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Review—Two Different Multiple Photosynthetic Reaction Centers Using Either Zinc Porphyrinic Oligopeptide-Fulleropyrrolidine or Free-Base Porphyrinic Polypeptide-Li+@C$_{60}$ Supramolecular Complexes

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An overview of two successful examples of photosynthetic reaction center models combined with light-capturing antenna chromophores is presented. In the first example, supramolecular complexes are formed between flexible zinc porphyrinic oligopeptides and fulleropyrrolidinol bearing either a pyridine or imidazole functionalized C60 via a coordination bond plus π–π interactions. The excited energy migration occurs between porphyrin units followed by charge separation. The charge separation (CS) lifetimes of the supramolecular oligopeptide complexes have been elongated by increasing the generation of the porphyrins, enabling us to attain the longest lifetime (0.84 ms) for the P(ZnP)8-ImC60 supramolecular system in PhCN solution at 298 K, ever reported for supramolecular complexes. In the second example, free-base porphyrin polypeptides (P(H2P)n; n = 4 and 8) form supramolecular complexes with Li@C60 in PhCN, in which the binding is much stronger than C60. Efficient energy migration occurs between porphyrins in P(H2P)n. The triplet CS states derived from 3Li@C60 had long lifetimes due to spin-forbidden back electron transfer. The triplet CS lifetime becomes longer upon increasing the number of H2P due to the charge migration among porphyrins. The present study provides valuable insight into the energy and electron transfer processes leading to long-lived charge separated states in artificial photosynthetic antenna-reaction center models.

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In natural photosynthetic systems, the solar energy is collected by pigment molecules attached to the light harvesting complexes. In these units, the chlorophylls are held in a favored spacing and orientation by fairly short α-helical polypeptides.1–3 When a photon hits one of the chlorophylls, the absorbed energy spreads extremely rapidly to the others until the reaction center is reached within the cell membrane where the solar energy is converted into chemical energy used by the cell to grow. In this way, the energy contained in a single photon is conducted in a very short time and with minimal loss of energy from the point where it is absorbed to where it is needed.

Effective effort has so far been devoted to mimicking light harvesting and charge separation processes in natural photosynthesis.4–20 Multiporphyrin arrays have been employed as light-harvesting units.21–30 Light-harvesting and charge-separation units have been combined by coordination bonds between metallochlorophylls and electron acceptor moieties bearing Lewis base ligands.36–44 Thus, metal centres were required for the construction of supramolecular complexes between porphyrins acting as light harvesting units and electron acceptors containing Lewis base ligands for coordination to the metal centres.36–44 To mimic the natural light harvesting antennae, we synthesized 15 years ago a family of porphyrin-functionalized α-polypeptide such as the octamer (Fig. 1).34 A peptidic backbone was chosen owing to the remarkable ability of peptides to establish secondary Structures such as α-helices, that renders reasonable to expect, beyond a certain degree of oligomerisation, a helical spatial arrangement of the chromophores. Such a conformation should induce an overlap of the porphyrins, which may then undergo sufficient electronic coupling to promote a good exciton migration along the set of porphyrins. It has been the first results presented at the ECS meeting in 2003. Since then, these polypeptides have been extensively studied and many properties have been highlighted.46–51 In the present mini-review, we focus on the preparation and study of multiple reaction centers.

Mimicking the natural photosynthetic process requires both synthesis of the light-harvesting and charge-separation units, which should be linked by covalent or noncovalent bonding.5–12,52–54 Non-covalent bonding is certainly more favorable as compared to covalent bonding from the viewpoint of mimicking the biological photosystems, in which non-covalent interaction plays a pivotal role to regulate the energy- and electron-transfer processes, and easy access toward construction of a variety of donor-acceptor systems.16–19,54–60 A variety of multiporphyrin arrays have so far been utilized as the light-harvesting units to mimic efficient energy transfer in the photosynthesis.60,61,62–64 Among multi-porphyrin arrays, porphyrin oligomers with oligopeptidic backbones in which porphyrins are held in a favored spacing and orientation by fairly short helical oligopeptides have merited special attention in relevance to the natural light-harvesting complex in which chlorophyll and oligopeptides are well-organized in the protein.65,66 We highlight in this mini-review the elaboration of two different multiple photosynthetic reaction centres using either zinc
porphyrinic oligopeptide–fulleropyrrolidine or free-base porphyrinic polypeptide–Li @ C60 supramolecular complexes.

