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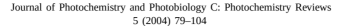
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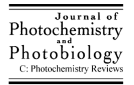
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### Review

## Intermolecular and supramolecular photoinduced electron transfer processes of fullerene–porphyrin/phthalocyanine systems

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### **Abstract**

The attainment of a better understanding of the dependence of photoinduced electron transfer reaction rates on the molecular structures of the donor and acceptor entities results in improving the capture and storage of solar energy. Here, the intermolecular and supramolecular electron transfer processes from electron donors (porphyrins (P), chlorophylls (Chl), phthalocyanines (Pc) and naphthalocyanines (Nc)) and their metal derivatives to electron acceptors (fullerenes such as  $C_{60}$  and  $C_{70}$ ) studied by nanosecond and picosecond laser flash photolysis techniques in polar and nonpolar solvents are reviewed. For intermolecular systems in polar solvents, photoinduced electron transfer takes place via the excited triplet states of  $C_{60}/C_{70}$  or via the excited triplet states of P/Pc/Nc, yielding solvated radical ions in polar solvents; thus, the back electron transfer rates are generally slow. In the case of the supramolecular dyads and triads formed by axial coordination, hydrogen bonding, crown ether complexation, or rotaxane formation, the photoinduced charge separation takes place mainly from the excited singlet state of the donor; however, the back electron transfer rates are generally quite fast. The relations between structures and photochemical reactivities of these novel supramolecular systems are discussed in relation to the efficiency of charge separation and charge recombination. © 2004 Japanese Photochemistry Association. Published by Elsevier B.V. All rights reserved.

Keywords: Porphyrins; Phthalocyanines; Fullerenes; Photoinduced electron transfer; Charge separation; Charge recombination; Self-assembly; Supramolecules; Intermolecular interactions

### **Contents**

1.	Intro	luction	80
2.	Intern	nolecular electron transfer	81
	2.1.	Fullerenes-tetraphenylporphyrins	81
	2.2.	Fullerenes-octaethylporphyrins	84
	2.3.	Fullerene-chlorophylls	87
	2.4.	Fullerenes-phthalocyanine/naphthalocyanine	88
3.	Photo	pinduced electron transfer in supramolecular fullerene-porphyrin/phthalocyanines systems.	91
	3.1.	Fullerene–porphyrin systems coordinated via axial ligation	91
	3.2.	Two-point binding supramolecular triads with electron donor	96
	3.3.	Fullerene-porphyrin coordinated systems: control over distance and orientation	98
	3.4.	Fullerene-bisporphyrin coordinated triads	99
	3.5.	Fullerene–porphyrin/phthalocyanine assembly systems	101
4.	Sumr	nary	102
Ack	nowle	dgements	102
Ref	erences		102

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### 1. Introduction

The process of photoinduced electron transfer (PET) is of great importance in chemistry and biology [1–15]. One of the most important goals of chemistry during the past century has been the construction and development of molecular and supramolecular-based artificial solar energy harvesting systems that have the ability to absorb light from the sun and convert it to useful and storable forms. One way to store solar energy is in the form of chemical energy, as plants do efficiently during photosynthesis. However, for building efficient artificial solar energy converting systems for this purpose, there are certain requirements that must be met: (i) the light must be captured by antenna molecules and/or sensitizers, leading to "excited states;" (ii) the absorption of the light must result in transfer of an electron to the acceptor entity; (iii) the electron transfer must be directional; and (iv) the lifetimes of the excited states must be long enough for electron transfer to take place. Constructing chemical systems possessing the characteristics listed above has been a very challenging goal for chemists over the past two decades.

Intermolecular PET is a simple process in which an electron is transferred from an electron-donating species (D) to an electron-accepting species (A), producing the radical cation of the donor ( $D^{\bullet+}$ ) and the radical anion of the acceptor ( $A^{\bullet-}$ ), when one of these species is photoexcited [2,15]. If these charged species are utilized as electrons and holes to drive electrical current or promote chemical reactions before back electron transfer leading to the initial states of the reactants occurs (Fig. 1), the light energy is effectively converted into electrical or chemical energy.

A critical factor in PET lies in the successful matching of D and A with suitable electrochemical and photophysical properties for the occurrence of such an exothermic ET [2,15,16,17]. Knowledge of the excited state energies of the chromophores and the redox potentials of D and A is thus an essential requirement for investigating PET processes. The majority of research on the photochemistry of porphyrins is an attempt to mimic the photosynthetic processes, in which

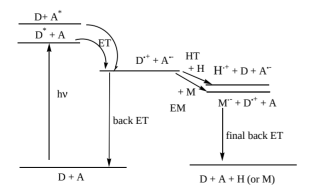


Fig. 1. Schematic energy diagrams for photoinduced ET processes in bimolecular donor–acceptor systems: HT refers to hole transfer step in the presence of hole acceptor (H) and EM refers to an electron mediation step in the presence of an electron mediator (M).

porphyrins have been widely employed as sensitizers and as electron donors [18–20]. As electron-acceptors, benzo-quinones and methyl viologens have been used to generate photocurrent and hydrogen evolution [21–23]. Covalently connected porphyrin–quinone dyads and triads have been synthesized to realize long lifetimes of the charge separated states [24–31].

Since the fullerenes were discovered and preparation methods were developed, fullerenes have been utilized as photosensitizers and electron acceptors [32,33]. Fullerenes (C<sub>60</sub>/C<sub>70</sub>) exhibit a number of characteristic electronic and photophysical properties, which make them promising candidates for the investigation of PET processes. Some of these characteristics are [32-39]: (i) fullerenes have first reduction potentials comparable to that of benzoquinone [40,41]. Since fullerenes can reversibly accept up to six electrons in electrochemical measurements, and in principle can act as electron accumulators [40,41], there are possibilities to realize a multiple photoreduction process. (ii) In terms of transient absorption spectral features, the singlet excited states of fullerenes (C<sub>60</sub> and C<sub>70</sub>) give rise to characteristic singlet-singlet absorptions in the visible and near-IR region [32,33,42-44]. Once generated, the excited singlet states (1.65–1.75 eV) are subject to a rapid and quantitative intersystem crossing process, with a lifetime of 0.9–1.3 ns, to the energetically low lying triplet excited states (1.45-1.55 eV) with lifetimes longer than 40 µs [32,33,44]. (iii) The triplet-triplet absorption spectrum of C<sub>60</sub> shows a maximum in the visible region  $(740 \,\mathrm{nm}; \,\varepsilon = 18,000 \,\mathrm{M}^{-1} \,\mathrm{cm}^{-1})$  [40]; in the case of  $C_{70}$ , the triplet-triplet absorption spectrum appears at 980 nm, with  $\varepsilon = 4000 \,\mathrm{M}^{-1} \,\mathrm{cm}^{-1}$  [45]. (iv) A more practical aspect of C<sub>60</sub> and C<sub>70</sub> concerns the optical absorption spectra of their  $\pi$ -radical anions, such as  $C_{60}^{\bullet-}$  and  $C_{70}^{\bullet-}$ , which show narrow bands in the near-IR region, around 1080 and 1380 nm, respectively, serving as diagnostic probes for their identification [32,33,45–47]. Furthermore, these isolated absorptions allow an accurate analysis of inter- and intramolecular ET dynamics of C<sub>60</sub> and C<sub>70</sub>, even in the presence of porphyrins and phthalocyanines, which have wide absorptions in the visible region. For this purpose, it is very important to develop techniques to measure the transient absorption spectra in the near-IR region [50,51]. (v) Fullerene-based electron donor-acceptor dyads exhibit relatively rapid photoinduced charge-separation (CS) and relatively slow charge-recombination (CR) due to the low reorganization energy of fullerenes [48–52]. Achieving a long-lived CS state after photoexcitation is the key to realizing artificial photosynthesis in supramolecular systems.

Porphyrins form a ubiquitous class of naturally occurring molecules. The UV-Vis absorption spectrum of the highly conjugated porphyrin macrocycle exhibits an intense feature (extinction coefficient > 200,000) at about 400 nm (the Soret band) followed by several weaker absorptions (Q bands) at higher wavelengths (from 450 to 750 nm), which are changed by the peripheral substituents on the porphyrin

ring and insertion of metal atoms into the center of the porphyrin ring. The extensively conjugated  $\pi$ -systems of porphyrins increase their electron-donor abilities, so that they are suitable for efficient ET in the ground and excited states. The electronic excited states of porphyrins survive long enough in the singlet and triplet states to provide a high probability to interact with molecules before deactivation [53–58]. Porphyrins are involved in a wide variety of important biological processes, ranging from oxygen transport to photosynthesis [4,53–58]. The role of porphyrins in photosynthetic mechanisms indicates a good capability to mediate visible photon-electron conversion processes. Porphyrins and related macrocycles such as phthalocyanines provide an extremely versatile synthetic base for a variety of materials for applications in many disciplines of chemistry and physics, such as opto-electronics, electrochemistry, catalysis, data storage, and solar cells [4,10,58–60]. Porphyrins and metalloporphyrins have also been examined for a variety of applications as sensors, which clearly represent an important class of chemo-responsive materials [10,22,61]. The stability of mono- and di-cation porphyrin  $\pi$ -radicals makes these systems especially interesting for photoionization processes, closely related to the so-called special pair reaction center of photosynthesis [4,9].

Fullerene–porphyrin mixed systems have recently become an active area of research for the generation of photocurrent [62–64]. To reveal the elemental processes, including electron transfer and electron-mediation process in addition to energy transfer (EN), there are several studies available in the literature [65–71]. Supramolecular systems composed of functionalized fullerenes that are coordinated to the central metal of the porphyrin have been studied to mimic the photosynthetic system [72–81]. The covalently connected fullerene–porphyrin dyads and triads were extensively investigated with the purpose of generating photocurrent, in addition to their unique photophysical and photochemical properties [48–52].

In the first part of the present review, we focus on the intermolecular ET between fullerenes ( $C_{60}$  and  $C_{70}$ ) with porphyrins, chlorophylls, phthalocyanines and naphthalocyanines to reveal the fundamental photochemical features of these systems. In the second part, we summarize the photochemical behavior of supramolecular assemblies, in which functionalized fullerenes are noncovalently interacting with porphyrins and phthalocyanines.

### 2. Intermolecular electron transfer

The simplest way to prepare the intermolecular system is by mixing electron acceptors (fullerenes) with electron donors (porphyrins, chlorophylls, phthalocyanines, and naphthalocyanines) in a suitable solvent. The electron transfer events can be monitored by observing the radical ions by means of nanosecond transient absorption spectra in the visible and near-IR regions, with which the ET mechanism

and ET kinetics can be characterized. We have organized this section into four parts: The first part deals with the ET processes of  $C_{60}$  and  $C_{70}$  with tetraphenylporphyrin ( $H_2$ TPP) bearing different substituents on the phenyl rings. The second part covers the ET processes of  $C_{60}$  and  $C_{70}$  with metal octaethylporphyrins (MOEP, where  $M=H_2$ , Pd, Ni, Co, V=O, Mg, Zn and Cu) to probe the effect of metal ions in the porphyrin cavity. The third part deals with the ET processes of  $C_{60}$  and  $C_{70}$  with chlorophylls (Chls) to reveal the role in natural systems. Finally, the fourth part deals with the ET processes of  $C_{60}$  and  $C_{70}$  with phthalocyanines (Pc) and naphthalocyanines (Nc) to probe structural effects of electron donors.

### 2.1. Fullerenes-tetraphenylporphyrins

Recently, we studied the electron transfer process of  $C_{60}$  with tetraphenylporphyrin (H<sub>2</sub>TPP) bearing different substituents on the phenyl rings to probe the substituent effects on the rates of the electron transfer process. It was reported that the photophysical and photochemical properties of porphyrins are affected by the substituents [82–85].

We employed free-base tetraphenylporphyrin ( $H_2$ TPP), tetra(p-hydroxyphenyl)-porphyrin ( $H_2$ THPP) and tetra(p-aminophenyl)porphyrin ( $H_2$ TAPP) and tetra(p-methoxyphenyl) porphyrin ( $H_2$ TMPP) as electron donors (Fig. 2) with  $C_{60}$  as an electron acceptor. This study was carried out in benzonitrile (BN) via triplet states of porphyrins ( ${}^3H_2$ TPPs\*) by observing the transient spectra in the wide spectral range from 400 to 1500 nm.

The absorption spectra of H<sub>2</sub>TPP, H<sub>2</sub>THPP and H<sub>2</sub>TAPP are shown in Fig. 3. The absorption bands of the porphyrins with electron-donating substituents are shifted to longer wavelength compared with those of H<sub>2</sub>TPP (Fig. 3). Such red shifts might originate from the narrowing of the band gap energy, which is caused by an increase in the HOMO energy level with electron-donating substituents, as can be interpreted according to Gouterman's four orbital model [86].

Absorption spectra of  $C_{60}$  and  $C_{70}$  are shown in Fig. 4; the absorbance in the visible region of  $C_{60}$  is weaker than those of  $H_2$ TPP derivatives. The absorption spectra of the mixture of either of  $H_2$ TPP,  $H_2$ THPP, or  $H_2$ TAPP with  $C_{60}$  are the same as the summation of the spectra of the corresponding components, suggesting that the interaction between  $C_{60}$  and the substituted porphyrins in the ground state is weak.

By photoexcitation of  $H_2TPP$  (0.1 mM) in deaerated BN using a 550 nm laser, the transient absorption spectrum obtained immediately after the laser pulse exhibited absorption bands at 450 and 780 nm, which are assigned to the triplet state of  $H_2TPP$  ( ${}^3H_2TPP^*$ ) [69–71]. In the presence of  $C_{60}$ , the generation of  $C_{60}$  was observed by a build-up of the absorption at 1080 nm at 10  $\mu$ s [32,33,45–47] that parallels a concomitant decay of  ${}^3H_2TPP^*$  (Fig. 5a). It was difficult to observe clearly the  $H_2TPP^{\bullet+}$  at 650 nm, because of the overlap with the depletion and emission of  $H_2TPP$ . In

Fig. 2. Structures of C<sub>60</sub>, C<sub>70</sub> and meso-tetraphenylporphyrins.

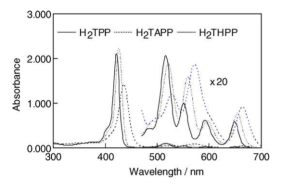


Fig. 3. Steady state absorption spectra of  $H_2$ TPP,  $H_2$ TAPP, and  $H_2$ THPP in BN; concentration = 0.007 mM.

the case of  $H_2TAPP$  as an electron donor to  $C_{60}$ , the transient spectrum (Fig. 5b) showed the bands at 460, 630, and 740 nm immediately after the laser exposure. These three bands are clearly assigned to  ${}^3H_2TAPP^*$ . With the decay of  ${}^3H_2TAPP^*$ , the rise of  $C_{60}^{\bullet-}$  was observed at 1080 nm. Interestingly, the broad absorption bands in the 600–1400 nm region with maxima at 580, 780, and 1200 nm, which are attributed to  $H_2TAPP^{\bullet+}$ , were observed in the spectrum at 10  $\mu$ s. Similarly, with the decays of  ${}^3H_2THPP^*$  at 460, 620, and 680 nm, the rise of  $H_2TPP^{\bullet+}$ , showing absorptions at 630, 680 and 950 nm, was observed in addition to  $C_{60}^{\bullet-}$  at 1080 nm (Fig. 5c).

Furthermore, the contribution of the triplet states of porphyrins to the ET process was confirmed by the  $O_2$  effect.

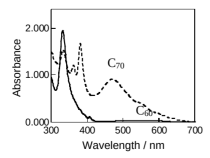


Fig. 4. Steady state absorption spectra of  $C_{60}$  and  $C_{70}$ ; concentration = 0.1 mM.

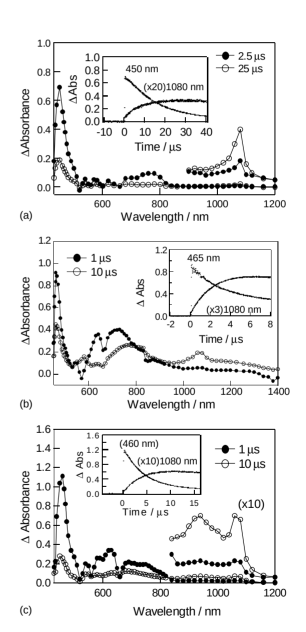


Fig. 5. Transient absorption spectra obtained by 550 nm laser photolysis of (a)  $\rm H_2TPP$  (0.1 mM), (b)  $\rm H_2TAPP$  (0.1 mM), and (c)  $\rm H_2THPP$  in the presence of  $\rm C_{60}$  (0.1 mM) in Ar-saturated BN.

