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Intramolecular charge transfer character in tetrathiafulvalene-annulated porphyrinoids: Effects of core modification and protonation†

Ramababu Bolligarla,a Masatoshi Ishida,a,b Vijayendra S. Shetti,a Kazuhisa Yamasumi,b Hiroyuki Furuta[b] and Chang-Hee Lee*a,

Synthesis, characterization, photophysical and electrochemical properties of novel tetrathiafulvalene (TTF)-annulated core-modified porphyrin (1) and its expanded rubyrin analogue (2) are described. The sulfur core modifications effected in 1 and 2 allow a feasible intramolecular charge transfer from the TTF fragments to central conjugated core as inferred from comparative spectroscopic and electrochemical measurements. DFT calculations also support the intramolecular charge transfer nature of 1 and 2 upon excitation. Further electronic perturbation of the TTF-annulated porphyrins was achieved by protonation, giving rise to drastic change in the optical feature with an extremely low energy band in the NIR region. The pronounced electron acceptor ability of the macrocyclic core of the dicationic species (H$_2$TTF$^{2+}$ and H$_2$2$^{2+}$) resulted in the thermally excited electron transfer occurring at room temperature as elucidated by EPR spectroscopy.

Introduction

Tetrathiafulvalene (TTF)-based molecular materials have received tremendous attentions over the years because of the excellent physical nature such as superconductivity, redox, magnetic and photophysical properties. Extensive applications with electron donor-acceptor conjugates including nonlinear optical materials, photovoltaic solar cells, organic field-effect transistors and metal cation sensors have been thus developed. A TTF molecule is indeed known to be involved in rich redox chemistry; TTFs can exist in any of three distinct stable redox states such as TTF$^0$, TTF$^+$ and TTF$^{2+}$. Over the past decade, the synthetic TTF derivatives hybridized with other π-electron rich acceptor molecules such as calixpyrroles and porphyrins would thus be expected to create intriguing systems with an essence of designed donor-acceptor (D-A) scaffolds.

The inherent charge transfer character present in a various TTF-annulated porphyrins (e.g., 3-6) conjugated through their β-pyrrolic positions has been investigated to gain insight into the structure-property relationship by varying the number and position of TTF fragments on the conjugated core as well as the size of π-network of the acceptor unit (Chart 1). Such donor-acceptor annulated ensembles reported previously have been thought to allow stronger electronic coupling interactions between the donor and acceptor components due to the controlled structural natures; compact distances, parallel orientations and rigidified skeletons. For further fine tuning of the photo-induced charge transfer process, reduction potentials of acceptor (porphyrin) units seem to be critical variables.

The core-modifications of porphyrins and expanded porphyrins by replacing core NH with other hetero atoms (S, Se, Te and O) can alter their inherent structural and electronic properties, while the basic aromatic nature is preserved as similar to their parent pyrrolic congeners. The overall photophysical (e.g., reasonably larger emission quantum yields and longer excited state dynamics) and electrochemical (e.g., facile reduction potentials) properties are expected to function as good acceptor units in these D-A ensembles. Furthermore, the sulfur core modified analogues has been shown the good photochemical stabilities as well as suitable HOMO and LUMO energy levels.

In this study, hybrid macrocycles composing of photocative core-modified porphyrinoids and electroactive TTFs fused at the pyrrole positions, namely TTF-annulated dibithiaporhyn (1) and tetrathiarubyrin (2) were synthesized and characterized as the first examples of donor-acceptor TTF-annulated, core modified porphyrins (Chart 1). The intramolecular charge transfer character from the TTF fragments to the porphyrin core was examined by comparable spectroscopic techniques and electrochemical measurements. The visualization of their electron flow upon an excitation was shown by DFT-based theoretical calculations. Furthermore, protonation, known as the ubiquitous phenomenon in the nitrogen-incorporated
porphyrinoids, drastically altered their electronic properties of the TTF-core modified porphyrins. Due to the enhanced electron acceptor ability of the macrocyclic core of the dicationic species (H$_3$T$^{2+}$ and H$_2$T$^{2+}$), an intermolecular thermal electron transfer occurred at room temperature along with emergence of lower energy NIR absorptions.