Results and Discussion

Multiple photosynthetic reaction centers using zinc porphyrinic oligopeptide–fulleropyrrolidine supramolecular complexes.—We first synthesized multiple photosynthetic reaction centers composed of light-harvesting multiporphyrin units and charge-separation units, which have been combined by using non-covalent binding including coordination bond and π–π interaction. Zinc porphyrinic oligopeptides with various numbers of porphyrin units [P(ZnP)n; n = 2, 4, 8]44,45,49,65,66 are used as light harvesting multiporphyrin units, which are bound to electron acceptors of fulleropyrrolidine bearing a pyridine (PyC60)67 or imidazole coordinating ligand (ImC60)66,68 as shown in Fig. 2.

The laser flash photolysis of the supramolecular complexes of P(ZnP)6 with PyC60 and ImC60 has revealed the formation of the long-lived charge-separated (CS) state. The details of synthesis of porphyrin–peptide oligomers have been reported previously65,66. The UV–visible spectra of P(ZnP)6 in benzonitrile (PhCN) at 298 K changed upon addition of PyC60, where the Soret band is red-shifted with an isosbestic point at 436 nm. The absorbance change exhibits a saturation behavior with increasing PyC60 concentration. This indicates that PyC60 forms a 1:1 supramolecular complex with the ZnP moiety of P(ZnP)6.69 It has well been established that such nitrogenous bases readily bind to zinc porphyrins with a 1:1 stoichiometry.68

As P(ZnP)6 forms the supramolecular complex with PyC60, the fluorescence of P(ZnP)6 is strongly quenched by the intra-supramolecular electron transfer from the singlet excited state of the ZnP moiety (ZnPπ) in P(ZnP)6 to bound PyC60 (i.e., static quenching) and not by an intermolecular electron transfer from ZnP6 to unbound PyC60 (i.e., dynamic quenching).69 In fact, if the fluorescence quenching is analyzed by the Stern-Volmer equation of bimolecular electron transfer [Eq. 1], the rate constant of bimolecular photo-induced electron transfer is estimated as 6 × 10^13 M^−1 s^−1, which is significantly larger than the diffusion rate constant in PhCN (5.6 × 10^7 M^−1 s^−1).

\[ (I_0/I) = 1 + k_{sv}[PyC_{60}] \]  

The values of the formation constants (K) of zinc porphyrinic oligopeptides and C60 derivatives (PyC60 vs ImC60) determined by the absorption spectral change are listed in Table I. In the case of PyC60 as an electron acceptor, the K value increases with increasing number of zinc porphyrins in an oligopeptide unit. No supramolecular complex formation was observed in the case of zinc tetraphenylporphyrin with C60 derivatives67 or the porphyrinic oligopeptides with pristine C60 in PhCN.48,49 The strong association between P(ZnP)4 and C60Py may originate from the strong π–π interaction between two zinc porphyrins and C60Py in addition to the axial coordination of C60Py to the zinc porphyrin. In the case of ImC60, however, the highest K value was obtained for the P(ZnP)4-ImC60 complex. This indicates that ImC60 is much more strongly bound by the oligopeptide, P(ZnP)4, than PyC60.

It should be noted that the apparent formation constants determined from the fluorescence quenching are significantly larger than those determined from the UV–vis spectral change, and the difference in such association constants increases with increasing generation of porphyrinic oligopeptides. This indicates that the excited energy migration between the porphyrin units occurs efficiently prior to the electron transfer to the bound C60 moiety. Such difference in the formation constants is more enhanced in the P(ZnP)4-ImC60 supramolecular complexes as compared to that of PyC60 (Table I). For example, an extremely efficient energy transfer may occur in the P(ZnP)4-ImC60 system judging from the large difference in the K values determined by the absorption change and by the fluorescence quenching (1.5 × 10^5 vs 3.3 × 10^7 M^−1). We have also checked the structural change of the oligopeptidic backbone when the porphyrinic oligopeptide forms the supramolecular complex with C60 derivatives by the circular dichroism spectroscopy, which exhibited the Cotton effect originating from the porphyrin Soret band at 431 nm. The occurrence of photoinduced electron transfer in the supramolecular complex in PhCN was confirmed by the transient absorption spectra of the supramolecular complex of P(ZnP)6 with PyC60 using nanosecond laser flash photolysis. The laser photoexcitation at 561 nm of the supramolecular complex of P(ZnP)6 with PyC60 results in the formation of the CS state as indicated by the transient absorption spectra in Fig. 3a. The absorption band due to ZnP− is clearly observed at 1000 nm together with that due to ZnP+.68

