7, 11, and 16 in CHCl₃ at 25 °C under Ar. (b) Stern–Volmer constants of DP_6 , DP_{12} , and DP_{24} in the presence of Py_2F_1 (red), Py_2F_2 (green), and Py_2F_3 (blue).

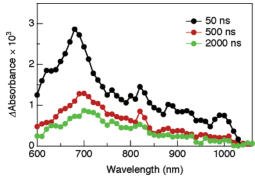


Figure 5. Nanosecond transient absorption spectra at 20 °C of a CH_2Cl_2 solution of DP_6 (1.7 \times 10⁻⁵ M) containing 3 equiv of Py_2F_3 upon photoexcitation at 532 nm.

 $(K_{SV}, \text{ Figure 4b})^{11}$ evaluated for the combination of DP_m (m =6, 12, and 24) and Py_2F_n (n = 1-3), were roughly comparable to their average binding affinities K (Figure 1c), indicating that the fluorescence quenching of DP_m is caused by the coordination of electron-accepting Py₂F_n. By means of transient absorption spectroscopy, we confirmed the occurrence of an electron transfer from the photoexcited zinc porphyrin units in DP_m to the fullerene units in ligating Py₂F_n. Excitation of a CH₂Cl₂ solution of DP₆ (1.7 \times 10⁻⁵ M) at 532 nm in the presence of 3 equiv of Py₂F₃ resulted in a transient absorption spectrum (Figure 5) with bands at around 680 and 1000 nm assignable to the cation and anion radicals of the zinc porphyrin and fullerene units, respectively. Broad absorption bands, observed at 750-900 nm, are due to the excited triplet states of these two units. Since the fluorescence spectral profiles of $DP_m \supset Pv_2F_n$ such as DP₆⊃Py₂F₃ under the conditions employed here did not show any sign of energy transfer from the photoexcited zinc porphyrin units in DP_m to the fullerene units of Py_2F_n , 11,14 we conclude that the quenching of the zinc porphyrin fluorescence observed for $DP_m \supset Py_2F_n$ (Figure 4) is mostly due to the intracomplex photoinduced electron transfer between them.

Time-resolved emission spectroscopy was employed for investigating the charge-separation event in $DP_m \supset Py_2F_n$. Upon excitation at 420 nm in CH_2Cl_2 at 20 °C, the fluorescence of DP_6 (1.0 × 10⁻⁶ M) in the presence of 3 equiv of fullerenefree bipyridine displayed a monoexponential decay profile at 590 nm with a lifetime (τ_f) of 1.6 ns. On the other hand, when

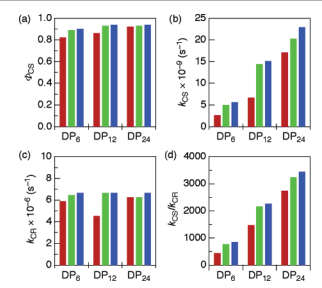


Figure 6. (a) Charge-separation quantum yields ($Φ_{CS}$), (b) charge-separation rate constants (k_{CS}), (c) charge-recombination rate constants (k_{CR}), and (d) k_{CS}/k_{CR} of DP₆, DP₁₂, and DP₂₄ in the presence of Py₂F₁ (red), Py₂F₂ (green), and Py₂F₃ (blue) in CH₂Cl₂ at 20 °C.