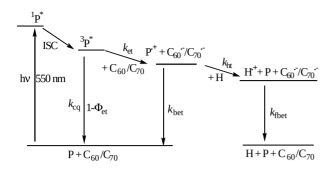


Fig. 6. Energy diagram for electron transfer by photoexcitation of P, which represents porphyrins, phthalocyanines, and chlorophylls, in the presence of  $C_{60}/C_{70}$  in polar solvents.

On addition of  $O_2$ , the decay of  ${}^3H_2TPP^*$  was accelerated owing to energy transfer to  $O_2$ ; consequently, the formation of  $C_{60}^{\bullet-}$  and  $H_2TPPs^{\bullet+}$  was suppressed. These observations suggest that the ET process takes place from  ${}^3H_2TPP^*$  to  $C_{60}$  (Fig. 6).

A more detailed picture of the kinetic event was observed in the time profiles, from which the rate constants of the bimolecular quenching  $(k_q)$  of  ${}^3H_2TPP^*$  were evaluated by monitoring the first-order decays of  ${}^3H_2TPPs^*$  as a function of  $C_{60}$  concentrations under the condition of  $[{}^3H_2TPPs^*] \ll [C_{60}]$ . The first-order rate constant for the decays of  ${}^3H_2TPPs^*$  in the presence of  $C_{60}$  is referred to as  $k_{1st}$  in Eq. (1).

$$k_{1st} = k_0 + k_q[C_{60}] \tag{1}$$

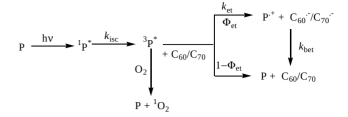
where  $k_0$  is referred to as a rate constant for the decay of  ${}^3\mathrm{H}_2\mathrm{TPPs}^*$  in the absence of  $\mathrm{C}_{60}$ . The linear dependence of the observed  $k_{1\mathrm{st}}$  values on  $[\mathrm{C}_{60}]$  gives the rate constant for bimolecular quenching  $(k_{\mathrm{q}})$ , as summarized in Table 1. The  $k_{\mathrm{q}}$  values of  ${}^3\mathrm{H}_2\mathrm{TPPs}^*-\mathrm{C}_{60}$  are in the range of  $(1.1-1.3)\times 10^9\,\mathrm{M}^{-1}\,\mathrm{s}^{-1}$ , although larger  $k_{\mathrm{q}}$  values were expected for substituted porphyrins, because of the lower oxidation potential  $(E_{\mathrm{ox}})$  values of substituted porphyrins,  $\mathrm{H}_2\mathrm{TMPP}$  (0.98 V),  $\mathrm{H}_2\mathrm{THPP}$  (0.75 V), and  $\mathrm{H}_2\mathrm{TAPP}$  (0.48 V) compared to  $\mathrm{H}_2\mathrm{TPP}$  (1.05 V) versus  $\mathrm{Fc/Fc}^+$  [87–89].

For  ${}^3H_2TPP^*$ , the absorption of  $C_{60}^{\bullet-}$  was not overlapped with those of the radical cation of  $H_2TPP^{\bullet+}$ ; therefore we can evaluate the maximal concentration of  $C_{60}^{\bullet-}$  from the reported molar extinction coefficient  $(14,000 \, M^{-1} \, cm^{-1})$  at

Table 1 Quenching rate constants  $(k_{\rm q})$  of the excited triplet states of substituted tetraphenylporphyrins ( $^3{\rm TPPs}^*$ ) by fullerene ( $C_{60}$ ) and back electron transfer rate constants  $(k_{\rm bet})$  between  $C_{60}^{\bullet-}$  and  $P^{\bullet+}$  in Ar-saturated benzonitrile (BN)

Systems	$k_{\rm q} \ ({\rm M}^{-1}{\rm s}^{-1}) \ (\times 10^9)$	$k_{\text{bet}} \ (M^{-1} \ s^{-1}) \ (\times 10^9)$
$^{3}\text{H}_{2}\text{TAPP*-C}_{60}$	1.1	2.4
$^3$ H <sub>2</sub> THPP* $-$ C <sub>60</sub>	1.4	3.5
$^{3}H_{2}TMPP^{*}-C_{60}$	1.3	5.5
$^{3}\text{H}_{2}\text{TPP*}-\text{C}_{60}{}^{a}$	1.1	4.9

<sup>&</sup>lt;sup>a</sup>  $\Phi_{\text{et}} = 0.26$  and  $k_{\text{et}} = 2.9 \times 10^8 \,\text{M}^{-1} \,\text{s}^{-1}$  [69].



Scheme 1. Routes for ET process occurring by the photoexcitation of electron donors (P) in the presence of fullerenes ( $C_{60}/C_{70}$ ) in BN; P is an abbreviation for porphyrins, chlorophylls, phthalocyanines, and naphthalocyanines.

1080 nm) [45–47,69–71]. In contrast, the initial concentration of  ${}^{3}\text{H}_{2}\text{TPP}^{*}$  was also evaluated from the molar extinction coefficient (20,000 M $^{-1}$  cm $^{-1}$  at 450 nm) [90]. Thus, the efficiency of ET via  ${}^{3}\text{H}_{2}\text{TPP}^{*}$  can be evaluated from the ratio of  $[C_{60}^{\bullet-}]_{\text{max}}/[{}^{3}\text{H}_{2}\text{TPP}^{*}]_{\text{initial}}$ , which usually saturates at appropriately high concentrations of  $[C_{60}]$ . The saturated ratio can be made equal to the quantum yield ( $\Phi_{\text{et}}$ ); for  ${}^{3}\text{H}_{2}\text{TPP}^{*}$ – $C_{60}$ , the  $\Phi_{\text{et}}$  value was evaluated to be 0.26 [71]. The rate constants for electron transfer ( $k_{\text{et}}$ ) can be evaluated with Eq. (2) [91–93]:

$$k_{\rm et} = \Phi_{\rm et} k_q \tag{2}$$

Since the  $\Phi_{\rm et}$  values are less than unity, there may be bimolecular deactivation processes of  ${}^3{\rm H}_2{\rm TPP}^*$  other than the ET process. As such a bimolecular deactivation process of  ${}^3{\rm H}_2{\rm TPP}^*$ , collisional quenching can be considered, as shown in Scheme 1, since energy transfer processes were not observed.

For  ${}^3H_2TAPP^*-C_{60}$ ,  ${}^3H_2THPP^*-C_{60}$ , and  ${}^3H_2TMPP^*-C_{60}$ , the molar extinction coefficients were not exactly evaluated; thus, the values of  $\Phi_{\rm et}$  and  $k_{\rm q}$  have not yet been obtained. Thus, the substituent effect on values of  $\Phi_{\rm et}$  and  $k_{\rm q}$  is not clear at this moment.

In the long time scale measurements, it was clearly observed that  $C_{60}^{\bullet-}$  begins to decay slowly after reaching the maximal absorbance. The decay time profile was fitted with second-order kinetics, suggesting that a bimolecular back ET process ( $k_{\text{bet}}$ ) from  $C_{60}^{\bullet-}$  to TAPP<sup>•+</sup> takes place after these radical ions are solvated separately into free radical ions in a polar solvent such as BN.

$$\frac{1}{\Delta A_t} = \frac{1}{\Delta A_0} + \frac{k_{\text{bet}}}{\varepsilon} t \tag{3}$$

From the slopes of the line of the second-order plot of  $1/\Delta A_t$  versus time (t), the ratio of the back ET rate constant  $(k_{\rm bet})$  to the molar extinction coefficient of the radical ions  $(\varepsilon)$  can be obtained. On employing the reported extinction coefficient  $(\varepsilon_{\rm A})$  of  $C_{60}^{\bullet-}$ , the  $k_{\rm bet}$  values were evaluated, as listed in Table 1. The  $k_{\rm bet}$  values of the substituted systems  $C_{60}^{\bullet-}$ – $H_2$ TAPP $^{\bullet+}$ ,  $C_{60}^{\bullet-}$ – $H_2$ THPP $^{\bullet+}$ , and  $C_{60}^{\bullet-}$ – $H_2$ TMPP $^{\bullet+}$  are much smaller than that of  $C_{60}^{\bullet-}$ – $H_2$ TPP $^{\bullet+}$ , suggesting the delocalization of the hole in the substituted porphyrins,

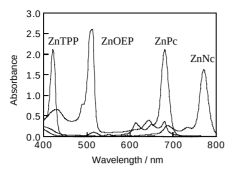


Fig. 7. Steady state absorption spectra of zinc tetraphenylporphyrin (ZnTPP), zinc octaethylporphyrin (ZnOEP), zinc phthalocyanine (ZnPc) and zinc naphthalocyanines (ZnNc) in BN: concentration = 0.007 mM.

which is supported by the appearance of the longer wavelength absorption band of  $H_2TAPP^{\bullet+}$ ,  $H_2THPP^{\bullet+}$ , and  $H_2TMPP^{\bullet+}$ . In general, the  $k_{bet}$  values seem to be close to the diffusion-controlled limit ( $k_{diff}$ ) in BN, which means that  $C_{60}^{\bullet-}$  and  $H_2TPP^{\bullet+}$  are long lived, although the lifetimes were dependent on their concentration [89]. Since the concentrations of  $C_{60}^{\bullet-}$  and  $H_2TPP^{\bullet+}$  are considerably lower than that of the reactant [ $C_{60}$ ], the observed decay rates of the backward process are far smaller than that of the forward process, even though  $k_{bet} \gg k_{et}$ .

The PET process was investigated between MTPP (M =  $H_2$ , Zn, Cu) and fullerenes ( $C_{60}/C_{70}$ ) in polar solvents by applying the 532 nm nanosecond laser photolysis method [69,70]. As shown in Fig. 7, the absorption peaks of ZnTPP are almost the same as those of  $H_2$ TPP. The ET process was followed via both  $^3$ MTPP\* and  $^3C_{60}^*/^3C_{70}^*$  by controlling the excitation molecules by their absorbance at 532 nm.

By employing the laser light at 532 nm, selective excitation of  $C_{70}$  was possible, since the absorption intensity at 532 nm of  $C_{70}$  is much higher than those of MTPP. The transient absorption of  ${}^3C_{70}^*$  appeared at 980 nm immediately after the laser light excitation of  $C_{70}$  in the presence of excess MTPP. With concomitant decay of  ${}^3C_{70}^*$ , the transient absorption band of  $C_{70}^{\bullet-}$  appeared at 1380 nm in the near-IR region giving evidence of ET from ZnTPP to  ${}^3C_{70}^*$  in BN. Similarly, ET from ZnTPP to  ${}^3C_{60}^*$  was possibly investigated at high concentration of  ${}^3C_{60}^*$  to permit the selec-

Table 2 Quenching rate constants  $(k_q)$ , electron transfer quantum yield  $(\Phi_{et})$  electron transfer rate constants  $(k_{et})$  of fullerene–MTPP systems in BN

Systems	$k_{\rm q}  ({ m M}^{-1}  { m s}^{-1}) \ ( imes 10^9)$	$\Phi_{ m et}$	$k_{\rm et}  ({\rm M}^{-1}  {\rm s}^{-1})$ (×10 <sup>8</sup> )
<sup>3</sup> C <sub>70</sub> *–ZnTPP	2.2	0.35	7.7
$^{3}\text{C}_{60}^{*}Z$ –ZnTPP	4.3	0.26	1.1
<sup>3</sup> C <sub>60</sub> *–CuTPP	2.1	0.13	2.7
$^3$ ZnTPP* $-$ C $_{70}$	4.7	0.15	7.0
$^3$ ZnTPP* $-$ C $_{60}$	4.0	0.12	4.8

tive excitation of  ${}^3C_{60}^*$ . Fig. 8 shows the schematic energy diagram of the ET process from the ground state of ZnTPP to  ${}^3C_{60}^*/{}^3C_{70}^*$ .

The quantum yields  $(\Phi_{\rm et})$  via  ${}^3{\rm C}_{60}^*$  and  ${}^3{\rm C}_{70}^*$  can be evaluated from the ratio of  $[{\rm C}_{60}^{\bullet-}]_{\rm max}/[{}^3{\rm C}_{60}^*]_{\rm initial}$  and  $[{\rm C}_{70}^{\bullet-}]_{\rm max}/[{}^3{\rm C}_{70}^*]_{\rm initial}$  in the range of 0.2–0.4, which also suggests the presence of the collisional quenching of  ${}^3{\rm C}_{60}^*$  and  ${}^3{\rm C}_{70}^*$  without ET (Scheme 2). The  $k_{\rm et}$  values for  ${}^3{\rm C}_{60}^*$  and  ${}^3{\rm C}_{70}^*$  from ZnTPP and CuTPP were evaluated from  $k_{\rm q}$   $\Phi_{\rm et}$ , as listed in Table 2.

In concentrated solutions of ZnTPP in which it was photoexcited selectively by 532 nm laser light, ET takes place from  ${}^3$ ZnTPP\* to the ground state of  $C_{60}$  and  $C_{70}$ , producing  $C_{60}^{\bullet-}$  and  $C_{70}^{\bullet-}$  (Scheme 1). The electron transfer rate constants and efficiencies were evaluated, listed in Table 2. Although there were slight differences in the  $k_q$ ,  $\Phi_{et}$ , and  $k_{et}$  values between  $C_{60}$  and  $C_{70}$  as well as between ZnTPP and  $H_2$ TPP, it can be considered that they are substantially similar values. This implies that the rigorous removal of  $C_{70}$  from mixtures of  $C_{60}$  and  $C_{70}$  is not necessary.

### 2.2. Fullerenes-octaethylporphyrins [70,93]

This section covers the ET process of metal octaethyl-porphyrins (MOEP, where  $M=H_2$ , Pd, Ni, Co, V=O, Mg, Zn and Cu) with fullerenes ( $C_{60}/C_{70}$ ) to reveal the effect of the metal ions in the porphyrin cavity (Fig. 9). Steady-state absorption bands of ZnOEP appear at longer wavelengths than those of ZnTPP, as shown in Fig. 7. The insertion of metal atoms into OEP usually strongly changes the visible absorption spectra; i.e., the Q-bands of  $H_2OEP$  (498,

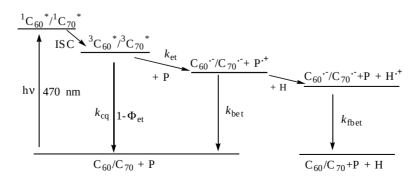


Fig. 8. Energy diagram for electron transfer by photoexcitation of fullerenes  $(C_{60}/C_{70})$  in the presence of porphyrins (P) in polar solvents.

$$C_{60}/C_{70} \xrightarrow{hv} {}^{1}C_{60}^{*/1}C_{70}^{*} \xrightarrow{k_{isc}} {}^{3}C_{60}^{*/3}C_{70}^{*} \xrightarrow{+P} \xrightarrow{k_{et}} C_{60}^{*/7}C_{70}^{*} + P^{+}$$

$$O_{2} \downarrow \qquad \qquad \downarrow k_{bet}$$

$$C_{60}/C_{70} + {}^{1}O_{2}$$

Scheme 2. Routes for the ET process occurring by the photoexcitation of fullerenes ( $C_{60}/C_{70}$ ) in the presence of porphyrins (P) in BN.

532, 567, and 521 nm), PdOEP (512, 546 nm), NiOEP (517, 552 nm), CuOEP (526, 562 nm), (V=O)OEP (534, 572 nm) and MgOEP (544 and 579 nm).

Selective excitation of C<sub>70</sub> is possible even in the presence of excess MOEP by 470 nm laser light irradiation. In the transient absorption spectra obtained immediately after the laser excitation, the absorption band of  ${}^{3}C_{70}^{*}$  at 980 nm was solely observed; with concomitant decay of  ${}^{3}C_{70}^{*}$ , the absorption of  $C_{70}^{\bullet-}$  appeared at 1380 nm with the absorption of MOEP<sup>•+</sup> at 650 nm, indicating that the ET process takes place in the same manner, as shown in Fig. 8.

The ET quantum yields  $(\Phi_{et})$  of  $C_{70}^{\bullet-}$  formation via  ${}^{3}C_{70}^{*}$  were evaluated from  $[C_{70}^{\bullet -}]_{max}/[{}^{3}C_{70}^{*}]_{initial}$  at appropriately high concentrations of [MOEP]. The  $\Phi_{\rm et}$  values via  ${}^{3}\text{C}_{70}^{*}$  varied with the central metal according to the following order; PdOEP > MgOEP > ZnOEP > (V=O)OEP > CoOEP > NiOEP > CuOEP. The change in the donor abilities of the MOEPs may be explained mainly by their  $E_{ox}$ values. The observed  $\Phi_{\rm et}$  values are less than unity, suggesting that there are some deactivation routes (e.g., collisional quenching and/or an encounter complex). The possibility of EN from  ${}^{3}\text{C}_{70}^{*}$  to MOEP in BN solution is quite low, because the rise of <sup>3</sup>MOEP\* was not observed. Thus, the deactivation process may be attributed to collisional quenching.