A kinetic analysis of the observed CS decay is derived from the first-order plots at different initial CS concentrations (Fig. 3b). The second-order kinetics would be involved for the decay time profile. In fact, the corresponding second-order plots (Fig. 3c) are clearly non-linear and the initial slope varies depending on the CS concentration. Thus, the observed decay is ascribed to back electron transfer in the supramolecular complex rather than intermolecular back electron transfer between ZnP and PyC60.

The CS lifetimes of the supramolecular complexes of other porphyrins [P(ZnP)6; n = 2, 4] and fullerene derivatives (PyC60 and ImC60) were also determined and listed in Table I. The CS lifetime...
becomes longer with increasing generation of porphyrinic oligopeptides. This may result from efficient hole migration between the porphyrin units following photo-induced electron transfer in the supramolecular complex. Thus, we have succeeded in attaining the longer CS lifetime by increasing the generation of porphyrinic oligopeptides. Such a long-lived CS state was also confirmed by the ESR measurements under photo-irradiation of the supramolecular complex of P(ZnP)₈ with PyC₆₀ or ImC₆₀ in frozen PhCN at 173 K. Under photo-irradiation, the isotropic ESR signals corresponding to the zinc porphyrin radical cation and fullerene radical anion are clearly observed at ₀ = 2.002 and 2.000, respectively.

In conclusion, multiple photosynthetic reaction centers combined with light-capturing antenna chromophores have successfully been constructed using supramolecular complexes formed between flexible porphyrinic oligopeptides and fulleropyrrolidine bearing a pyridine (PyC₆₀) or imidazole (ImC₆₀) ligand via a coordination bond plus π–π interaction. The excited energy migration occurs between porphyrin units followed by charge separation. The CS lifetimes of the supramolecular oligopeptide complexes have been elongated by increasing the generation of the porphyrins, enabling us to attain the longest lifetime (0.84 ms) for the P(ZnP)₈-ImC₆₀ supramolecular system in PhCN solution at 298 K ever reported for supramolecular complexes.

### Table 1. Formation constants (Kₐ), lifetimes and quantum yield of the CS state of the supramolecular complexes in PhCN at 298 K.

<table>
<thead>
<tr>
<th>Porphyrin</th>
<th>Acceptor</th>
<th>UV–vis Fluorescence</th>
<th>τ₀/mea</th>
<th>τ_CS/mea</th>
</tr>
</thead>
<tbody>
<tr>
<td>P(ZnP)₈</td>
<td>PyC₆₀</td>
<td>2.3 × 10⁴</td>
<td>1.5 × 10⁵</td>
<td>0.70</td>
</tr>
<tr>
<td>P(ZnP)₄</td>
<td>PyC₆₀</td>
<td>5.2 × 10³</td>
<td>1.2 × 10⁵</td>
<td>0.40</td>
</tr>
<tr>
<td>P(ZnP)₂</td>
<td>PyC₆₀</td>
<td>4.0 × 10³</td>
<td>9.0 × 10⁴</td>
<td>0.34</td>
</tr>
<tr>
<td>P(ZnP)₈</td>
<td>ImC₆₀</td>
<td>1.5 × 10⁴</td>
<td>3.3 × 10⁵</td>
<td>0.84</td>
</tr>
<tr>
<td>P(ZnP)₄</td>
<td>ImC₆₀</td>
<td>5.9 × 10⁴</td>
<td>8.5 × 10⁴</td>
<td>0.38</td>
</tr>
</tbody>
</table>

a) Experimental error within 5%.

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Multiple photosynthetic reaction centers of porphyrinic polypeptide–Li⁺@C₆₀ supramolecular complexes. — In the latter example, metal centers were required for the construction of supramolecular complexes between porphyrins acting as light harvesting units and electron acceptors containing Lewis base ligands for coordination to the metal centers. As such there have been only a few reports on supramolecular complexes of free baseporphyrin arrays and electron acceptors without Lewis base ligands, in which the binding is rather weak. We thus focused on the construction of supramolecular complexes of free base porphyrin polypeptides (P(H₂P)ₙ; n = 4 and 8) with lithium ion-encapsulated C₆₀ (Li⁺@C₆₀) in which binding is much stronger than the case of C₆₀ in benzonitrile (PhCN). The photo-dynamics were studied by femtosecond and nanosecond laser-induced transient absorption and fluorescence lifetime measurements.