fullerene-appended Py₂F₁ was used in place of Py₂, a fast-decay fluorescing component with τ_f of 320 ps appeared in addition to the slow-decay component ($\tau_{\rm f}=1.1$ ns). This fast-decay component obviously originates from the charge-separation process caused by the photoinduced electron transfer from DP₆ to Py₂F₁. In the presence of Py₂F₂ or Py₂F₃ instead of Py₂F₁, the lifetime of the fast-decay component of DP₆ became much shorter; $\tau_f = 170 \text{ (Py}_2F_2)$ and 150 ps (Py₂F₃). In contrast with DP₆⊃Py₂, DP₁₂⊃Py₂ and DP₂₄⊃Py₂ exhibited a biexponential fluorescence decay at 590 nm possibly due to a dense packing of the zinc porphyrin units (vide ante) on the dendrimer surface. 11 Use of Py_2F_n in place of Py_2 to allow complexation with DP₁₂ or DP₂₄ resulted in an additional decay component with a much shorter τ_f value, owing to the electron transfer from DP_m to Py_2F_n . 11 On the basis of the fluorescence decay profiles, the charge-separation rate constants (k_{CS}) and quantum yields (Φ_{CS}) were evaluated according to the following equations by comparison of the shortest τ_f components of $DP_m \supset Py_2F_n$ with average τ_f values of $DP_m \supset Py_2$:

$$\begin{split} k_{\mathrm{CS}} &= \left(\frac{1}{\tau_{\mathrm{f}}}\right)_{\mathrm{DP_{\mathrm{m}}} \supset \mathrm{Py}_{2}\mathrm{F_{\mathrm{n}}}} - \left(\frac{1}{\tau_{\mathrm{f}}}\right)_{\mathrm{DP_{\mathrm{m}}} \supset \mathrm{Py}_{2}} \\ \Phi_{\mathrm{CS}} &= \left[\left(\frac{1}{\tau_{\mathrm{f}}}\right)_{\mathrm{DP_{\mathrm{m}}} \supset \mathrm{Py}_{2}\mathrm{F_{\mathrm{n}}}} - \left(\frac{1}{\tau_{\mathrm{f}}}\right)_{\mathrm{DP_{\mathrm{m}}} \supset \mathrm{Py}_{2}} \right] / \left(\frac{1}{\tau_{\mathrm{f}}}\right)_{\mathrm{DP_{\mathrm{m}}} \supset \mathrm{Pv}_{2}\mathrm{F}} \end{split}$$

The Φ_{CS} values thus evaluated are all high, ranging from 0.82 to 0.94 (Figure 6a). Thus, the photoinduced electron transfer in $DP_m \supset Py_2F_n$ occurred very efficiently, irrespective of the magnitudes of m and n. Of further interest, the k_{CS} value increases following the order of $Py_2F_1 < Py_2F_2 < Py_2F_3$ and enhances, to a much greater extent, from DP_6 to DP_{12} and then to DP_{24} (Figure 6b). For example, the k_{CS} value of 2.3×10^{10} s⁻¹, as evaluated for $DP_{24} \supset Py_2F_3$, is an order of magnitude greater than that of $DP_6 \supset Py_2F_1$ (0.26 \times 10¹⁰ s⁻¹) and obviously the largest among those of the $DP_m \supset Py_2F_n$ family. These trends indicate an interesting possibility that dense packing of both the zinc porphyrin and fullerene units on the dendrimer surface

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plays an essential role in facilitating the charge-separation process. As expected, nondendritic zinc porphyrin reference P_1 in the presence of Py_2F_n showed much smaller k_{CS} values such as 0.031×10^{10} (n = 1), 0.12×10^{10} (n = 2), and 0.13×10^{10} s⁻¹ (n = 3).