![Chart 1](image)

**Experimental Section**

**Instruments.** $^1$H-NMR (300 MHz) and $^{13}$C-NMR (100 MHz) spectra were recorded on a Bruker spectrometer. Chemical shifts (δ-scale, ppm) were referenced to the residual solvent peaks (δ of 5.32 for proton and 53.8 for carbon for dichloromethane). MALDI-TOF mass spectra were recorded on a Voyager-DE STR spectrometer using dithranol (1,8,9-trihydroxyanthracene) as a matrix. High-resolution mass spectra (HR-MS) were recorded on a Bruker spectrometer. Chemical shifts (δ-scale, ppm) were referenced to the residual solvent peaks of 5.32 for proton and 53.8 for carbon for dichloromethane.

Synthesis of C$_8$S-TTF-pyrrole: 4,5-Bis(hexylthio)-1,3-dithiole-2-thione (1.89 g, 5.154 mmol) and 5-tosyl-5H-[1,3]dithiolo[4,5-c]pyrrol-2-one (0.7 g, 2.248 mmol) were suspended in triethylposphite (15.0 mL) and the reaction mixture was refluxed for 20 min. The yellow reaction mixture was refluxed for 20 min. Then cooled to room temperature, and then filtered the cold solution. The obtained yellow solid was washed with hexane for several times to remove an impurity of symmetrical TTF compound. The compound can be used directly without any further purification. Yield: 1.05 g, ca. 74%.

$^1$H NMR (300 MHz, CDCl$_3$): δ 0.88 (t, 6H, CH$_3$), 1.25-1.33 (m, 8H, CH$_2$), 1.34-1.44 (m, 4H, CH$_2$), 1.56-1.66 (m, 4H, CH$_2$), 2.42 (s, 3H, ArCH$_3$), 2.80 (t, 4H, SCH$_2$), 6.93 (s, 2H, α-H), 7.30 (d, J = 8.0 Hz, 2H, Ar-H), 7.73 (d, J = 8.0 Hz, 2H, Ar-H) ppm.

A suspension of tosyl protected TTF pyrrole (0.4 g, 0.635 mmol) in anhydrous THF-MeOH (1:1 v/v, 20 mL) was degassed by N$_2$ for 15 min before addition of sodium methoxide (30% solution in MeOH, 2.35 mL) in one portion. The yellow reaction mixture was refluxed for 20 min. Then cooled to room temperature, and then filtered the cold solution. The obtained yellow solid was washed with hexane for several times to remove an impurity of symmetrical TTF compound. The compound can be used directly without any further purification. Yield: 1.05 g, ca. 74%.

**Density Functional Theory (DFT) Calculations.** Theoretical calculations were performed with the Gaussian09 program suite using a supercomputer. All calculations were carried out using the density functional theory (DFT) method with Becke’s three-parameter hybrid exchange functionals and the Lee-Yang-Parr correlation functional (B3LYP) employing the 6-31G(d) basis set for all atoms.

Chart 1. Chemical structures of TTF-annulated porphyrins and the higher derivative.
General Synthetic procedure for TTF-annulated porphyrins
A dichloromethane solution (360 mL) of the corresponding thiophene (8) and bithiophene (9) diols (0.5 mmol), respectively, and TTF annexulated pyrrole (7) (0.5 mmol) was purging with N₂ gas before adding acid catalyst, BF₃·Et₂O (0.25 mmol). The reaction mixture was stirred for 1 hour. DDQ (1.5 mmol) was then added to the mixture and stirred further for 30 min in air. Triethylamine (0.3 mL) was added to quench the reaction. The crude residues thus obtained were recrystallized in MeOH solvent by sonication and filtered. The desired porphyrin compound 1 was purified over silica gel eluted with CH₂Cl₂, and rubyrin compound 2 was eluted chloroform:hexane (1:1) on basic alumina column chromatography. The samples for experiments were further purified by recrystallization in CHCl₃/MeOH solvents.

TTF-annulated dithiaporphyrin 1: Yield: 28%; ³¹H NMR (300 MHz, CDCl₃): δ 0.90 (t, J = 7.0 Hz, 12H, CH₃), 1.26-1.32 (m, 16H, CH₂), 1.38-1.47 (m, 8H, CH₂), 1.59-1.69 (q, J = 7.4 Hz, 8H, SCH₂), 2.76 (s, 12H, ArCH₃), 2.82 (t, J = 7.4 Hz, 8H, SCH₂), 7.66 (d, J = 7.8 Hz, 8H, Ar-H), 7.94 (d, J = 7.8 Hz, 8H, Ar-H), 9.48 (s, 4H, β-H); ¹³C NMR (100 MHz, CDCl₃) δ 147.8, 147.3, 141.8, 139.2, 136.8, 135.8, 132.8, 132.5, 132.1, 129.5, 129.3, 127.8, 36.3, 31.3, 29.7, 28.2, 22.5, 21.7, 14.0 ppm. HRMS (FAB) m/z = 1520.2729 (found), 1520.2724 (Calcd. for C₅₀H₃₆N₈S₄). Error: +0.3 ppm. UV/Vis [in Toluene, λmax/nm (log ε)]: 545 (5.54), 521 (4.62), 572 (3.66), 636 (3.63), 693(3.65).