Free base porphyrin polypeptide (P(H₂P)ₙ) were synthesized and characterized as reported previously. Upon mixing a PhCN solution of Li⁺@C₆₀ with that of P(H₂P)ₙ, the intensity of the Soret band decreased with an increasing concentration of Li⁺@C₆₀. A Soret band was slightly red-shifted from 425 to 427 nm by the addition of Li⁺@C₆₀ to the solution. From the net absorption change at 425 nm in which the absorption due to Li⁺@C₆₀ was subtracted, a linear Benesi–Hildebrand plot was obtained, indicating that each porphyrin unit of P(H₂P)ₙ forms a 1:1 supramolecular charge-transfer complex with Li⁺@C₆₀ independently with approximately the same binding constant of the unit of M⁻¹. The binding constant at 298 K was determined from the intercept and the slope to be 2.1 × 10⁴ M⁻¹, which is significantly larger than that of C₆₀ (5.3 × 10³ M⁻¹). The stronger binding of Li⁺@C₆₀ as compared with C₆₀ may result from the stronger electron acceptor ability of Li⁺@C₆₀, which facilitates the charge-transfer interaction.
as reported for the stronger charge-transfer binding of Li$^+@C_{60}$ with corannulene.$^{73}$ Similarly the binding constant of the supramolecular complex of P(H$_2$P)$_4$ was determined from the spectral titration to be $6.2 \times 10^3$ M$^{-1}$. In the case of P(H$_2$P)$_2$ and P(H$_2$P)$_1$, however, the spectral change was too small to be able to determine the binding constants accurately. The binding constants are summarized in Table II. Thus, multiporphyrins may facilitate charge transfer interactions through the encapsulation of Li$^+@C_{60}$ by multiple porphyrins.

Fluorescence of P(H$_2$P)$_8$ was quenched by the addition of a PhCN solution of Li$^+@C_{60}$. A Benesi-Hildebrand plot for fluorescence quenching afforded the apparent binding constant of the supramolecular complex of P(H$_2$P)$_8$ with Li$^+@C_{60}$ being $8.5 \times 10^4$ M$^{-1}$, which is significantly larger than the value determined from the absorption spectral titration ($2.1 \times 10^4$ M$^{-1}$). Similarly the apparent binding constant of P(H$_2$P)$_4$ with Li$^+@C_{60}$ was determined from the fluorescence quenching to be $1.6 \times 10^4$ M$^{-1}$, which is also larger than the value determined from the absorption spectral titration (Table II). Such larger apparent binding constants determined from the fluorescence quenching than those determined from the absorption spectral titration indicate that energy migration occurs between the singlet excited state of H$_2$P (H$_2$P*) and adjacent H$_2$P in P(H$_2$P)$_n$ (n = 4 and 8).

The energy migration rate constant was determined by the fluorescence lifetime measurements of P(H$_2$P)$_8$ with Li$^+@C_{60}$. The fluorescence of $^1$P(H$_2$P)$_8$* exhibited a single exponential decay with the lifetime of 11 ns. In the presence of Li$^+@C_{60}$, the decay of $^1$P(H$_2$P)$_8$* is well analyzed by three exponentials with lifetimes of 100 ps, 3.0 ns and 11 ns. The fast component corresponds to the lifetime of $^1$H$_2$P* in the supramolecular complex with Li$^+@C_{60}$-95% of the fast component agrees with that predicted by the binding constant determined from the absorption spectral titration. The third component has the same lifetime as that without Li$^+@C_{60}$-corresponding to the fluorescence lifetime of $^1$(H$_2$P)$_8$*. The second component corresponds to energy migration between $^1$H$_2$P* and the adjacent H$_2$P. 5% of the sum of the first and second components agrees with that predicted from the binding constant determined from fluorescence quenching.