In Table 3, it is shown that the  $\Phi_{\rm et}$  values gradually increase with decreasing  $E_{ox}(D/D^+)$ , except for (V=O)OEP. The free energy changes ( $\Delta G_{\text{et}}^0$ ) for ET from <sup>3</sup>P to C<sub>60</sub> were calculated from the Rehm-Weller relation [16,17].

$$\Delta G_{\text{ef}}^0 = E_{\text{ox}}(D/D^{\bullet +}) - E_{\text{red}}(A^{\bullet -}/A') - E_{\text{T}} - E_{\text{c}}$$
 (4)

where  $E_{\rm ox}({\rm D/D}^{\bullet+})$ ,  $E_{\rm red}({\rm A}^{\bullet-}/{\rm A})$ ,  $E_{\rm c}$  and  $E_{\rm T}$  refer to the oxidation potential of the donor, the reduction potential of the acceptor  $(C_{60}/C_{70})$ , the Coulomb term, and the triplet energy of the excited species, respectively. It was also observed that the  $k_{\rm et}$  values increase with decreasing  $\Delta G_{\rm et}^0$  values along the curve calculated by the semiempirical Rehm-Weller plot [16,17]. For systems with very negative  $\Delta G_{\rm et}^0$  values, the  $k_{\rm et}$ values are close to  $k_{\text{diff}}$  in BN [89]. The  $k_{\text{bet}}$  values in BN listed in Table 3 are also close to  $k_{\text{diff}}$ , because the free energy change ( $\Delta G_{\text{bet}}^0$ ) for back ET from  $C_{70}^{\bullet-}$  to MOEP $^{\bullet+}$ , which can be calculated from Eq. (5), are all very negative.

$$\Delta G_{\text{bet}}^0 = E_{\text{ox}}(D/D^{\bullet+}) - E_{\text{red}}(A^{\bullet-}/A) - E_{\text{c}}$$
 (5)

By laser excitation of C<sub>70</sub> in the presence of MOEP in toluene, no ET process was observed. The EN process from <sup>3</sup>C<sub>70</sub>\* to MOEP in toluene was also not observed; this observation is reasonable, because  $E_{\rm T}$  ( $^3{\rm C}_{70}^*=1.54\,{\rm eV}$ ) is slightly lower than  $E_{\rm T}$  ( $^3$ MOEP\* = 1.60–1.90 eV). Thus, the formation of the triplet exciplex  ${}^3[C_{70}^{(\delta-)}\cdots MOEP^{(\delta+)}]^*$ would be expected to be dominant in nonpolar solvents. The formation of such triplet exciplexes in nonpolar solvents has been reported during the quenching of <sup>3</sup>MTPP by various quinones in toluene [94]. Thus, the formation of  ${}^{3}[C_{70}^{(\delta-)}\cdots MOEP^{(\delta+)}]^{*}$  could provide a possible explanation for the near diffusion-controlled triplet quenching rate constant of  ${}^{3}C_{70}^{*}$  in the presence of MOEP in toluene.

Kinetic analysis of the <sup>3</sup>C<sub>70</sub>\*-MgOEP system in toluene–BN mixtures afforded valuable information about the dissociation of  ${}^3[C_{70}^{(\delta-)}\cdots \text{MOEP}^{(\delta+)}]^*$  into the

Metal octaethylporphyrins (MOEP) ( M = Pd, Mg, Zn, Co, Ni, Cu and (V=O) 6-yl-methylene)hydrazine (DTQH)

N,N-diphenyl-N-(1,2,3,4-tetrahydro-quinoline-

Fig. 9. Structure of metal octaethylporphyrins (MOEP) and a hole-transfer reagent (DTQH).

Table 3 Oxidation potentials  $(E_{\rm ox})$ , and free energy changes  $(\Delta G_{\rm et}^0)$ , and kinetic parameters  $(k_{\rm q}, \Phi_{\rm et} \text{ and } k_{\rm et})$  for the ET process from MOEP via  ${}^3{\rm C}_{70}{}^*$  in BN;  $k_{\rm bet}$  between  ${\rm C}_{70}{}^{\bullet-}$  and MOEP $^{\bullet+}$ 

System	$E_{\rm ox}$ (V)	$\Delta G_{\mathrm{et}}^{0} \; (\mathrm{kJ}  \mathrm{mol}^{-1})$	$k_{\rm q} \ ({\rm M}^{-1} \ {\rm s}^{-1})$	$\overline{arPhi_{ ext{et}}}$	$k_{\rm et} \ ({\rm M}^{-1} \ {\rm s}^{-1})$	$k_{\text{bet}}  (M^{-1}  \text{s}^{-1})$
<sup>3</sup> C <sub>70</sub> *–PdOEP	0.44	-66.5	$2.2 \times 10^{9}$	0.74	$1.6 \times 10^{9}$	$3.2 \times 10^{9}$
<sup>3</sup> C <sub>70</sub> *–MgOEP	0.53	-57.7	$2.4 \times 10^{9}$	0.52	$1.2 \times 10^{9}$	$4.7 \times 10^9$
<sup>3</sup> C <sub>70</sub> *–ZnOEP	0.63	-48.1	$2.9 \times 10^{9}$	0.40	$1.1 \times 10^{9}$	$9.0 \times 10^{9}$
<sup>3</sup> C <sub>70</sub> *–NiOEP	0.64	-47.3	$2.7 \times 10^{9}$	0.32	$8.7 \times 10^{8}$	$6.5 \times 10^9$
<sup>3</sup> C <sub>70</sub> *–CoOEP	0.68	-43.5	$2.2 \times 10^{9}$	0.39	$8.5 \times 10^{8}$	$4.0 \times 10^{9}$
<sup>3</sup> C <sub>70</sub> *–CuOEP	0.85	-27.2	$2.0 \times 10^{9}$	0.21	$4.3 \times 10^{8}$	$8.0 \times 10^{9}$
${}^{3}\text{C}_{70}^{*}$ -(V=O)OEP	0.96	-17.2	$1.8 \times 10^{9}$	0.40	$7.2 \times 10^{8}$	$4.6 \times 10^{9}$
${}^{3}\text{C}_{60}^{*}Z\text{-NiOEP}$	0.64	-43.5	$3.5 \times 10^{9}$	0.11	$3.9 \times 10^{8}$	$1.2 \times 10^{9}$
${}^{3}\text{C}_{60}^{*}Z\text{CoOEP}$	0.68	-27.2	$3.3 \times 10^{9}$	0.11	$3.6 \times 10^{8}$	$7.8 \times 10^{9}$
$^{3}\text{C}_{60}^{*}Z$ –CuOEP	0.85	-18.8	$2.6 \times 10^9$	0.06	$1.6 \times 10^{8}$	$9.7 \times 10^9$

Table 4 Kinetic parameters  $(k_{\rm q}, \Phi_{\rm et}, \text{ and } k_{\rm et})$  for the ET process via  ${}^3{\rm C}_{70}{}^*$  in the presence of (V=O)OEP and  $k_{\rm bet}$  between  ${\rm C}_{70}{}^{\bullet-}$  and (V=O)OEP $^{\bullet+}$  in Ar-saturated BN:toluene (Tol) mixtures

Solvents (BN:Tol)	$k_{\rm q}~({ m M}^{-1}~{ m s}^{-1})$	$\Phi_{ m et}$	$k_{\rm et} \ ({ m M}^{-1} \ { m s}^{-1})$	$k_{\text{bet}} \ (\mathbf{M}^{-1} \ \mathbf{s}^{-1})$
25:75	$3.3 \times 10^{9}$	_	_	_
50:50	$3.3 \times 10^{9}$	0.25	$8.3 \times 10^{8}$	$9.7 \times 10^{9}$
60:40	$3.3 \times 10^{9}$	0.30	$9.8 \times 10^{8}$	$7.4 \times 10^{9}$
75:25	$3.0 \times 10^{9}$	0.35	$1.1 \times 10^{9}$	$5.6 \times 10^{9}$
87:13	$2.8 \times 10^{9}$	0.39	$1.1 \times 10^{9}$	$5.3 \times 10^{9}$
100:0	$2.6 \times 10^{9}$	0.40	$1.1 \times 10^{9}$	$4.5 \times 10^9$

solvent-separated ion-pair (SSIP) or into free radical ions in solution. In the region of toluene-rich content (toluene > 75%), the rapid decay of the transient absorption band of  ${}^3C_{70}^*$  was observed with formation of  $C_{70}^{\bullet-}$ , similar to that of 100% toluene solution. Thus, it is assumed that less polar solvents retard the dissociation of the triplet exciplex into SSIPs or into free radical ions in solution. In the region BN > 25%, the dissociation of the triplet exciplex was confirmed by observing the absorption bands of  $C_{60}^{\bullet-}/C_{70}^{\bullet-}$  and MOEP $^{\bullet+}$ . This is reasonably interpreted by the stabilization of the SSIP and free radical ions in a polar medium. In polar solvents, on assuming that lifetimes of  ${}^3[C_{70}^{(\delta-)}\cdots \text{MOEP}^{(\delta+)}]^*$  are very short, the  $\Phi_{\text{et}}$  and  $k_{\text{et}}$  values can be evaluated in a similar manner to those of BN, as listed in Table 4. In both BN and toluene–BN, the decay of  $C_{60}^{\bullet-}/C_{70}^{\bullet-}$  and MOEP $^{\bullet+}$  was fitted with second-order

kinetics, suggesting that the radical ions are present as free ion radicals or SSIP. The evaluated  $k_{\text{bet}}$  values (Table 4) seem to increase slightly with increasing toluene fraction. This finding suggests that the fraction of SSIP increases with toluene fraction, resulting in the increase of  $k_{\text{bet}}$  values.

By employing laser light at 560 nm, selective excitation of MOEP was possible even in the presence of  $C_{60}$  and  $C_{70}$ ; thus, the ET process via  $^3\text{MgOEP}^*$  was confirmed by observing the decay of the absorption bands of  $^3\text{MOEP}^*$  at 440 nm and the concomitant rise of  $C_{60}^{\bullet-}$  at 1080 nm and  $C_{70}^{\bullet-}$  at 1380 nm. A possibility of the ET process via  $^1\text{MOEP}^*$  is excluded due to the slow rise of  $C_{60}^{\bullet-}$  and  $C_{70}^{\bullet-}$ . The kinetic parameters for the  $^3\text{MOEP}^*-C_{60}$  systems in BN are listed in Table 5. The  $k_q$  values via  $^3\text{MOEP}^*$  are almost the same as those via  $^3C_{70}^*/^3C_{60}^*$ , which is reasonable on the basis of their similar  $\Delta G_{\text{et}}^0$  values. It is remarkable that the  $\Phi_{\text{et}}$  values via  $^3\text{MOEP}^*-C_{70}$  systems seem to be higher than those of the  $^3\text{MOEP}^*-C_{60}$  systems; the difference can be explained by the difference in the  $E_{\text{red}}$  values between  $C_{60}$  (-0.51 V versus SCE in BN) and  $C_{70}$  (-0.43 V versus SCE in BN).

The transient absorption spectrum observed after laser excitation of MgOEP in the presence of C<sub>70</sub> and DTQH, which is well known as a hole shifter, confirmed the hole shift process from MgOEP<sup>•+</sup> to DTQH, generating DTQH<sup>•+</sup>. Fig. 10 shows the transient absorption spectra observed by the selective excitation of MgOEP in the presence of C<sub>70</sub> and DTQH in BN. At 0.5 μs, the sharp band at 440 nm of <sup>3</sup>MOEP\* was observed, showing rapid decay

Table 5 Kinetic parameters ( $k_q$ ,  $\Phi_{et}$ , and  $k_{et}$ ) for ET from  $^3\text{MOEP}^*$  to  $C_{60}/C_{70}$  in Ar-saturated BN;  $k_{bet}$  between  $C_{60}^{\bullet-}/C_{70}^{\bullet-}$  and  $\text{MOEP}^{\bullet+}/C_{70}^{\bullet-}$  and  $\text{MOEP}^{\bullet-}/C_{70}^{\bullet-}/C_{70}^{\bullet-}$  and  $\text{MOEP}^{\bullet-}/C_{70}^{\bullet-}/C_{70}^{\bullet-}$  and  $\text{MOEP}^{\bullet-}/C_{70}^{\bullet-}/C_{70}^{\bullet-}$  and  $\text{MOEP}^{\bullet-}/C_{70}^{\bullet-}/C_{70}^{\bullet-}$  and  $\text{MOEP}^{\bullet-}/C_{70}^{\bullet-}/C_{70}^{\bullet-}$  and  $\text{MOEP}^{\bullet-}/C_{70}^{\bullet-}/C_{70}^{\bullet-}/C_{70}^{\bullet-}$ 

System	$k_{\rm q} \ ({\rm M}^{-1} \ {\rm s}^{-1})$	$\overline{\Phi_{ m et}}$	$k_{\rm et} \ ({\rm M}^{-1} \ {\rm s}^{-1})$	$k_{\text{bet}} \ (M^{-1} \ s^{-1})$
3PdOEP*-C <sub>60</sub>	$3.2 \times 10^{9}$	0.47	$1.5 \times 10^9$	$7.3 \times 10^9$
$^{3}$ ZnOEP* $-$ C <sub>60</sub>	$3.7 \times 10^9$	0.28	$1.0 \times 10^{9}$	$8.4 \times 10^{9}$
$^{3}$ MgOEP*- $C_{60}$	$3.1 \times 10^{9}$	0.21	$6.5 \times 10^{8}$	$9.9 \times 10^{9}$
$^{3}(V=O)OEP^{*}-C_{60}$	$3.2 \times 10^9$	0.19	$6.0 \times 10^{8}$	$4.5 \times 10^{9}$
<sup>3</sup> PdOEP*-C <sub>70</sub>	$3.3 \times 10^9$	0.60	$2.0 \times 10^{9}$	$3.3 \times 10^{9}$
$^{3}$ ZnOEP*- $C_{70}$	$3.0 \times 10^{9}$	0.49	$1.5 \times 10^9$	$8.9 \times 10^{9}$
$^{3}$ MgOEP*- $C_{70}$	$2.0 \times 10^{9}$	0.40	$8.0 \times 10^{8}$	$4.8 \times 10^{9}$
<sup>3</sup> (V=O)OEP*-C <sub>70</sub>	$2.0 \times 10^{9}$	0.25	$5.0 \times 10^{8}$	$4.5 \times 10^{9}$

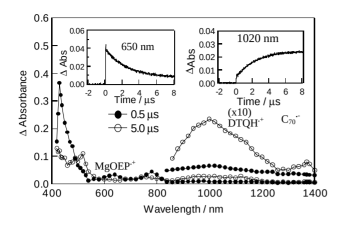


Fig. 10. Transient absorption spectra observed by  $550\,\mathrm{nm}$  laser excitation of MgOEP (0.1 mM) in the presence of  $C_{70}$  (0.1 mM) and DTQH (5 mM) in Ar-saturated BN. Inset: time profiles at 650 and  $1020\,\mathrm{nm}$  [93].

by ET to  $C_{70}$ , producing  $C_{70}^{\bullet-}$  at 1380 nm. Although, the absorption of MgOEP<sup>•+</sup> appeared at 640 nm in the absence of DTQH, in its presence, the rapid decay of MgOEP++ was observed, with concomitant rise of DTQH<sup>•+</sup> in the 800-1300 nm region. Thus, photosensitized ET occurs at first from  ${}^3MgOEP^*$  to  $C_{70}$ , yielding  $C_{70}^{\bullet-}$  and  $MgOEP^{\bullet+}$ , followed by the hole shift from MgOEP<sup>•+</sup> to DTOH, yielding DTQH<sup>•+</sup>, as summarized in Scheme 3. Such a hole shift is possible when the  $E_{ox}$  value of the hole shift reagent DTQH ( $E_{ox} = 0.32 \text{ V}$  versus SCE) is lower than that of MgOEP ( $E_{ox} = 0.53 \text{ V}$  versus SCE). The final back ET rate constant ( $k_{\rm fbet}$ ) was evaluated as  $6.1 \times 10^8 \, {\rm M}^{-1} \, {\rm s}^{-1}$  by following the long time decay profiles of DTQH $^{\bullet+}$  and C<sub>70</sub> $^{\bullet-}$ , which obey second-order kinetics. The decay of  $C_{70}^{\bullet-}$ in the presence of DTQH<sup>•+</sup> ( $k_{\text{fbet}} = 6.1 \times 10^8 \,\text{M}^{-1} \,\text{s}^{-1}$ ) is slowed down compared with that in the presence of  $MgOEP^{\bullet +}$  ( $k_{bet} = 4.7 \times 10^9 M^{-1} s^{-1}$ ).