We also investigated the charge-recombination events in $DP_m \supset Py_2F_n$, using the decay profiles in CH_2Cl_2 of the transient absorption band at 1000 nm due to C_{60} (Figure 5). In every case, the decay profile was satisfactorily fitted with twoexponential components, where the faster decay is attributed to the charge-recombination process, while the slower decay originates from the excited triplet species. The chargerecombination rate constants (k_{CR} , Figure 6c) thus obtained are all within a narrow range from 4.5×10^6 to 6.7×10^6 s⁻¹, and the lifetimes of the charge-separation state (τ_{RIP} ; 150–220 ns) are comparable to one another. 11 These trends are quite reasonable since the donor-acceptor distance in $DP_m \supset Pv_2F_n$ should not be much dependent on the degrees of steric congestion of the two ligating components but is primarily determined by the length of the spacer unit between the bipyridine and fullerene units in Py₂F_n. The successful determination of the k_{CS} values for $DP_m \supset Py_2F_n$ allowed us to compare the ratios of $k_{\rm CS}/k_{\rm CR}$, which have often been used as a measure for the excellence of photoinduced electron-transfer systems. 15 Of interest, the values of k_{CS}/k_{CR} (Figure 6d) are all large in a range from 450 ($DP_6 \supset Py_2F_1$) to 3400 ($DP_{24} \supset Py_2F_3$). In particular, the k_{CS}/k_{CR} ratio for $DP_{24} \supset Py_2F_3$ is more than an order of magnitude greater than those reported for precedent porphyrin-fullerene supramolecular dyads and triads.¹⁵ It is obvious that a larger number of the fullerene units in DP₂₄⊃Py₂F₃ could enhance the probability of the electron transfer from the zinc porphyrin units. However, in addition to this, one can also presume that an efficient energy migration along the densely packed zinc porphyrin array⁹ may enhance the opportunity for this electron transfer. When the degrees of fluorescence quenching $(1 - I/I_0)$ [I and I_0 ; fluorescence intensities with and without the quencher, respectively], observed for DP₆, DP₁₂, and DP₂₄ upon titration with Py₂F₃, are plotted against the ratio of the numbers of the fullerene and zinc porphyrin units ([fullerene]/ [zinc porphyrin]), 11 it is clear that the quenching of the excited singlet state of the zinc porphyrin units by Py₂F₃ is more efficient as m in DP_m is larger from 6 to 12 and then to 24 (Figure 7). For example, the degree of fluorescence quenching of 0.6 can be attained for DP_{24} only at [fullerene]/[zinc porphyrin] = 0.19, whereas it requires a much larger ratio of [fullerene]/[zinc porphyrin] such as 0.64 for DP₆. Since the average binding affinities of Py₂F₃ toward DP₆ and DP₂₄ are not much different

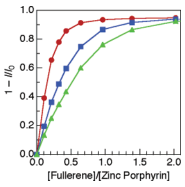


Figure 7. Plots of degrees of fluorescence quenching $(1 - I/I_0)$ versus [fullerene]/[zinc porphyrin] upon titration of DP₆ (green), DP₁₂ (blue), and DP₂₄ (red) ([zinc porphyrin] = 3.6×10^{-6} M) with Py₂F₃ in CHCl₃ at 25 °C. Concentrations of fullerene and zinc porphyrin units are given by $3 \times [\text{Py}_2\text{F}_3]$ and $m \times [\text{DP}_m]$, respectively.

from one another (Figure 1c), it is obvious that the energy migration along the zinc porphyrin array in DP_{24}^9 greatly facilitates the electron transfer to the fullerene units.

Conclusions

By means of a multivalent surface ligation of dendritic macromolecules, 16 we constructed, using DP_m (m = 6, 12, and24) and Py_2F_n (n = 1-3), a photoactive layer, consisting of electron-donating zinc porphyrin and electron-accepting fullerene arrays on the dendrimer surface. Upon increment of the numbers of these donor and acceptor units, the electron transfer reaction was remarkably facilitated, while the recombination of the resulting charge-separated state remained virtually intact. Consequently, among the $DP_m \supset Py_2F_n$ family, $DP_{24} \supset Py_2F_3$ accommodating 24 zinc porphyrin units and roughly 30 fullerene units on the dendrimer surface (Table 1) achieved the largest ratio of the charge-separation to charge-recombination rate constants (3400), which is even 1 order of magnitude greater than those of precedent examples. Application of this molecular design strategy to the development of optoelectronic materials is one of the subjects worthy of further investigations.

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Supporting Information Available: Details for synthesis, characterization, and spectral data (PDF). This material is available free of charge via the Internet at http://pubs.acs.org. See any current masthead page for ordering information and Web access instructions.

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