Femtosecond laser-induced transient absorption spectra of P(H$_2$P)$_8$ were recorded in the absence and presence of Li$^+@C_{60}$ (Fig. 5). In each case, only the singlet-singlet absorption of $^1$P(H$_2$P)$_8$* was observed, because H$_2$P was exclusively excited at the excitation wavelength (425 nm) in Fig. 5. The decay of $^1$P(H$_2$P)$_8$* exhibits three exponentials with lifetimes of 6 ps, 90 ps and 2 ns. The ratio of the fast component increased upon increasing the number of H$_2$P in P(H$_2$P)$_n$ (n = 4 and 8). In the case of P(H$_2$P)$_1$, however, no fast component was observed. The ratio of the fast component increased upon increasing the number of H$_2$P in P(H$_2$P)$_n$. 96% of the fast component of P(H$_2$P)$_8$ decreased upon decreasing the laser intensity (Fig. 5b). These results indicate that the fast component results from the singlet–singlet annihilation of two $^1$H$_2$P* moieties in P(H$_2$P)$_n$ (n = 4 and 8).

In the presence of Li$^+@C_{60}$ (20 mM), the decay of $^1$P(H$_2$P)$_8$* exhibited three exponentials with lifetimes of 6 ps, 90 ps and 2 ns. The first fast component resulted from the singlet-singlet

Figure 3. (a) Transient absorption spectra of P(ZnP)$_8$ (2.9 $\times$ $10^{-6}$ M) in the presence of PyC$_{60}$ (4.9 $\times$ $10^{-3}$ M) in deaerated PhCN at 298 K taken at 70 (solid line with black circles) and 350 ms (solid line with white circles) after laser excitation at 561 nm (4 mJ pulse$^{-1}$), respectively. (b) Time profiles of the absorption at 1000 nm due to PyC$_{60}$-- with different laser powers (4 and 1 mJ pulse$^{-1}$) at 298 K. Inset: first-order plots. (c) Second-order plots.
annihilation as observed without Li\(^+\)@C\(_{60}\). The second component (90 ps) may result from electron transfer from 1H\(_2\)P* to Li\(^+\)@C\(_{60}\) in the supramolecular complex. Because the transient absorption due to the charge-separated state (H\(_2\)P\(^+\) and Li\(^+\)@C\(_{60}\)\(^-\)) was not observed in Fig. 5a, where only the singlet-singlet absorption of 1H\(_2\)P\(^*\) is seen, back electron transfer may be much faster than the forward electron transfer. The third component may correspond to the energy migration between 1H\(_2\)P\(^*\) and the adjacent H\(_2\)P, because its lifetime agrees with the lifetime of energy migration in the fluorescence lifetime measurements. In the presence of high concentration of Li\(^+\)@C\(_{60}\) (120 mM), the component of electron transfer from 1H\(_2\)P\(^*\) to Li\(^+\)@C\(_{60}\) in the supramolecular complex increased. The three component decay was also observed for P(H\(_2\)P)\(_4\) and P(H\(_2\)P)\(_2\) with Li\(^+\)@C\(_{60}\). In the case of P(H\(_2\)P)\(_1\), only single exponential decay with a lifetime of 11 ns was observed in the absence and presence of Li\(^+\)@C\(_{60}\) because the binding of Li\(^+\)@C\(_{60}\) to P(H\(_2\)P)\(_1\) was negligible.

Nanosecond laser-induced transient absorption spectra of P(H\(_2\)P)\(_8\) with Li\(^+\)@C\(_{60}\) (40 mM) at the excitation wavelength of 532 nm are shown in Fig. 6, where the transient absorption band at 730 nm due to the triplet excited state of Li\(^+\)@C\(_{60}\) (3Li\(^+\)@C\(_{60}\)*) is observed together with the absorption band at 1035 nm due to

Table II. Association constant (K, M\(^{-1}\)) of P(H\(_2\)P)\(_n\) with Li\(^+\)@C\(_{60}\) and C\(_{60}\) in PhCN determined from UV–vis absorption and fluorescence spectral titration.

<table>
<thead>
<tr>
<th>Porphyrin</th>
<th>Li(^+)@C(_{60})</th>
<th>Fluorescence</th>
<th>C(_{60})</th>
<th>Fluorescence</th>
</tr>
</thead>
<tbody>
<tr>
<td>P(H(_2)P)(_1)</td>
<td>1.1 \times 10(^4)</td>
<td>1.1 \times 10(^4)</td>
<td>1.1 \times 10(^4)</td>
<td>1.1 \times 10(^4)</td>
</tr>
<tr>
<td>P(H(_2)P)(_2)</td>
<td>1.6 \times 10(^3)</td>
<td>1.6 \times 10(^3)</td>
<td>1.6 \times 10(^3)</td>
<td>1.6 \times 10(^3)</td>
</tr>
<tr>
<td>P(H(_2)P)(_4)</td>
<td>2.1 \times 10(^3)</td>
<td>2.1 \times 10(^3)</td>
<td>2.1 \times 10(^3)</td>
<td>2.1 \times 10(^3)</td>
</tr>
<tr>
<td>P(H(_2)P)(_8)</td>
<td>6.2 \times 10(^3)</td>
<td>6.2 \times 10(^3)</td>
<td>6.2 \times 10(^3)</td>
<td>6.2 \times 10(^3)</td>
</tr>
</tbody>
</table>