### 2.3. Fullerene-chlorophylls [95]

In photosynthesis, the chlorophylls, Chls (close cousins of metalloporphyrins) play key roles in absorbing light energy over a wide spectral range and converting the light energy into the highly directional transfer of electrons. Green plants employ chlorophylls and magnesium—chlorins as the chromophores to harvest light. The investigations of oxidation—reduction reactions photosensitized by Chls in

$$\begin{pmatrix} C_{60}/C_{70} & ET & P^+ & HT & DTQH \\ k_{bet} & k_{fbet} & k_{ht} & \\ C_{60}/C_{70} & 3_P^* & P & DTQH^+ \end{pmatrix}$$

Scheme 3. Routes for the electron-transfer/hole shift cycle start with photoexcitation of P in the presence of  $C_{60}/C_{70}$  and DTQH in Ar-saturated BN.

vitro are of great importance to elucidate the mechanism of the primary photoreactions in photosynthesis [96,97].

One of the specific features of Chls is related to the quenching of their photoexcited states by compounds with high electron affinity via an ET process [99,100]. The quenching of excited states of Chls by quinones has been widely studied as a simple model system for the primary photoinduced CS in the chloroplast. It has been demonstrated by the flash photolysis and ESR techniques that various quinones quench <sup>3</sup>Chls\*. By ESR measurements, the signal of the semiquinone  $(Q^{\bullet-})$  was observed [98,99]. Also, by applying laser flash photolysis measurements, the intermediates Q<sup>•-</sup> and Chls<sup>•+</sup> were observed by the light excitation of Chls. The main problem frequently faced in the flash photolysis measurements is the overlap of the absorptions of the intermediates, which leads to difficulties in the interpretation of the mechanisms and quantitative analysis of the rates and yields of the ET processes. The absorption region of the Chls strongly overlaps with the absorption band of Q • at 435 nm. In addition, the absorption of <sup>3</sup>Chls\* masks most of the absorption band of the Chls<sup>•+</sup> in the visible region. In contrast with quinones, the transient absorptions of  $C_{60}^{\bullet-}$  and  $C_{70}^{\bullet-}$  in the near-IR region make it easy to study quantitatively the elemental steps in the PET processes [95].

A considerable insight into the details of the ET process in the systems of Chl-a/Chl-b and  $C_{60}/C_{70}$  via  $^3$ Chls\* can be obtained by applying 640 nm laser light, which selectively excites Chl-a/Chl-b. The transient absorption bands appeared at 480–500 nm, which are assigned unambiguously to  $^3$ Chls\* [96–100].

In the presence of  $C_{60}/C_{70}$ , the ET processes from  ${}^3\mathrm{Chl-a}^*/{}^3\mathrm{Chl-b}^*$  to  $C_{60}/C_{70}$  were observed by recording the diagnostic peaks of  $C_{60}^{\bullet-}/C_{70}^{\bullet-}$  in the near-IR region by the excitation of Chl-a/Chl-b. The second-order quenching rate constant  $(k_q)$ , ET quantum-yield  $(\Phi_{\mathrm{et}})$ , and ET rate constant  $(k_{\mathrm{et}})$  were evaluated, as in Table 6, in which the  $k_{\mathrm{et}}$  values from  ${}^3\mathrm{Chl-a}^*$  to  $C_{60}/C_{70}$  are slightly larger than the corresponding values from  ${}^3\mathrm{Chl-b}^*$  to  $C_{60}/C_{70}$ . The difference in the  $E_{\mathrm{ox}}$  and  $E_{\mathrm{T}}$  values between Chl-a and Chl-b may be responsible for the difference in the  $k_{\mathrm{et}}$  values. Moreover, the presence of the electron-withdrawing group (–CHO) decreases the electron-donor ability of Chl-b compared to Chl-a, with its electron-donating methyl group.

In benzene as a nonpolar medium, although rapid quenching of  ${}^3\mathrm{Chl}\text{-}a^*/{}^3\mathrm{Chl}\text{-}b^*$  by  $\mathrm{C}_{60}/\mathrm{C}_{70}$  was observed, both ET and EN processes from  ${}^3\mathrm{Chl}\text{-}a^*/{}^3\mathrm{Chl}\text{-}b^*$  to  $\mathrm{C}_{60}/\mathrm{C}_{70}$  are ruled out. This may be quite reasonable, because the  $E_{\mathrm{T}}$  values of Chl-a/Chl-b are lower than those of  $\mathrm{C}_{60}$  (1.5 eV). As a reason for the observed rapid quenching of  ${}^3\mathrm{Chl}\text{-}a^*/{}^3\mathrm{Chl}\text{-}b^*$  by  $\mathrm{C}_{60}$  in BZ, exciplex formation or collision deactivation may be considered.

By employing the excitation wavelength of the laser light at 532 nm, which selectively excites  $C_{70}$ , the ET process from Chl-a/Chl-b to  $^3C_{70}^*$  was also confirmed in polar solvents. The transient spectra exhibited the absorption band

Systems	$\Delta G_{\mathrm{et}}^{0} \; (\mathrm{kJ}  \mathrm{mol}^{-1})$	$k_{\rm q} \ ({\rm M}^{-1} \ {\rm s}^{-1})$	$\overline{arPhi_{ ext{et}}}$	$k_{\rm et} \ ({ m M}^{-1} \ { m s}^{-1})$	$k_{\text{bet}} \ (M^{-1} \ \text{s}^{-1})$
<sup>3</sup> Chl-a*-C <sub>60</sub> -BN	-31.4	$1.9 \times 10^{9}$	0.44	$8.4 \times 10^{8}$	$1.0 \times 10^{10}$
<sup>3</sup> Chl-a*-C <sub>70</sub> -BN	-39.6	$2.2 \times 10^{9}$	0.43	$9.4 \times 10^{8}$	$7.2 \times 10^9$
$^{3}$ Chl-b*- $C_{60}$ -BN	-27.5	$1.9 \times 10^{9}$	0.20	$3.8 \times 10^{8}$	$4.5 \times 10^{9}$
<sup>3</sup> Chl-b*-C <sub>70</sub> -BN	-34.9	$2.4 \times 10^{9}$	0.26	$6.1 \times 10^{8}$	$4.8 \times 10^{9}$
$^{3}$ Chl-a* $-$ C <sub>60</sub> $-$ BZ	_	$2.2 \times 10^{9}$	_	_	_
<sup>3</sup> Chl-a*-C <sub>70</sub> -BZ	_	$2.5 \times 10^9$	_	_	_
<sup>3</sup> C <sub>70</sub> *–Chl-a–BN	-55.7	$2.2 \times 10^{9}$	0.37	$8.1 \times 10^{8}$	$8.8 \times 10^{9}$
<sup>3</sup> C <sub>70</sub> *-Chl-b-BN	-45.3	$2.5 \times 10^{9}$	0.39	$9.7 \times 10^{8}$	$4.8 \times 10^{9}$
<sup>3</sup> C <sub>70</sub> *–Chl-a–BZ	_	$3.4 \times 10^{9}$	_	_	_

Table 6 Free energy changes ( $\Delta G_{\rm et}^0$ ) and kinetic parameters ( $k_{\rm q}$ ,  $\Phi_{\rm et}$ , and  $k_{\rm et}$ ) for the ET process between Chl-a/Chl-b and ( $C_{60}/C_{70}$ ) in BN and BZ

of  ${}^3C_{70}^*$  at 980 nm, which decayed with the concomitant formations of  $C_{70}^{\bullet-}$  and Chl-b $^{\bullet+}$  at 1380 and 780 nm, respectively.

The high electron-donor abilities of Chl-a/Chl-b to  ${}^3C_{70}^*$  and  ${}^3Chl$ -a\*/ ${}^3Chl$ -b\* to  $C_{60}/C_{70}$  are in good agreement with the similarly high donor abilities of MgOEP and ZnOEP, as shown in Tables 3 and 5. We have come to conclude that Chls have similar electron-donor ability, in spite of their long chain, electron-withdrawing substituents and Mg(II) central atom.

### 2.4. Fullerenes—phthalocyanine/naphthalocyanine [70,101]

Phthalocyanines (Pc) are a class of organic compounds that have attracted great attention because of their unique properties, such as semiconductivity, photoconductivity, photochemical reactivity, chemical stability, electrochromism, bio-organic and catalytic activity and their various applications in technology [102–109].

Several studies have been performed to examine the photophysical properties as well as the potential for ET from metal phthalocyanines (MPc;  $M=H_2$  and Zn in Fig. 11) to electron acceptor molecules. It has been reported that the photosensitivity of ZnPc in a polymeric binder is increased by the addition of  $C_{60}$  [110,111]. From photoemission experiments,  $C_{60}$  and  $C_{70}$  are expected to be appropriate electron-accepting materials when they are brought into contact with Pc in solids [63,64].

Fig. 11. Tetra-t-butylphthalocyanines.

In 1997, we reported a detailed study on the intermolecular ET between  $C_{60}/C_{70}$  and MPc via  ${}^3C_{60}^*/{}^3C_{70}^*$  by applying 532 nm nanosecond laser light in a polar solvent [70]. The selective excitation of  $C_{60}/C_{70}$  was possible in the presence of MPc, because MPc does not show any absorption at 532 nm at all. For example, the  $E_{ox}$  value of ZnPc generating ZnPc\*+ is +0.8 V versus SCE in BN; thus, the ET process is possible from ZnPc to  ${}^{3}C_{60}^{*}/{}^{3}C_{70}^{*}$ . Indeed, excitation of C<sub>60</sub>/C<sub>70</sub> in BN gives rise to the rapid formation of  $C_{60}^{\bullet-}/C_{70}^{\bullet-}$  at 1080 nm/1380 nm, and the rise of ZnPc $^{\bullet+}$  at 840 nm, with concomitant decay of  ${}^{3}C_{60}^{*}/{}^{3}C_{70}^{*}$ at 740 nm/980 nm. In contrast, in nonpolar solvents, although rapid quenching of  ${}^{3}C_{60}^{*}/{}^{3}C_{70}^{*}$  was observed in the presence of MPc, no evidence of formation of the radical ions was obtained by the nanosecond transient spectra, indicating absence of ET in the nonpolar solvent, because the energy levels for the radical ions are significantly raised. In the case of MPc, the evidence for EN was obtained by the rise of <sup>3</sup>MPc\* at 490 nm, which is reasonable because of the lower  $E_{\rm T}$  of MPc compared to that of  ${}^{3}{\rm C}_{60}^{*}/{}^{3}{\rm C}_{70}^{*}$ . These rate parameters are summarized in Table 7. It is noteworthy to mention that the ET quantum yield  $(\Phi_{et})$  via  ${}^{3}C_{70}^{*}$  (0.75) was found to be higher than that via  ${}^{3}C_{60}^{*}$  (0.50). Also, we proved that ZnPc acts as a stronger electron donor than H<sub>2</sub>Pc [70]. In BZ, the energy transfer occurs predominantly even for ZnPc.

Further continuation of this study [101] was performed for the ET process via  $^3\text{MPc}^*$  to  $\text{C}_{60}/\text{C}_{70}$  by applying a 670 nm laser, which selectively excites MPc in the presence of excess  $\text{C}_{60}/\text{C}_{70}$ . In contrast to the porphyrins, we observed photoionization character of ZnPc in polar media. That is, the transient absorption spectrum with 670 nm laser light of ZnPc (0.1 mM) in BN exhibited absorption bands at 490 and 840 nm, corresponding to  $^3\text{ZnPc}^*$  and  $\text{ZnPc}^{\bullet+}$ , respectively. This indicated the occurrence of photoionization via  $^3\text{ZnPc}^*$  with a quantum yield ( $\Phi_{\text{ion}}$ ) less than 0.1 in BN. In the case of  $^3\text{H}_2\text{Pc}^*$ , such photoionization did not occur even in polar solvents such as BN.

In the presence of  $C_{60}/C_{70}$ , the transient absorption spectra observed by excitation of ZnPc exhibited growth of the absorption bands of ZnPc $^{\bullet+}$  at 840 nm and  $C_{60}^{\bullet-}/C_{70}^{\bullet-}$  at 1080 nm/1380 nm, accompanied by a concurrent decay of

 $1.8 \times 10^{8}$ 

 $k_{\rm et}~({
m M}^{-1}~{
m s}^{-1})$  $k_{\rm en}~({\rm M}^{-1}~{\rm s}^{-1})$  $k_{\rm q}~({\rm M}^{-1}~{\rm s}^{-1})$  $\Phi_{\mathrm{et}}$  $\Phi_{\mathrm{en}}$  $^3$ C $_{60}$ \*-ZnPc-BN  $1.6 \times 10^{9}$ 0.50  $8.0 \times 10^{8}$ 0.50  $8.0 \times 10^8$  $^{3}C_{60}^{*}$ -ZnPc-BZ  $2.1 \times 10^{9}$  $2.1 \times 10^9$ 0.00 1.00  ${}^{3}C_{60}^{*}-H_{2}Pc-BN$  $3.0 \times 10^{9}$  $5.4 \times 10^8$  $2.5 \times 10^8$ 0.18 0.82  $^{3}C_{60}^{*}-H_{2}Pc-BZ$  $2.8 \times 10^{9}$ 0.00 0.00  $2.8 \times 10^{9}$  $^3C_{70}^*$ -ZnPc-BN  $1.7 \times 10^{9}$  $1.3 \times 10^{9}$  $4.0 \times 10^8$ 0.75 0.25  $^{3}\text{C}_{70}^{*}$ -ZnPc-BZ  $1.8 \times 10^{9}$ 0.00 1.00  $1.8 \times 10^{9}$  $^{3}\text{C}_{70}^{*}\text{-H}_{2}\text{Pc-BN}$  $7.0 \times 10^8$  $3.5 \times 10^{9}$  $2.8 \times 10^{8}$ 0.20 0.80

0.00

Table 7 Kinetic parameters  $(k_q, \Phi_{\text{et}}, \text{ and } k_{\text{et}})$  for electron transfer (ET) and energy transfer (EN) between  ${}^3C_{60}^*/{}^3C_{70}^*$  and MPc in Ar-saturated BN and BZ

 $^{3}\text{C}_{70}^{*}\text{-H}_{2}\text{Pc-BZ}$ 

 $^3$ ZnPc\*. Furthermore, the rise rate of ZnPc\*+ was quite similar to that of  $C_{60}$ \*- $/C_{70}$ \*-. These findings indicate that ET takes place from  $^3$ ZnPc\* to  $C_{60}$ , with  $\Phi_{et}=$  ca. 0.8, accompanied by only a small contribution from photoionization. The rate parameters are summarized in Table 8.

 $1.8 \times 10^{9}$ 

It was found that the  $k_{\rm q}$  value of the  ${}^3{\rm ZnPc^*-C_{60}}$  system (8.7 × 10<sup>8</sup> M<sup>-1</sup> s<sup>-1</sup>) is 300 times higher than that for the  ${}^3{\rm H_2Pc^*-C_{60}}$  system (2.9 × 10<sup>6</sup> M<sup>-1</sup> s<sup>-1</sup>). Moreover, we observed that the electron donor ability of  ${}^3{\rm ZnPc^*}$  to  ${\rm C_{60}}$  ( $\Phi_{\rm et}=0.77$ ) is 10 times higher compared to  ${}^3{\rm H_2Pc^*}$  ( $\Phi_{\rm et}=0.07$ ). In order to confirm whether such a large difference in electron donor ability between  ${}^3{\rm ZnPc^*}$  and  ${}^3{\rm H_2Pc^*}$  was specific to the spherical fullerene molecule, we examined ET from  ${}^3{\rm ZnPc^*}$  or  ${}^3{\rm H_2Pc^*}$  to benzoquinone (BQ) as a representative of flat small molecules. The results revealed that the higher electron donor ability is a general characteristic of  ${}^3{\rm ZnPc^*}$  and  ${}^3{\rm H_2Pc^*}$ , but not a specific characteristic of the fullerene acceptors.