a) Too small to determine accurately. b) Taken from Ref. 48.

in Fig. 5a, where only the singlet-singlet absorption of 1H\(_2\)P\(^*\) is seen, back electron transfer may be much faster than the forward electron transfer. The third component may correspond to the energy migration between 1H\(_2\)P\(^*\) and the adjacent H\(_2\)P, because its lifetime agrees with the lifetime of energy migration in the fluorescence lifetime measurements. In the presence of high concentration of Li\(^+\)@C\(_{60}\) (120 mM), the component of electron transfer from 1H\(_2\)P\(^*\) to Li\(^+\)@C\(_{60}\) in the supramolecular complex increased. The three component decay was also observed for P(H\(_2\)P)\(_4\) and P(H\(_2\)P)\(_2\) with Li\(^+\)@C\(_{60}\). In the case of P(H\(_2\)P)\(_1\), only single exponential decay with a lifetime of 11 ns was observed in the absence and presence of Li\(^+\)@C\(_{60}\) because the binding of Li\(^+\)@C\(_{60}\) to P(H\(_2\)P)\(_1\) was negligible.

Nanosecond laser-induced transient absorption spectra of P(H\(_2\)P)\(_8\) with Li\(^+\)@C\(_{60}\) (40 mM) at the excitation wavelength of 532 nm are shown in Fig. 6, where the transient absorption band at 730 nm due to the triplet excited state of Li\(^+\)@C\(_{60}\) (3Li\(^+\)@C\(_{60}\)*) is observed together with the absorption band at 1035 nm due to
Li\(^+\)@C\(_{60}\). The decay of the absorption at 730 nm coincides with the appearance of Li\(^+\)@C\(_{60}\). Thus, electron transfer from P(H\(_2\)P)\(_8\) to Li\(^+\)@C\(_{60}\) occurs to produce the triplet charge-separate (CS) state of P(H\(_2\)P)\(_8\)\(^+\) and Li\(^+\)@C\(_{60}\). The transient absorption due to P(H\(_2\)P)\(_8\)\(^+\) is overlapped with that of Li\(^+\)@C\(_{60}\) in the 600–700 nm region. The photoexcitation of 532 nm resulted in the excitation of Li\(^+\)@C\(_{60}\) leading to the formation of Li\(^+\)@C\(_{60}\) via intersystem crossing. The decay of the absorbance at 1035 nm due to Li\(^+\)@C\(_{60}\) obeyed first-order kinetics with the lifetime of 210 ms (Fig. 6b). Thus, back electron transfer from Li\(^+\)@C\(_{60}\) to P(H\(_2\)P)\(_8\) occurs in the supramolecular complex. The lifetime is long because of the spin-forbidden back electron transfer in the triplet CS state.

Similarly, the triplet CS states derived from Li\(^+\)@C\(_{60}\) were observed for P(H\(_2\)P)\(_8\) with Li\(^+\)@C\(_{60}\). The energy migration in the supramolecular complexes of P(H\(_2\)P)\(_n\) with Li\(^+\)@C\(_{60}\) are summarized in Table III together with the lifetimes of the triplet CS states. The CS lifetime increases upon increasing the number of H\(_2\)P in P(H\(_2\)P)\(_n\). Such an increase in the CS lifetime may result from the charge migration between H\(_2\)P and the adjacent H\(_2\)P in P(H\(_2\)P)\(_n\). The energy diagram of photoinjection of P(H\(_2\)P)\(_n\) with Li\(^+\)@C\(_{60}\) is given in Scheme 1. The energies of the excited state of porphyrin and Li\(^+\)@C\(_{60}\) were determined by UV–vis, fluorescence and phosphorescence spectral measurements. The energy of the CS state was determined to be 0.82 eV in PhCN from the one-electron oxidation potential of porphyrin (+0.96 V vs SCE) and reduction potential of Li\(^+\)@C\(_{60}\) (+0.14 V vs SCE) by the electrochemical measurements. Femtosecond laser excitation of the supramolecular complex of P(H\(_2\)P)\(_n\) with Li\(^+\)@C\(_{60}\) at 425 nm resulted in electron transfer from H\(_2\)P to the bound Li\(^+\)@C\(_{60}\) to produce the singlet CS state and energy transfer from H\(_2\)P to the adjacent H\(_2\)P, followed by electron transfer to Li\(^+\)@C\(_{60}\), when it is bound H\(_2\)P. The singlet CS state decays rapidly to the ground state. Nanosecond laser excitation at 532 nm resulted in the formation of the triplet excited state of excess Li\(^+\)@C\(_{60}\), followed by intersystem crossing to produce Li\(^+\)@C\(_{60}\), to which intermolecular electron transfer occurs from H\(_2\)P to produce the triplet CS state with a long lifetime because of the spin-forbidden back electron transfer to the ground state.