The  $\Delta G_{\rm et}^0$  values evaluated for ET occurring from  $^3{\rm ZnPc^*}$  and  $^3{\rm H_2Pc^*}$  to  ${\rm C}_{60}$  were found to be -19.7 and -29.7 kJ mol $^{-1}$ , respectively. Similarly for BQ,  $\Delta G_{\rm et}^0$  values were calculated for  $^3{\rm ZnPc^*}$  (-13.4 kJ mol $^{-1}$ ) and  $^3{\rm H_2Pc^*}$  (-17.2 kJ mol $^{-1}$ ). These negative  $\Delta G_{\rm et}^0$  values imply that the  $k_{\rm et}$  values via  $^3{\rm ZnPc^*}$  and  $^3{\rm H_2Pc^*}$  to  ${\rm C}_{60}$  (or  ${\rm C}_{70}$  and BQ) are all close to the  $k_{\rm diff}$  value ( $5.6 \times 10^9$  M $^{-1}$  s $^{-1}$  in BN). However, the  $\Phi_{\rm et}$  values indicated that the electron acceptor ability of  ${\rm C}_{70}$  is slightly lower than that of  ${\rm C}_{60}$ , although the calculated  $\Delta G_{\rm et}^0$  values did not predict such a

tendency. The lower electron acceptor ability of  $C_{70}$  compared to  $C_{60}$  is also shown for  $^3H_2Pc^*$  (Table 8). In nonpolar solvents, although quenching of  $^3MPc^*$  was observed, there was no evidence for ET and/or EN in the transient absorption spectra. The decay time profile of  $C_{60}^{\bullet-}$  observed in the longer time scale obeyed second-order kinetics in BN, indicating that  $ZnPc^{\bullet+}$  and  $C_{60}^{\bullet-}$  recombine after being solvated as free radical ions. The  $k_{bet}$  values from  $C_{60}^{\bullet-}$  and  $C_{70}^{\bullet-}$  to  $ZnPc^{\bullet+}$  in polar solvents were evaluated as being close to  $k_{diff}$ .

1.00

Recently, we reported the ET process in the systems composed of oxotitanium(IV) tetra-t-butyl-phthalocyanine ((Ti=O)Pc in Fig. 11) with  $C_{60}$  and  $C_{70}$  to examine the effect of the metal ion on the electron transfer process [112]. (Ti=O)Pc has been widely applied to the photoelectric conversion system, although it has low solubility in organic solvents. The absorption peaks, Soret and Q-bands, of (Ti=O)Pc were observed at 350 and 620–705 nm, respectively. The fluorescence time profile shows a single exponential decay, giving a lifetime of 5.1 ns. No dynamic quenching of  $^1$ (Ti=O)P\* was observed on addition of  $C_{60}$  or  $C_{70}$ , even in polar solvents.

The transient absorption spectra observed by the selective excitation of (Ti=O)Pc with laser light at 355 nm in Ar-saturated toluene exhibited the rapid decay of an absorption band at 1400 nm, with a rate of ca.  $1 \times 10^8 \, \mathrm{s}^{-1}$ , which may be assigned to the  $S_1$ – $S_n$  transition of (Ti=O)Pc. With the decay band at 1400 nm, the growth of a band at 1300 nm

Table 8 Kinetic parameters ( $k_q$ ,  $\Phi_{et}$ , and  $k_{et}$ ) for ET between ZnPc–BQ and C<sub>60</sub>/C<sub>70</sub> in Ar-saturated BN and BZ;  $k_{bet}$  between C<sub>60</sub>•-/C<sub>70</sub>•- and ZnPc•+/BQ•+,  $k_q$ 

System	$k_{\rm q}~({ m M}^{-1}~{ m s}^{-1})$	$arPhi_{ m et}$	$k_{\rm et} \ ({\rm M}^{-1} \ {\rm s}^{-1})$	$k_{\text{bet}} \ (M^{-1} \ s^{-1})$
<sup>3</sup> ZnPc*-C <sub>60</sub> -BN	$8.7 \times 10^{8}$	0.77	$6.7 \times 10^{8}$	$3.4 \times 10^{9}$
$^{3}$ ZnPc* $-$ C <sub>60</sub> $-$ BZ	$6.3 \times 10^8$	0.00	_a	_a
$^{3}$ ZnPc* $-$ C <sub>70</sub> $-$ BN	$1.6 \times 10^{9}$	0.48	$7.7 \times 10^{8}$	$3.3 \times 10^{9}$
$^{3}$ ZnPc* $-$ C <sub>70</sub> $-$ BZ	$1.0 \times 10^{9}$	0.00	_a	_a
<sup>3</sup> ZnPc* –BQ–BN	$3.3 \times 10^{8}$	0.83	$2.7 \times 10^{8}$	$3.5 \times 10^{9}$
$^{3}$ H <sub>2</sub> TBPc* $-$ C <sub>60</sub> $-$ BN	$2.7 \times 10^{6}$	0.07	$1.9 \times 10^{5}$	$5.0 \times 10^{9}$
$^{3}\text{H}_{2}\text{Pc}^{*}$ $-\text{C}_{60}$ –BZ	$4.9 \times 10^{6}$	0.00	_a	_a
$^{3}\text{H}_{2}\text{Pc}^{*}$ $-\text{C}_{70}$ –BN	<10 <sup>5</sup>	_b	_b	_b
$^{3}\text{H}_{2}\text{Pc}^{*}$ -BQ-BN	< 10 <sup>5</sup>	_b	_b	_b

a No PET was observed.

<sup>&</sup>lt;sup>a</sup> No PET was observed.

<sup>&</sup>lt;sup>b</sup> Too weak to observe or too slow to observe.

$$(\text{Ti=O})\text{Pc} + {^{3}\text{C}_{60}}^{*}/{^{3}\text{C}_{70}}^{*} \underbrace{\overset{k_{\text{ass}}}{\overset{k_{\text{ass}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}}{\overset{k_{\text{cq}}}}}}{\overset{k_{\text{cq}}}$$

Scheme 4. Routes for the electron-transfer process occurring by the photoexcitation of C<sub>60</sub>/C<sub>70</sub> in the presence of (Ti=O)Pc in BN.

was observed. Since the species with absorption at 1300 nm has a long lifetime ( $\tau = 67 \,\mu s$ ), this species was attributed to  $^3$ (Ti=O)Pc\*. By the selective excitation of  $C_{70}$  and  $C_{60}$  in a nonpolar solvent, the EN process from  $^3C_{70}^*$  and  $^3C_{60}^*$  to (Ti=O)Pc was confirmed with rate constants of  $3.3 \times 10^9$  and  $2.0 \times 10^9 \, M^{-1} \, s^{-1}$ , respectively. This is the same tendency observed for MPc such as ZnPc and  $H_2$ Pc in Table 6 [70].

In polar BN solvent, the ET process from (Ti=O)Pc to  ${}^3C_{60}^*$  was confirmed by observing the decay of  ${}^3C_{60}^*$  at 750 with a concomitant rise of (Ti=O)Pc ${}^{\bullet +}$  at 880 nm and  ${}^6C_{60}^{\bullet -}$  at 1080 nm. Similarly, the ET process was confirmed from (Ti=O)Pc to  ${}^3C_{70}^*$ . The  $k_{1st}$  value of the rise of  ${}^3C_{60}^{\bullet -}$  was found to be smaller than the  $k_{1st}$  value for the decay of  ${}^3C_{60}^*$ , which might suggest some process intermediate between the decay and rise, such as triplet exciplex formation (Scheme 4).

The  $\Delta G_{\rm et}^0$  for ET from (Ti=O)Pc to  ${}^3{\rm C}_{60}^*$  was evaluated to be 0.64 eV by the Rehm–Weller equation. The  $\Phi_{\rm et}$  value via  ${}^3{\rm C}_{60}^*$  was evaluated as 0.2. Such a low  $\Phi_{\rm et}$  value for (Ti=O)Pc- ${}^3{\rm C}_{60}^*$  system compared to ZnPc- ${}^3{\rm C}_{60}^*$  system suggests the presence of a deactivation process of  ${}^3{\rm C}_{60}^*$  without ET; i.e., collisional quenching.

We have also investigated the ET process of the zinc tetra-*t*-butyl-naphthalocyanine (ZnNc in Fig. 12) and C<sub>60</sub>/C<sub>70</sub> systems via <sup>3</sup>ZnNc\* to probe the structural effect on the ET process [113]. The steady-state absorption spectra of ZnNc appear at 780 nm (Fig. 7), which is at a longer wavelength compared to ZnTPP and ZnPc. Since C<sub>60</sub> and C<sub>70</sub> have no appreciable absorption intensity at 650 nm, ZnNc can be selectively excited by the 670 nm laser light. The transient absorption spectra of ZnNc in BN (Fig. 13a) exhibited the intense absorption bands of <sup>3</sup>ZnNc\* at 600 and 770 nm accompanied by the weak growth of ZnNc\*+ at 970 nm. As shown in the inset of Fig. 13a, the slow rise of ZnNc\*+ at 970 nm seems to be a mirror image of

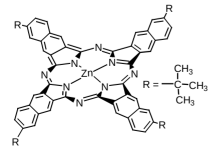


Fig. 12. Zinc tetra-t-butylnaphthalocyanine.

the slow decay of  ${}^3ZnNc^*$  at 600 nm, suggesting that the photoionization takes place via  ${}^3ZnNc^*$ . In the presence of  $C_{60}$ , the ET process from  ${}^3ZnNc^*$  to  $C_{60}$  was confirmed by the growth of the absorption bands of  $ZnNc^{\bullet+}$  at 970 nm and  $C_{60}^{\bullet-}$  at 1080 nm, accompanied by a concurrent decay of the absorption band of  ${}^3ZnNc^*$  at 600 and 770 nm (Fig. 13b). A similar ET process was observed in the case of  $C_{70}$  with  ${}^3ZnNc^*$  (Fig. 13c), in which the growth of the ab-

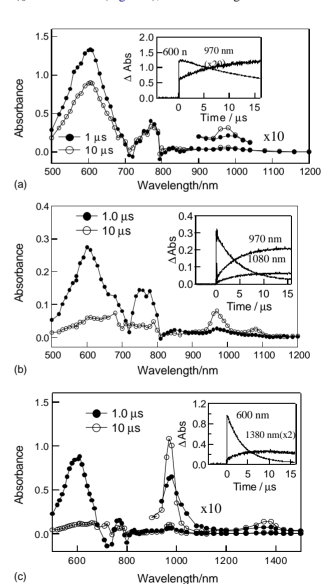


Fig. 13. Transient absorption spectra obtained by  $650\,\mathrm{nm}$  laser light of ZnNc (0.1 mM): (a) in the absence and (b) in the presence of  $C_{60}$  (0.1 mM) and (c)  $C_{70}$  (0.1 mM) in Ar-saturated BN. Inset: time profiles.

sorption bands of ZnNc\*+ (970 nm) and  $C_{70}$ \*- (1380 nm) was accompanied by the decay of  ${}^3$ ZnNc\* (600 nm). These findings indicate that ET occurs from  ${}^3$ ZnNc\* to  $C_{60}/C_{70}$ , with a small contribution from photoionization in polar solvents. The weaker absorption intensity of  $C_{70}$ \*- at 1380 nm (Fig. 13c) compared to that of  $C_{60}$ \*- at 1080 nm (Fig. 13b) was attributed to the smaller extinction coefficient ( $\varepsilon$ ) value of  $C_{70}$ \*- compared to that of  $C_{60}$ \*-. The second-order quenching rate constants ( $k_q$ ) for ET from  ${}^3$ ZnNc\* to  $C_{60}/C_{70}$  were evaluated. It was found that the  $k_q$  value for the  ${}^3$ ZnNc\*- $C_{60}$  system (1.45 × 10<sup>9</sup> M<sup>-1</sup> s<sup>-1</sup>) is much higher compared to that of the  ${}^3$ ZnNc\*- $C_{70}$  system (1.3 × 10<sup>8</sup> M<sup>-1</sup> s<sup>-1</sup>).

Summarizing the results of intermolecular ET of the  $C_{60}/C_{70}$ –MP (MTPP/Chl/MPc/MNc) systems, the following conclusions were drawn:

- (i) By changing the excitation wavelength and concentration, it was possible to change the ET routes from (a)  ${}^3\text{MP*}$  to  $\text{C}_{60}/\text{C}_{70}$  and (b) MP to  ${}^3\text{C}_{60}{}^*/{}^3\text{C}_{70}{}^*$ . The  $\Phi_{\text{et}}$  values of route (b) are usually higher than those of route (a) in polar solvents.
- (ii) In all cases, the  $\Phi_{et}$  values of ZnP–C<sub>60</sub>/C<sub>70</sub> systems are significantly higher compared to H<sub>2</sub>P–C<sub>60</sub>/C<sub>70</sub> in polar solvents.
- (iii) In the case of MOEP, the observed  $\Phi_{et}$  values of M = Pd, Zn, and Mg are larger than those of Co, Ni, and Cu for  ${}^3C_{60}^*$ .
- (iv) Chls have high electron donor abilities, similar to ZnTPP and MgOEP.
- (v) (Ti=O)Pc shows a reactivity quite different from others such as (V=O)OEP.
- (vi) MP, MPc and  $C_{60}/C_{70}$  absorbing wide visible region may be useful for applications such as photosynthetic solar energy conversion. For this purpose, the most important observation is the similar electron acceptor ability of  $C_{70}/^3C_{70}^*$  to  $C_{60}/^3C_{60}^*$ , which suggests that it is unnecessary to rigorously remove  $C_{70}$  from the crude samples of  $C_{60}$ .

### 3. Photoinduced electron transfer in supramolecular fullerene-porphyrin/phthalocyanines systems

Supramolecular systems composed of porphyrin and fullerene moieties have received much attention from researchers in recent years [73–82]. These systems are composed of porphyrin and fullerene derivatives functionalized in such a way that the two entities are able to diffuse together and reversibly bind in solution. The modes of binding most often employed include  $\pi$ – $\pi$  interactions, electrostatic attraction, hydrogen bonding, and axial ligation via a nitrogen-based ligand to the metal center of the metalloporphyrin.

The self-assembled donor-acceptor systems offer several advantages over intermolecular systems. First, the relative orientation of the donor and acceptor can be controlled, in some cases. This is quite important, since ET rates are dependent upon orbital overlap and distance between the donor and acceptor moieties. Second, for intermolecular systems, ET is a diffusion-controlled process, while for supramolecular systems this process is only partially governed by diffusion rates, but also by binding strength and concentration. Also, for intermolecular systems, the entity that is excited usually has enough time to undergo the ISC process from the singlet excited state to the triplet excited state before colliding with a donor or acceptor. Therefore, most intermolecular systems undergo ET via the triplet excited state. However, for self-assembled systems, in which the conditions have been properly adjusted so that a sufficient amount (>99%) of complexed donor-acceptors are present in solution, the excited species usually do not have enough time to undergo the ISC process. Therefore, most self-assembled systems undergo ET from the short-lived singlet excited state. Third, since the binding of the donor-acceptor complex is reversible in nature, after ET occurs, the individual charge-separated species (D<sup>•+</sup> and A<sup>•-</sup>) can diffuse away from each other, creating a long-lived SSIP in a sufficiently polar medium; thus, increasing the lifetime of the CS state.

### 3.1. Fullerene-porphyrin systems coordinated via axial ligation

The porphyrin macrocycle is capable of binding a variety of transition metals within its central cavity, thus leaving the positions axial to the plane of the porphyrin ring available for binding with a variety of ligands. In 1999, three different research groups studied systems composed of  $C_{60}$  functionalized with a coordinating ligand capable of axially ligating to the metalloporphyrin metal center (MTPP; M = Zn and Ru) [73–81,113–115].

A system composed of a pyridine-appended  $C_{60}$  (py $\sim C_{60}$ ) axially ligated to ZnTPP through the pyridine nitrogen (Fig. 14a) was studied by D'Souza and co-workers [114]. Upon complexation of py $\sim$ C<sub>60</sub> to ZnTPP, (the symbol  $\sim$ refers to a covalent bond) the optical absorption spectrum experienced a red shift of the Soret band. A control experiment using a phenyl-appended C<sub>60</sub> derivative exhibited no such spectral shift, confirming that axial coordination occurs through the pyridine nitrogen, but not through the pyrrolidine nitrogen. The binding constant (K) was obtained from absorption spectral data using the Scatchard method to be  $7337\,M^{-1}$  for ZnTPP $\leftarrow$ py $\sim$ C<sub>60</sub>, (the symbol  $\leftarrow$  refers to coordination bond). The estimated ET rate for the system from steady-state fluorescence quenching studies was found to be  $(2.4 \pm 0.3) \times 10^8 \,\mathrm{s}^{-1}$ , which is smaller than those of covalently bonded ZnTPP~C<sub>60</sub> dyads [50–52].

Two similar systems composed of a pyridine-appended  $C_{60}$  molecule axially ligated to MTPP (M = Zn and Ru in Fig. 14b) were studied by Pasimeni and co-workers [72,73]. The Ru based system exhibited photochemistry that was strongly solvent-dependent in nature. In a nonpolar solvent

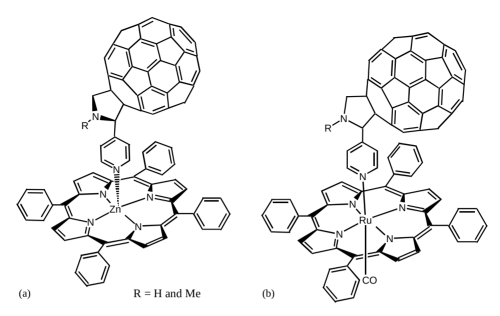


Fig. 14. Structure of the (a) zinc tetraphenylporphyrin:(4-pyridyl)fulleropyrrolidine and (b) ruthenium tetraphenylporphyrin:(4-pyridyl)fulleropyrrolidine dyads.

such as toluene, the excitation of the RuTPP leads primarily to the formation of  ${}^3RuTPP^*$  through a rapid ISC process following the initial excitation. After employing a 532 nm laser light pulse, the transient absorption spectrum of the dyad (RuTPP $\leftarrow$ py $\sim$ C<sub>60</sub>) at 85 ps exhibited a broad absorption maximum at 710 nm with a shoulder at 820 nm. The 710 nm feature corresponds to the absorption of the  ${}^3C_{60}^*$  moiety. These findings suggest intramolecular energy transfer from  ${}^3RuTPP^*$  to the py $\sim$ C<sub>60</sub> moiety. The dyad had a  $\Phi_{\rm ISC}$  value of 0.65 and lifetime of 43  $\mu$ s.

$$[^{3}RuTPP^{*}\leftarrow py\sim C_{60}] \rightarrow [RuTPP\leftarrow {}^{3}py\sim C_{60}^{*}]$$
 (6)

Upon changing to more polar solvents such as BN or dichloromethane, a marked change in the photochemical behavior of the system was observed. Upon irradiation with a laser light pulse at 532 nm of the dyad in BN, the Q-bands were bleached out, and a set of new peaks at 565, 610, and 670 nm were observed in the transient absorption spectrum. These peaks correspond to RuTPP $^{\bullet+}$ . Also, a peak at 1010 nm appeared in the near-IR region of the spectrum, which decayed after about 50  $\mu$ s. This peak corresponds to the formation of  $C_{60}^{\bullet-}$ . These data support the mechanism of ET via  $^3$ RuTPP $^*$  in BN.

$$[^{3}RuTPP^{*} \leftarrow py \sim C_{60}] \rightarrow [RuTPP^{\bullet +} \leftarrow py \sim C_{60}^{\bullet -}]$$
(7)

The same experiment in dichloromethane yielded different results. The data are consistent with the ET mechanism and the formation of the radical ion-pair, as was observed in BN; however, the CR process occurred much faster (<4 ns). This is attributed to the ability of BN to compete with the pyridine ligand for the axial coordination site. This promotes bond cleavage of the Ru—pyridine bond, thus creating a SSIP,

which slows down the CR process. However, in solvents not capable of axially ligating to the Ru metal center, this process does not occur and hence leads to a very fast charge recombination [73].

A similar system was studied for ZnTPP and  $C_{60}$  with pyridine axially ligated to the Zn atom. Transient absorption spectra recorded in toluene and dichloromethane after a laser light flash (532 nm) yielded similar results as compared to the ruthenium-based dyad. Broad absorptions appeared at 715, 960, and 1010 nm, corresponding to ZnTPP $^{\bullet+}$  and  $C_{60}^{\bullet-}$ , respectively [74]. These data indicate that the system undergoes ET from  $^1$ ZnTPP $^*$  to  $C_{60}$ .

$$[^{1}\text{ZnTPP}^{*} \leftarrow \text{py} \sim \text{C}_{60}] \rightarrow [\text{ZnTPP}^{\bullet +} \leftarrow \text{py} \sim \text{C}_{60}^{\bullet -}]$$
 (8)

Investigation of the same dyad system in BN using transient absorption methods revealed the formation of a broad absorption around 730 and 1010 nm, in which the latter corresponds to the formation  $C_{60}^{\bullet-}$ . However, the formation  $C_{60}^{\bullet-}$  was revealed via two routes; a slow component was due to ET from  $^3$ ZnTPP\*, while the fast component was due to ET from  $^1$ ZnTPP\*. This can be rationalized by considering that BN is a coordinating solvent capable of competing with  $C_{60}$  pyridine for the axial coordination site. Thus, both inter- and intramolecular ET mechanisms are possible pathways.

Our research groups recently performed a systematic study on donor–acceptor systems composed of  $C_{60}$  bearing nitrogen-based ligands (o-pyridyl, m-pyridyl, p-pyridyl, N-phenyl imidazole) axially ligated to ZnTPP (Fig. 15) [79]. UV-Vis spectral data were used to determine the binding constants (K) for each  $C_{60}$  derivative with ZnTPP. The trend observed for the K values was: o-pyridyl  $\ll m$ -pyridyl  $\ll p$ -pyridyl  $\ll N$ -phenyl imidazole. Thermodynamic param-

Ligand <sup>a</sup>	$K^{b}$ $(M^{-1})$	$\Delta G \text{ (kJ mol}^{-1})$	$\Delta H \text{ (kJ mol}^{-1}\text{)}$	$\Delta S (J K^{-1} \text{ mol}^{-1})$
Pyridine	7750	-22.1	-27.3	-17.1
N-Phenyl imidazole	18620	-24.0	-32.4	-28.2
1	c	_	_	_
<b>2</b> <sup>b</sup>	7740	-22.1	-26.1	-12.3
3	7660	-22.1	-26.7	-15.4
4	7170	-22.0	-26.9	-16.5
5	16110	-23.6	-31.9	-28.1

Table 9
Formation constant, K and thermodynamic parameters for self-assembled zinc tetraphenylporphyrin–fulleropyrrolidine conjugates in o-DCE

eters for these systems were also evaluated, as summarized in Table 9. These data suggested that both enthalpy and entropy changes contribute to the overall free energy change for the self-assembly of the systems studied.

X-ray structural and ab initio computational studies on of the supramolecules involving p-pyridyl derivatized fulleropyrrolidine and ZnTPP were also performed [116]. In the studied solid state structure, the zinc-to-axially coordinated pyridyl nitrogen distance was found to be 2.158 Å which is compatible with a 2.075 Å average distance of the porphyrin ring Zn–N bonds. Importantly, the center-to-center distance between the porphyrin zinc ion and fullerene was ca. 9.53 Å. Additional intermolecular interactions between the zinc porphyrin and the  $C_{60}$  unit that is not directly coordinated to the zinc were also observed in the crystal packing.

Steady-state fluorescence experiments were performed in o-dichlorobenzene (o-DCB) on the dyad systems. Stern–Volmer plots were used to calculate the bimolecular quenching constants ( $k_q^S$ ). These values were found to be 2–3 orders of magnitude greater than would be expected for a diffusion-controlled process. In BN, a coordinating solvent, the quenching constant decreases significantly, probably due to the competition between the solvent and the  $C_{60}$  derivative for the axial coordination site.

Time-resolved fluorescence experiments showed a similar trend to that of the steady-state data. Upon addition of the  $C_{60}$ -coordinating derivatives to a solution containing ZnTPP the quenching of  $^1ZnTPP^*$  was accelerated (monitored at

Fig. 15. Structure of the fullerene derivatives functionalized to bear pyridine and imidazole ligands used to complex with zinc porphyrin.

600 nm). Upon addition of 3, the CS rate constant ( $k_{cs}$ ) and quantum yield ( $\Phi_{cs}$ ) increased only slightly. However, for 5, these values increased greatly (Table 10). Also, the quenching process was found to consist of both a slow and a fast process, probably due to bound and unbound species present in solution.

Time-resolved transient absorption spectra were recorded to confirm the mechanism of CS for the studied dyads. Transient absorption spectra of 5 with ZnTPP in o-DCB (530 nm laser excitation) exhibit a peak at 1000 nm after 0.01 µs, which is attributed to  $C_{60}^{\bullet-}$ . This peak then decays very rapidly ( $<10^8 \,\mathrm{s}^{-1}$ ), indicating that CS occurs from  $^1\mathrm{ZnTPP}^*$ and decays very rapidly under these experimental conditions. The transient absorption spectra in BN showed very different features. Here, the absorption bands of <sup>3</sup>ZnTPP\* at 840 and 700 nm decay quickly and the rise of the transient absorption band at 1020 nm was attributed to  $C_{60}^{\bullet-}$ . These data indicate that, for the dyad system composed of 5 and ZnTPP in BN, the ET process occurs mainly from the triplet excited state (<sup>3</sup>ZnTPP\*), with an estimated ET rate constant  $(k_{\rm et}^{\rm T})$  of  $2.5 \times 10^7 \, {\rm M}^{-1} \, {\rm s}^{-1}$ . The different photochemical pathways are shown in the energy diagrams in Fig. 16.

Table 10 Fluorescence lifetimes ( $\tau$ ), CS rate constants ( $k_{\text{CS}}^{\text{singlet}}$ ) and quantum yields ( $\Phi_{\text{CS}}^{\text{singlet}}$ ) for ZnTPP $\leftarrow$ fulleropyrrolidine supramolecular dyads in o-DCB

Compounda	Solvent	$\tau_{\rm f}$ (ns) (fraction %)	$k_{\rm CS}^{\rm singlet} \ ({\rm s}^{-1})^{\rm b}$	$\Phi_{\mathrm{CS}}^{\mathrm{singlet}\;\mathrm{c}}$
ZnTPP	DCB	2.10 (100)	_	_
ZnTPP← <b>3</b> (1:1)	DCB	1.88 (100)	$5.6 \times 10^{7}$	0.10
ZnTPP← <b>3</b> (1:6)	DCB	1.85 (100)	$6.3 \times 10^{7}$	0.12
ZnTPP← <b>5</b> (1:1)	DCB	0.058 (30)	$1.7 \times 10^{10}  d$	$0.97^{d}$
		2.00 (70)	$5.3 \times 10^{9} \mathrm{e}$	0.61 <sup>e</sup>
ZnTPP← <b>5</b> (1:6)	DCB	0.058 (50)	$1.7 \times 10^{10} \mathrm{d}$	0.97 <sup>d</sup>
		2.00 (50)	$8.6 \times 10^{9} \mathrm{e}$	$0.84^{e}$

<sup>&</sup>lt;sup>a</sup> For abbreviations see Fig. 15.

<sup>&</sup>lt;sup>a</sup> For abbreviations see Fig. 15.

<sup>&</sup>lt;sup>b</sup> At 298 K.

<sup>&</sup>lt;sup>c</sup> No appreciable binding was observed.

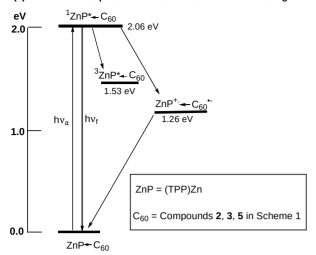
<sup>&</sup>lt;sup>b</sup>  $k_{\text{CS}}^{\text{singlet}} = (1/\tau_{\text{f}})_{\text{sample}} - (1/\tau_{\text{f}})_{\text{ref}}.$ 

 $<sup>^{\</sup>text{c}}\Phi_{\text{cs}}^{\text{singlet}} = [(1/\tau_{\text{f}})_{\text{sample}} - (1/\tau_{\text{f}})_{\text{ref}}]/(1/\tau_{\text{f}})_{\text{sample}}.$ 

d From the fast decay.

e Average value.

### (a) Intramolecular photochemical events: Noncoordinating solvent



### (b) Intermolecular photochemical events: Coordinating solvent

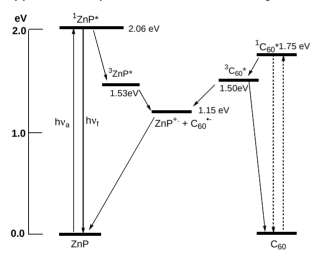


Fig. 16. Energy level diagram showing (a) intramolecular photochemical events of ZnTPP—fulleropyrrolidine dyads and (b) intermolecular photochemical events of ZnTPP—fulleropyrrolidine mixture.

Our research groups recently prepared and studied a supramolecular dyad system composed of an imidazole-appended  $C_{60}$  (5), which axially coordinates via the imidazole nitrogen to the central metal of ZnNc (Fig. 17) [113]. UV-Vis spectral data were used to confirm the formation of the self-assembled dyad system. Upon addition of 5 to a solution containing ZnNc (toluene), the absorbance band at 767 nm exhibited diminished intensity, and isosbestic points appeared at 675, 717, 752, and 791 nm. The K value for the dyad in toluene solution was determined to be  $6.2 \times 10^4 \, \mathrm{M}^{-1}$  by the Scatchard method. This value is an order of magnitude larger than that of the counterpart ZnTPP dyad system.

Steady-state fluorescence experiments were performed on the self-assembled dyad system. Upon addition of  $\mathbf{5}$  to a solution containing ZnNc (toluene or o-DCB), the emission bands at 781 and 812(sh) nm were gradually quenched to

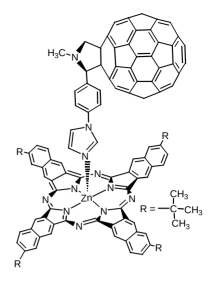


Fig. 17. Structure of the naphthalocyanine:fullerene conjugate self-assembled via axial ligation.

about 30% of the original intensity. Also, the emission band at 781 nm experiences a 3 nm blue shift compared to the original uncoordinated ZnNc. Stern–Volmer plots were used to determine the quenching constant for the dyad system. This value was determined to be four orders of magnitude larger than what would be expected for a diffusion-controlled process. This indicates that an intramolecular quenching process is the predominant quenching pathway for the dyad system.

Time-resolved fluorescence spectral studies were performed on the dyad system. The excited state of pure ZnNc had a lifetime of 2.42 ns. Upon addition of 5 to form the complex ( $5\rightarrow$ ZnNc) (toluene) the excited state decayed bi-exponentially, having both fast and slow components. The fast component had a lifetime of around 71 ps, while the slow component had a lifetime of around 2.14 ns. The slow component lifetime was similar to that obtained for uncomplexed ZnNc. The short-lived nature of the excited state of ZnNc in the complex ( $5\rightarrow$ ZnNc) suggests quenching via ET from the singlet excited state of ZnNc ( $^1$ ZnNc\*). The  $k_{cs}$  value was determined to be  $1.4\times10^{10}\,\mathrm{s}^{-1}$ , with a  $\Phi_{cs}$  value of 0.97 for the process (toluene).

Picosecond transient absorption spectra were obtained in order to determine the nature of the excited state photochemical reactions in this time domain. Upon excitation of the dyad with 388 nm laser light, new bands appeared at 710 and 985 nm in the time region of 10-200 ps (Fig. 18). These bands were attributed to the formation of  $ZnNc^{\bullet+}$ . The band expected at 1000 nm representing  $5^{\bullet-}$  was not observed, probably due to masking by the intense band at 985 nm. The  $k_{cs}$  value for the dyad was evaluated to be  $1.4 \times 10^{10}$  s<sup>-1</sup>, which is in agreement with the value determined by fluorescence lifetime measurements, while the  $k_{cr}$  value was evaluated to be  $8.5 \times 10^8$  s<sup>-1</sup>.

Nanosecond transient absorption spectra were also obtained for the self-assembled dyad. Upon excitation of the dyad (5→ZnNc) with 532 nm laser light, intense absorption

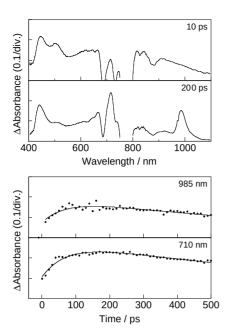


Fig. 18. Upper panels: transient absorption spectra of ZnNc  $(0.03 \, \text{mM})$  in the presence of  $C_{60}\text{Im}$   $(0.3 \, \text{mM})$  in toluene. Lower panels: time profiles at 985 and 710 nm [113].

bands were observed between 600 and 700 nm after 100 ns, corresponding to  ${}^{3}\text{ZnNc}^{*}$  and  ${}^{3}\textbf{5}^{*}$ . After 10 ns, absorption bands were observed in the region of 960–1000 nm, corresponding to the formation of the radical ion pair  $(\mathbf{5}^{\bullet-} \rightarrow \mathbf{ZnNc}^{\bullet+})$ . These absorption bands show quick rise-decay behavior, indicating a rapid CR process.