We can thus confirm that free base porphyrin polypeptides (P(H\(_2\)P)\(_n\); n = 4 and 8) form supramolecular complexes with Li\(^+\)@C\(_{60}\) in PhCN, in which the binding is much stronger than C\(_{60}\). Efficient energy migration occurs between porphyrins in P(H\(_2\)P)\(_n\). Although the lifetimes of the singlet CS states were too short to be detected, the triplet CS states derived from Li\(^+\)@C\(_{60}\) had long lifetimes because of the spin-forbidden back electron transfer. The triplet CS lifetime becomes longer upon increasing the number of H\(_2\)P due to the charge migration among porphyrins. The present study provides valuable insight into the energy and electron transfer in multiple reaction centers.

Conclusions

In conclusion, multiple photosynthetic reaction centers combined with light-capturing antenna chromophores have successfully been constructed using supramolecular complexes formed between

![Figure 6](image-url)
polypeptides (P(H2P)n; n is rather weak. Our second study concerned supramolecular complexes of free base porphyrin arrays with electron acceptors without Lewis base ligands, in which the binding energies of supramolecular complexes between porphyrins acting as light harvesting units and electron acceptors containing Lewis base ligands are much stronger than those observed in the free base porphyrins. As such there have been only a few reports on supramolecular complexes of free base porphyrins acting as light harvesting units and electron acceptors containing Lewis base ligands due to the charge migration among porphyrins.

In the first example, metal centers were required for the construction of supramolecular complexes between porphyrins acting as light harvesting units and electron acceptors containing Lewis base ligands for coordination to the metal centers. As such there have been only a few reports on supramolecular complexes of free base porphyrin arrays and electron acceptors without Lewis base ligands, in which the binding is much stronger than that observed in the free base porphyrins. Although the lifetimes of the singlet CS states were too short to be detected, the triplet CS states derived from Li+@C60 had long lifetimes because of the spin-forbidden back electron transfer. The triplet CS lifetime becomes longer upon increasing the number of H2P due to the charge migration among porphyrins.

Acknowledgments

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References


Table III. Rate constants of intra- and inter-molecular photo-induced electron transfer (kET and ktr) and quantum yields (ΦCS) of the charge-separated state.

<table>
<thead>
<tr>
<th>Porphyrin</th>
<th>kET, M⁻¹ s⁻¹</th>
<th>ktr, M⁻¹ s⁻¹</th>
<th>τCS, µs</th>
<th>ΦCS</th>
</tr>
</thead>
<tbody>
<tr>
<td>P(H2P)₁</td>
<td>7.7 × 10⁵</td>
<td>1.7 × 10⁸</td>
<td>~0.70</td>
<td></td>
</tr>
<tr>
<td>P(H2P)₂</td>
<td>6.7 × 10⁵</td>
<td>1.8 × 10⁸</td>
<td>~0.34</td>
<td></td>
</tr>
<tr>
<td>P(H2P)₄</td>
<td>9.5 × 10⁴</td>
<td>8.0 × 10⁸</td>
<td>160</td>
<td>0.16</td>
</tr>
<tr>
<td>P(H2P)₈</td>
<td>1.5 × 10⁴</td>
<td>1.5 × 10⁹</td>
<td>210</td>
<td>0.13</td>
</tr>
</tbody>
</table>

a) Bimolecular decay in the radical ion pair.

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References