The transient absorption spectra in coordinating solvents such as BN were quite different. Upon excitation of the dyad with 650 nm laser light, the spectra exhibited bands at 600 and 750 nm, which were attributed to appreciable population of  ${}^3ZnNc^*$  and  ${}^3S^*$ . These absorption bands of the triplet states decay, accompanied by the rise of the absorption bands in the 960–1000 nm region, which were attributed to the formation of the radical ion pairs,  $ZnNc^{\bullet+}$  and  $S^{\bullet-}$ . These data suggest that in a coordinating solvent the ET process takes place from  ${}^3ZnNc^*$  to  $S^*$ . The bimolecular ET rate constant ( $S^*$ ) was evaluated to be  $S^*$ . While the back ET rate constant ( $S^*$ ) was determined to be  $S^*$ .

Schuster and co-workers [117] also developed and studied a dyad system composed of a pyridine-appended  $C_{60}$  that axially ligates to the central metal of ZnTPP with a linear geometry (Fig. 19). Therefore, the donor–acceptor distance in the dyad is greatly increased. Due to the increased electron density around the pyridine nitrogen, it was shown to have a greater capacity to bind to the zinc central metal. This was confirmed by steady state fluorescence quenching experiments where the K value was determined to be  $7.4 \times 10^4 \,\mathrm{M}^{-1}$  in o-DCB. Also, these experiments show that fluorescence quenching occurred quite efficiently in both polar and nonpolar solvents.

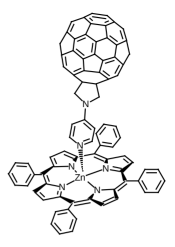


Fig. 19. Structure of the investigated zinc porphyrin-fullerene dyad.

Nierengarten and co-workers studied a similar dyad [115]. This dyad was composed of a pyridine-appended C<sub>60</sub> molecule (py~C<sub>60</sub>) synthesized by Bingel addition to the fullerene, which can axially coordinate to ZnTPP via the central metal atom (Fig. 20a). Steady state fluorescence was used to arrive at an association constant of  $K_a = 3000 \pm 4001 \mathrm{M}^{-1}$  for the complex in toluene. Upon excitation of ZnTPP in the presence of py $\sim C_{60}$ , the fluorescence of ZnTPP was quenched. This quenching was rationalized to occur as a result of the CS process from ZnTPP to py $\sim$ C<sub>60</sub>, owing to the very large rate constant observed ( $k_{\rm CS} > 5 \times 10^{10} \, {\rm s}^{-1}$ ). Also, the nonpolar nature of the solvent contributes to the observation of the EN process, since the CS state is unstable in nonpolar media relative to polar solvent media. EN processes in the system were found to have a slow and fast component. The fast component was thought to exist as a result of singlet-singlet energy transfer in the associated complex, while the slow component occurs via triplet-triplet energy transfer for the unassociated complex [115].

Other coordinated porphyrin–fullerene dyad systems have been developed to increase the donor–acceptor distance and thereby probe its effect on energy or electron transfer events. Guldi, Hirsch, and coworkers [74] developed a dyad system composed of a heterofullerene (C<sub>59</sub>N) appended with a pyridine moiety capable of axial ligation to the central metal of a zinc tetrakis(*p-tert*-butylphenyl)porphyrin entity (ZnTBPP) (Fig. 20b). In this system the pyridine linkage to C<sub>59</sub>N has an almost linear geometry, and thus the donor–acceptor distance is greatly increased relative to those systems previously discussed in this review.

Steady-state fluorescence data were used to determine the K value of the dyad (ZnTBPP $\leftarrow$ py $\sim$ C<sub>59</sub>N), which was around 10,000 M $^{-1}$  in non-coordinating solvents (toluene, o-DCB). Upon addition of increasing amounts of the C<sub>59</sub>N to ZnTBPP, the fluorescence of ZnTBPP was steadily quenched. Time-resolved fluorescence decay experiments were performed to probe the nature of the dyad

Fig. 20. Structure of axially coordinated zinc porphyrin:pyridine-appended fullerene conjugates.

excited state. Upon laser excitation at 337 nm, the decay profile displayed both a short-lived and long-lived component, with lifetimes of 0.15 and 1.9 ns, respectively. These experiments also revealed emission of  $C_{59}N$  at around 825 nm, indicating that the excited state of  $C_{59}N$  is being populated.

Transient absorption experiments were performed to determine the nature of the photochemical reactions for the studied dyad. Upon addition of the axially ligating py $\sim$ C<sub>59</sub>N to ZnTBPP in toluene and laser light excitation at 535 nm,  $^1$ ZnTBPP\* decayed very rapidly, with a lifetime of about 2.3 ns. Peaks appeared in the transient spectra at 735 and 940 nm. This indicates that, due to the rapid decay of the excited state, the deactivation takes place from  $^1$ ZnTBPP\* and populates C<sub>59</sub>N via a singlet–singlet energy transfer mechanism. The  $^1$ C<sub>59</sub>N\* state subsequently decayed to  $^3$ C<sub>59</sub>N\* via the ISC process with a rate constant of 1.0  $\times$  10<sup>9</sup> s $^{-1}$  which then decays further to the ground state, with a lifetime of 15  $\mu$ s.

However, in a more polar solvent such as *o*-DCB, the transient absorption spectra of the dyad were quite different. After laser light excitation of ZnTBPP (18 ps), <sup>1</sup>ZnTBPP\* decays, rapidly revealing peaks at 640 and 1020 nm, which correspond to ZnTBPP\*+ and C<sub>59</sub>N\*-¬, respectively. These data indicate that the photoinduced ET is the predominant photochemical reaction process in polar medium.

### 3.2. Two-point binding supramolecular triads with electron donor

Our research groups recently prepared and studied a novel triad system composed of a zinc porphyrin appended with hydrogen-bonding groups such as either a carboxylic acid or an amide group (ZnTPP $\sim$ COOH or ZnTPP $\sim$ NH<sub>2</sub>) and a C<sub>60</sub> molecule appended with a pyridine group and a

N,N-dimethylaniline (DMA) group acting as a secondary electron donor ( $C_{60}\sim$ DMA) (Fig. 21) [118]. The triad system is self-assembled via a "two-point" binding motif, where the pyridine group on the  $C_{60}$  axially ligates to the central metal of the zinc porphyrin, while the nitrogen of the fulleropyrrolidine ring hydrogen bonds with the hydrogen-bonding group attached to ZnTPP, either ZnTPP $\sim$ COOH or ZnTPP $\sim$ CONH<sub>2</sub>.

UV-Vis,  $^1$ H NMR, and ab initio B3LYP/3–21G(\*) computational studies were used to verify the integrity of the self-assembled triads. Upon addition of  $C_{60}\sim$ DMA to a solution containing either ZnTPP $\sim$ COOH or ZnTPP $\sim$ CONH<sub>2</sub>,

Fig. 21. Structure of the zinc porphyrin–fullerene-dimethylaminophenyl supramolecular triad bound by "two-point" hydrogen bonding-coordinate bonding.

a characteristic decrease and red shift of the Soret band were observed. Also, upon addition of either ZnTPP $\sim$ COOH or ZnTPP $\sim$ NH<sub>2</sub> to a solution of C<sub>60</sub> $\sim$ DMA, the pyridine and fullerpyrrolidine protons experienced an upfield shift caused by interaction with the ring current of the porphyrin ring, while the amide and carboxylic acid protons experienced a downfield shift due to hydrogen bonding interactions. These results clearly indicated the formation of the supramolecular triad system via the "two-point" binding motif. The K values for the triads C<sub>60</sub> $\sim$ DMA $\rightarrow$ ZnTPP $\sim$ COOH and C<sub>60</sub> $\sim$ DMA $\rightarrow$ ZnTPP $\sim$ COOH<sub>2</sub> were arrived at via Scatchard plots and were determined to be  $10 \times 10^4$  and  $3.1 \times 10^4$  M $^{-1}$ , respectively.

Steady-state fluorescence experiments were carried out on the supramolecular triad systems. Upon addition of  $C_{60}\sim DMA$  to a solution containing either ZnTPP $\sim COOH$  or ZnTPP $\sim NH_2$  in o-DCB, the fluorescence emission intensity of the ZnTPP moiety was quenched to about 30% of its original value. Also, a weak band at 710 nm, corresponding to the emission of  $C_{60}$ , was observed. Compared with the results of a control experiment where  $C_{60}\sim DMA$  was added to ZnTPP, the observed fluorescence showed that the amount of  $C_{60}\sim DMA$  needed to quench emission by the same extent as with ZnTPP $\sim COOH$  or ZnTPP $\sim NH_2$  was about 25% less. These data indicate that the efficiency of the charge separation process is increased as a result of the stronger "two-point" binding.

Time-resolved fluorescence emission experiments were also performed on the self-assembled triad systems. The singlet excited state lifetimes of ZnTPP, ZnTPP $\sim$ COOH, and ZnTPP $\sim$ CONH<sub>2</sub> were determined to be 1.92, 2.35, 1.97 ns, respectively. Upon complexation with C<sub>60</sub> $\sim$ DMA, fast fluorescence decay was observed, indicating that the CS process from the singlet excited states occurred. The fluorescence decay was found to be more efficient for the "two-point" bound systems as compared with the singly bound counterpart. These data indicate that the DMA moiety acts as a secondary electron donor and accelerates the CS process (Table 11).

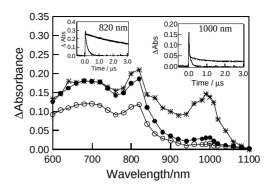


Fig. 22. Transient absorption spectra obtained by 532 nm laser light of  $C_{60}NDMA \rightarrow ZnTPP \sim CONH_2$  (0.1 mM, 1:1 eq.) in o-DCB: (\*\*) 0.01  $\mu$ s, ( $\bullet$ ) 0.25  $\mu$ s and ( $\bigcirc$ ) 2.5  $\mu$ s. Inset: time profiles of the 820 and 1000 nm bands [118].

Transient absorption spectra were obtained to determine the nature of the excited state photochemical reactions for the self-assembled dyads (Fig. 22). The transient absorption spectra of  $C_{60}\sim DMA:ZnTPP\sim NH_2$  obtained after 532 nm laser light flash after 0.01  $\mu$ s exhibited bands at 700, 870, and 1000 nm corresponding to  $^3C_{60}^*$ ,  $^3ZnTPP\sim NH_2^*$ , and  $C_{60}^{\bullet-}$ , respectively. The  $C_{60}^{\bullet-}$  band at 1000 nm was still observed in the transient absorption spectra after 0.25  $\mu$ s. This indicates that the DMA group aids in the stabilization of the CS state. The CS state exhibited both a fast ( $k=3.1\times 10^7~{\rm s}^{-1}$ ) and slow ( $k=3.5\times 10^5~{\rm s}^{-1}$ ) component for the CR process.

The relatively strong binding in the "two-point" bound triad system is thought to play a role in the increased stabilization of the CS state. A comparison of the ratio  $k_{\rm CS}/k_{\rm CR}$ , which reflects the ET efficiency, shows that as the K values increase for the "two-point" bound system, so does  $k_{\rm CS}/k_{\rm CR}$ . This indicates that for the more strongly bound systems the efficiency of the ET process increases. Also, the DMA group is thought to play a role in slowing down CR and thereby increasing the lifetime of the CS state for the triad system. Since the  $E_{\rm ox}$  value of the DMA group is relatively low,

Table 11 Binding constants (K), fluorescence lifetimes ( $\tau_f$ )<sup>a</sup>, CS rate constants ( $k_{CS}^{singlet}$ ), quantum yields ( $\Phi_{CS}^{singlet}$ ) and CR rate constants ( $k_{CR}$ ) for N,N-dimethylaminophenyl-zinc porphyrin  $\leftarrow$  C<sub>60</sub> triads in o-DCB

Complex	K (M <sup>-1</sup> )	τ <sub>f</sub> <sup>a</sup> (fraction %)	$k_{\rm CS}^{\rm singlet}$ (s <sup>-1</sup> )	$\Phi_{\mathrm{CS}}^{\mathrm{singlet_b}}$	$k_{\rm CR} \ ({\rm s}^{-1})$	$k_{\rm CS}/k_{\rm CR}$
$C_{60}$ py $\rightarrow$ ZnTPP	$0.77 \times 10^4$	1.85 ns (100)	$6.3 \times 10^{7}$	0.12	$3.0 \times 10^{7}$	2
$C_{60}{\sim}DMA{\rightarrow}ZnTPP$	$0.78\times10^4$	71 ps (27) 2.03 ns (73)	$1.4 \times 10^{10}$ b $1.5 \times 10^{8}$ c	0.97 <sup>b</sup> 0.22 <sup>c</sup>	$3.4\times10^7$	4
$C_{60}\sim$ DMA $\rightarrow$ ZnTPP $\sim$ CONH <sub>2</sub>	$3.1\times10^4$	69 ps (81) 1.60 ns (19)	$1.4 \times 10^{10b}$ $1.8 \times 10^{9c}$	0.98 <sup>b</sup> 0.77 <sup>c</sup>	$3.1\times10^7$	58
$C_{60}\sim$ DMA $\rightarrow$ ZnTPP $\sim$ COOH	$10\times10^4$	8 ps (75) 2.28 ns (25)	$1.3 \times 10^{11b}$ $1.8 \times 10^{9c}$	0.99 <sup>b</sup> 0.67 <sup>c</sup>	$2.3\times10^7$	43

<sup>&</sup>lt;sup>a</sup> The singlet lifetimes of ZnTPP, ZnTPP-CONH<sub>2</sub>, ZnTPP-COOH in deareated *o*-DCB were found to be 1.92, 1.97, and 2.35 ns (mono-exponential decay), respectively.

<sup>&</sup>lt;sup>b</sup> From the fast decay.

<sup>&</sup>lt;sup>c</sup> Average values.

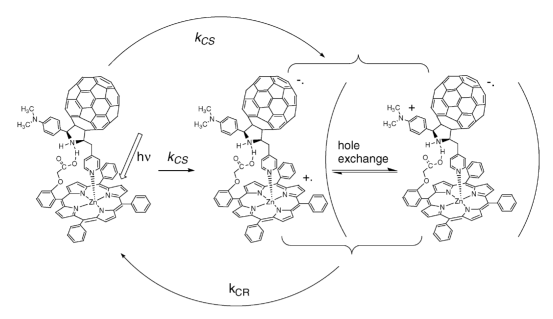


Fig. 23. Proposed mechanism of the photochemical charge stabilization in the self-assembled DMA $\sim$ C<sub>60</sub>Py $\rightarrow$ ZnP $\sim$ COOH triad.

after the initial charge separation generating  $ZnTPP^{\bullet+}$  and  $C_{60}^{\bullet-}$ , the DMA group donates an electron to  $ZnTPP^{\bullet+}$ . Thus, a hole shift occurs from  $ZnTPP^{\bullet+}$  to the DMA group, which is stable owing to the irreversible nature of the DMA oxidation, as shown in Fig. 23. These processes are thought to be the mechanism of the slow CR that is observed.

### 3.3. Fullerene-porphyrin coordinated systems: control over distance and orientation

The relative distance and orientation between an electron donor and acceptor in a dyad-type system greatly affect electron transfer rates and lifetimes. Recently, researchers have been developing dyad systems, where the distance and orientation of the donor and acceptor can be controlled, thereby allowing for their effects on the ET processes to be studied [33,36,48–52]. In this regard, we recently prepared an exotic dyad system composed of ZnTPP covalently linked with C<sub>60</sub> appended with a pyridine ring [119]. The pyridine ring (4-pyridyl or 3-pyridyl) on C<sub>60</sub> can either exist in the bound ("tail-on") or unbound ("tail-off") form by axial ligation to the ZnTPP (Fig. 24). The relative donor–acceptor proximity was varied by adjusting temperature or the concentration of a competing ligand (3-picoline). It was observed at room temperature that the Soret bands of the studied dyads were located around 433 nm, which is close to that of a

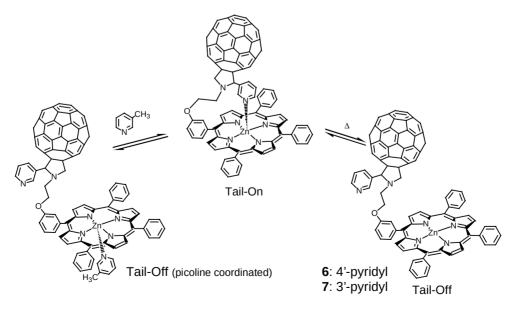


Fig. 24. Structure of zinc porphyrin-fullerene dyads utilizing 'tail-on' and 'tail-off' binding processes.

Table 12 Equilibrium constant (*K*) and thermodynamic parameters for the temperature-controlled 'tail-on' and 'tail-off' processes of zinc porphyrin–C<sub>60</sub> dyads in *o*-DCB

Dyad <sup>a</sup>	K <sup>b</sup> (M <sup>-1</sup> )	$\Delta G$ (kJ mol <sup>-1</sup> )	$\Delta H$ (kJ mol <sup>-1</sup> )	$\frac{\Delta S}{(\text{J K}^{-1} \text{ mol}^{-1})}$
7	17.29	-7.07	-19.17	-40.59
6	14.43	-6.62	-18.11	-38.52
$ZnP_m{\sim}py^c$	2.99	-2.72	-45.36	-138.6

<sup>&</sup>lt;sup>a</sup> For abbreviations see Fig. 24.

penta-coordinated ZnTPP-pyridine complex. Therefore, it was assumed that the dyads are predominantly in the "tail-on" form at room temperature. The *K* values for the dyads and thermodynamic parameters were determined by the Van't Hoff method using variable temperature UV-Vis spectral data. These data suggest that there are both entropic and enthalpic contributions to the overall binding process (Table 12).

Steady-state fluorescence experiments were performed on the dyads using the addition of picoline as a competing ligand to vary the amount of the "tail-on" and "tail-off" forms present in solution. Upon addition of excess picoline (10 eq.) to shift the equilibrium to the "tail-off" form, the emission intensity of the ZnTPP moiety increased by about 27%. This fluorescence emission behavior was most likely due to the increased distance between the donor and acceptor in the "tail-off" form and minor structural changes to the system.

Picosecond time-resolved fluorescence experiments were performed to investigate the nature of the excited state for the investigated dyads. In the case of the 4-pyridyl-substituted derivative, the "tail-on" form slightly accelerates the CS process. The reverse is observed for the 3-pyridyl-substituted derivative, suggesting that, in this case, the "tail-off" form facilitates structural changes that promote rapid CS process.

Nanosecond transient absorption spectra were recorded to elucidate the nature of the CS state. Bands corresponding to  ${}^{3}C_{60}^{*}$ ,  ${}^{3}ZnTPP^{*}$ , and  $C_{60}^{\bullet-}$  appeared in the transient absorption spectra at 700, 850, and 1000 nm, respectively. The triplet bands decayed with a lifetime of around 600 ns. The decay profile exhibited both fast and slow decay components indicating that there are two types of decay processes from the CS state. This was attributed to relaxation from the long-lived triplet CS state and also from the singlet CS state. Upon addition of picoline, a peak at 1000 nm was observed, which is ascribed to  $C_{60}^{\bullet-}$ , confirming that the CS process occurs for the "tail-off" form of the dyads. Also, the relative intensity of the peak at 1000 nm versus the peak at 700 nm is greater for the "tail-off" form than for "tail-on" form. This indicated that the slow decay component from  ${}^3C_{60}{}^*$  was a relatively minor pathway for the decay of the "tail-off" form of the CS state. Both the "tail-on" and "tail-off"

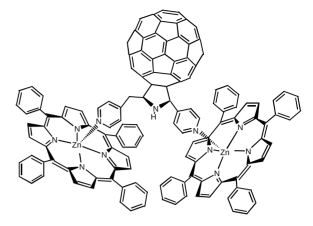


Fig. 25. Structure of the bis zinc porphyrin-fullerene triad self-assembled via axial coordination

forms undergo CS via <sup>1</sup>ZnTPP\*, although this process is slightly more efficient for the "tail-off" form of the dyads studied.

### 3.4. Fullerene-bisporphyrin coordinated triads

More recently we developed a system composed of a  $C_{60}$  unit appended with two pyridine units ( $C_{60} \sim py_2$ ) that axially ligate to two ZnTPP thus forming a self-assembled triad (Fig. 25) [120]. UV-Vis absorption data were used via the Scatchard method to determine the binding constant value for the system. This value was determined to be  $1.45 \times 10^4 \, \text{M}^{-1}$  which is considerably large as compared to the dyad counterpart analogue.

Steady-state fluorescence experiments were performed on the triad system. Upon addition of  $C_{60} \sim py_2$  to a solution of ZnTPP in o-DCB, the fluorescence was quenched to approximately 30% of its original intensity and red-shifted by 2–3 nm. The Stern–Volmer quenching constant ( $k_q$ ) was determined and found to be three times larger than that of a diffusion-controlled process, suggesting that intramolecular quenching is the primary quenching pathway for the system. Also, a weak emission band at 710 nm corresponding to the singlet emission of  $C_{60}$  was observed.

Time-resolved transient absorption spectra were recorded to confirm the nature of the excited state photochemical dynamics of the system. A solution of the triad in o-DCB was excited by laser light at 532 nm. Bands were observed at 760 and 850 nm corresponding to  ${}^{3}C_{60}^{*}$  and  ${}^{3}ZnTPP^{*}$ , respectively. Also, a weak band at 1000 nm corresponding to the  $C_{60}^{\bullet-}$  was observed. The quick rise and decay of this band indicates that charge separation takes place from the singlet excited state of ZnTPP.

The absorption spectra observed in coordinating solvents such as BN were quite different. Bands were observed at 770 and 840 nm, corresponding to  ${}^{3}C_{60}^{*}$  and  ${}^{3}ZnTPP^{*}$ . These bands were found to decay faster than what was observed for o-DCB. Also, a slow rise was observed in the 900–1100 nm region, corresponding to the formation of the  $C_{60}^{\bullet-}$ . These

b At 298.15 K.

<sup>&</sup>lt;sup>c</sup> Represents a pyridine-appended porphyrin.

data indicate that, in coordinating solvents, energy transfer from <sup>3</sup>ZnTPP\* to C<sub>60</sub> was the predominant process.

Another supramolecular triad was formed by a "covalent-coordinate" approach, where a free base porphyrin was functionalized to bear a  $C_{60}$  appended with a pyridine group ( $H_2TPP\sim C_{60}py$ ) capable of axial ligation with ZnTPP (Fig. 26) [80]. Steady state fluorescence experiments were performed on the supramolecular triad system. Upon addition of  $\bf 6$  or  $\bf 7$  to a solution containing ZnTPP, the two emission bands of ZnTPP at 598 and 646 nm were quenched, accompanied by the appearance of two new emission bands at 665 and 720 nm, corresponding to the emission of  $H_2TPP$ . These data indicate that an intramolecular quenching process is occurring within the self-assembled supramolecular triad system.

Picosecond time-resolved fluorescence spectral studies were performed on the covalently linked dyad systems. The short excited state lifetimes for the studied dyads (440 ps for 6 and 710 ps for 7) indicate that an intramolecular CS process takes place from  ${}^{1}H_{2}TPP^{*}$ . The  $k_{cs}$  values for the dyads were found to be  $2.2 \times 10^9 \,\text{s}^{-1}$  for **6** and  $2.6 \times 10^9 \,\text{s}^{-1}$  for 7. Time-resolved fluorescence spectra were also obtained for the self-assembled triad system with ZnTPP. For the triad systems, the fluorescence quenching of <sup>1</sup>ZnTPP\* was slightly accelerated; the lifetimes of <sup>1</sup>ZnTPP\* of **6**:ZnTPP and 7:ZnTPP were found to be 1.78 and 1.83 ns, respectively, while the lifetime of uncoordinated ZnTPP was 2.10 ns. Nanosecond transient absorption spectral studies were performed to determine the nature of the excited state photochemical reactions in the dyad and triad systems. The relative efficiency of intermolecular ET was evaluated by monitoring the absorbance ratio of the transient absorption bands at 700 and 1000 nm  $(A_{1000 \text{ nm}}/A_{700 \text{ nm}})$ . For dyad systems 6 and 7, this ratio was determined to be 0.45 and 0.32, respectively. Upon addition of 6:1 equivalents of 6 or 7 to

Fig. 26. Structure of H<sub>2</sub>TPP~C<sub>60</sub>Py self-assembled to ZnTPP triads.

ZnTPP, under conditions where almost all ZnTPP is coordinated, the absorbance ratio  $(A_{1000\,\mathrm{nm}}/A_{700\,\mathrm{nm}})$  increased to 0.50 and 0.52, respectively. These data indicate that the CS efficiency increases upon coordination of ZnTPP to the fullerpyrollidine entity of the dyads. Fig. 27 illustrates the different photochemical events in the studied triads.

In a coordinating solvent such as BN, the transient absorption spectra were quite different. Bands at 1000 and 620 nm exhibited a slow rise in the spectrum, while bands at 750

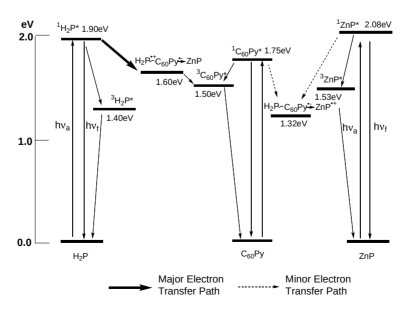


Fig. 27. Energy level diagrams showing the different photochemical events for supramolecular  $H_2TPP \sim C_{60}Py \rightarrow ZnTPP$  in o-DCB.

$$OC_{12}H_{25}$$
 $OC_{12}H_{25}$ 
 $OC_{12}H_{17}$ 
 $OC_{12}H_{17$ 

Fig. 28. Structures of crown ether complexed to fullerene bearing alkyl ammonium cation conjugate: (a) porphyrin and (b) phthalocyanine.

and 850 nm decayed at appreciable rates. These results suggest that, in coordinating solvents, intermolecular ET takes place from  ${}^{3}$ ZnTPP\*. The  $k_{\rm et}$  values were determined to be  $1.4 \times 10^{8}$  and  $1.9 \times 10^{8}$  M $^{-1}$  s $^{-1}$  for triads composed of **6** and **7**, respectively.

### 3.5. Fullerene-porphyrin/phthalocyanine assembly systems

Recently, Nierengarten and co-workers developed a system composed of a C<sub>60</sub> molecule appended with a quaternary ammonium unit and a free base porphyrin (H<sub>2</sub>TPP) functionalized to bear a crown ether unit (Fig. 28a) [121]. The two different entities self-assemble in solution via interaction between the quaternary ammonium and crown ether subunits. Steady-state fluorescence was used to determine the K value for the dyad system. The Benesi-Hildebrand method was applied to the data to determine a binding constant of about  $375,000 \,\mathrm{M}^{-1}$  for the system. This value is two orders of magnitude higher than those of most coordinating systems previously studied. This very strong binding was rationalized by the presence of  $\pi$ – $\pi$  stacking interactions between the porphyrin and C<sub>60</sub> units, in addition to hydrogen bonding. Evidence for  $\pi$ – $\pi$  stacking was provided by a <sup>1</sup>H NMR experiment where, upon addition of the C<sub>60</sub> derivative to the crown ether appended porphyrin, the β-pyrrole protons of the porphyrin ring experienced an upfield shift. These data indicate that the porphyrin ring must be interacting with the  $C_{60}$  spheroid. Also, steady-state fluorescence experiments indicate that the presence of the C<sub>60</sub> quaternary ammonium derivative quenches the singlet excited state of the porphyrin crown ether moiety efficiently.

Guldi et al. [122] prepared and studied self-assembled supramolecular dyad and triad systems composed of one or two phthalocyanine units (ZnPc or ZnPc–ZnPc) appended with a dibenzo-24-crown-8 unit capable of hydrogen bonding with a  $C_{60}$  molecule appended with a tertiary ammonium group ( $C_{60}$  amm) (Fig. 28b). Steady-state fluorescence

data were used to obtain the K values for the studied systems. These values were determined to be  $1.4 \times 10^4 \, \text{M}^{-1}$  for the dyad system (ZnPc–C<sub>60</sub>amm) and  $1.9 \times 10^4 \, \text{M}^{-1}$  for the triad system (ZnPc–C<sub>60</sub>amm–ZnPc).

Time-resolved fluorescence experiments were employed to determine the nature of the excited state photochemical reactions for the dyad and triad systems. The two pristine ZnPc derivatives exhibited an excited state lifetime of around 3.1 ns. Upon addition of  $C_{60}$ amm, a short-lived component with a lifetime of 0.28 ns became predominant, accompanied by depletion of the long-lived component. The fast decay of the excited state of  $^1\text{ZnPc}^*$  upon complexation with  $C_{60}$ amm suggests that intramolecular ET from  $^1\text{ZnPc}^*$  is the predominant quenching mechanism.

Transient absorption spectral studies were performed to determine the nature of the species produced during the

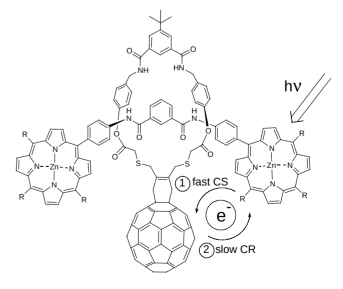


Fig. 29. Structure of the bis zinc porphyrin-fullerene rotaxane supramolecule.

course of the excited state photochemical reaction. Upon excitation with a 8 ns laser light flash, after 50 ns, bands at 1040 nm corresponding to  $C_{60}$  amm $^{\bullet-}$  and bands at 500 and 860 nm corresponding to  $ZnPc^{\bullet+}$  were observed. These data indicate that the formation of SSIP occurs with a high  $\Phi_{cs}$  value of about 0.90 and a lifetime ( $\tau_{cs}$ ) of 1.5  $\mu$ s.

Recently, Watanabe et al. [123] reported intrarotaxane electron transfer between ZnTPP and  $C_{60}$ -crown ether in a supramolecular system shown in Fig. 29. The time profile of the fluorescence decay of the rotaxane revealed a biexponential decay in BN. From this data, efficient CS ( $k_{cs} = 10^{10} \, \mathrm{s}^{-1}$ ) from  $^{1}$ ZnTPP\* to the fullerene entity was calculated. The evaluated rate of CR from nanosecond transient absorption spectral studies by monitoring the decay of  $C_{60}^{\bullet-1}$  was found to be  $5.5 \times 10^{6} \, \mathrm{s}^{-1}$ . This value corresponded to a lifetime  $\tau_{\rm RIP}$  of 180 ns for the CS state (radical ion pair). The observed slow CR suggested that this process was located in the Marcus-inverted region far more negative than the CS process.

### 4. Summary

The intermolecular ET processes from electron donors (porphyrins, chlorophyll, phthalocyanines, and naphthalocyanines) and their metal derivatives to electron acceptors, fullerenes (C<sub>60</sub>/C<sub>70</sub>) studied by nanosecond and picosecond laser flash photolysis techniques in polar and nonpolar solvents have revealed many interesting features. In polar solvents, the ET process takes place via the triplet excited states of the excited acceptor or excited donor, yielding solvated radical ion pairs. Because of the bimolecular encounter of solvated radical ion pairs, the back ET rates are found to be generally slow. The structure-photochemical reactivity probed in the intermolecularly interacting systems revealed dependence on the nature of the porphyrin/phthalocyanine macrocycle, the metal ion present in the porphyrin/phthalocyanine cavity, electron donor substituents on the macrocycle periphery, and the polarity of the solvent medium.

The porphyrins, phthalocyanines and fullerenes have been found to be excellent building blocks for supramolecular systems for the study of photoinduced CS reactions by using time-resolved ultrafast spectroscopic techniques. In the supramolecular conjugates formed by axial coordination, hydrogen bonding, crown-ether complexation, or rotaxane formation, the CS process occurs mainly from the singlet excited state of the donor. In contrast to the intermolecular back ET process, the back CR rates are found to be fast. In some of the conjugates studied, the predicted acceleration of the CS process and deceleration of the CR process have been clearly observed, mainly due to the small reorganization energies of fullerenes in electron transfer reactions. The photophysical properties of the porphyrin and fullerene moieties are shown to be tuned in a controlled manner upon coordination of metal centers. The nature of the linker between the donor and acceptor entities, the solvent, and the metal ions in the porphyrin cavity influences the overall photochemical reactivity. Studies on self-assembled supramolecular triads, tetrads, etc., are only in the beginning stages, and future studies will be anticipated to involve more complex systems targeted for better charge stabilization and also to perform specific light-driven photochemical processes. The supramolecular approach of building fullerene–porphyrin and fullerene–phthalocyanine conjugates is beginning to provide well-characterized donor–acceptor systems, which could eventually be used for the development of solar energy harvesting and opto-electronic devices such as sensors, switches, gates, etc.

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