TTF-annulated tetrathiarubyrin 2: Yield: 13%; ¹³H NMR (400 MHz, CDCl₃): δ 0.91 (t, J = 7.0 Hz, 12H, CH₃), 1.28-1.35 (m, 16H, CH₂), 1.42-1.50 (m, 8H, CH₂), 1.65-1.84 (q, J = 7.4 Hz, 8H, CH₂), 2.88 (t, J = 7.4 Hz, 8H, CH₂), 2.92 (s, 12H, ArCH₃), 7.85 (d, J = 7.6 Hz, 8H, Ar-H), 8.24 (d, J = 7.6 Hz, 8H, Ar-H), 10.26 (d, J = 5.0 Hz, 8H, β-H), 11.49 (d, J = 5.0 Hz, 8H, β-H); ¹³C NMR (100 MHz, CDCl₃) δ 144.0, 141.7, 141.2, 139.3, 138.4, 135.5, 133.2, 129.9, 129.3, 127.9, 124.8, 106.24, 36.4, 31.3, 29.8, 28.3, 22.5, 21.9, 14.0 ppm. HRMS (FAB) m/z = 1684.2478 (found), 1684.2479 (Calcd. for C₅₀H₃₆N₈S₄). Error: −0.1 ppm. UV/Vis [in Toluene, λmax/nm (log ε)]: 545 (6.02), 670 (4.61), 731 (4.91), 846 (4.16).

Results and Discussion

Synthesis and characterization of 1 and 2. The TTF-annulated dithiaporphyrin 1, and tetrathiarubyrin 2 were prepared by conventional one-pot condensation of TTF-annulated pyrrole (7) with the appropriate carbonyl (i.e., 2,5-bis(3-tolyl)hydroxymethyl)thiophene (8) or 5,5′-bis(3-tolyl)hydroxymethyl-2,2′-bithiophene (9) in the presence of boron trifluoride and followed by 2,3-dichloro-5,6-dicyano-p-benzoquinone (DDQ) oxidation. Macrocycles 1 and 2 were well characterized by spectroscopic means including NMR, Mass spectrometry and HPLC techniques (cf. ESI).

In the ¹H NMR spectra, the signal corresponding to β-thiophene-Hs appeared as a singlet at δ 9.48 ppm for 1 and as two doublets at 11.49 (J = 5.0 Hz) and 10.28 (J = 5.0 Hz) ppm for 2 are seen, which are indicative of typical aromatic character due to the diatropic ring current of the macrocycles (Fig. S2 and S5). In particular, it is concluded that a highly symmetric and the planar rectangle conformation of the core of 2 is thus present in solution as considering the characteristic NMR resonance patterns of the unsubstituted rubyrin (S₄TTR). In order to gain the structural insights of the macrocycle, the density functional theory (DFT) calculations were carried out by using the B3LYP/6-31G(d) levels. Compared to the structures of the unfunctionalized derivatives, S₂TTP and S₃TTR, the relatively coplanar conformations for 1 and 2 were demonstrated with mean deviation values (defined by porphyrin core atoms) of 0.087 and 0.332 Å, respectively (Fig. S10). The nucleus-independent chemical shift (NICS) values at their central values of the mean cores (e.g., −15.2 ppm for 1 and −14.1 ppm for 2) support the distinct aromaticity of 1 and 2, which are well consistent with the NMR spectroscopic observations.

Fig 1. Comparative UV-vis-NIR absorption spectra of the porphyrins between (a) 1 (red) and S₂TTP (black), and the rubyrins; (c) 2 (red) and S₃TTR (black) recorded in CH₂Cl₂. Insets show the magnified region of Q-band, respectively. Emission
Optical properties and DFT calculations. These 18 π- and 26 π-electron compounds, 1 and 2 exhibited the typical optical features in the UV-vis-NIR absorption spectroscopy. The redshifted Soret band and broad Q-bands tailing to the 1000 nm in 1 were seen in CH$_2$Cl$_2$ in comparison with those of the non-annulated S$_2$TPP (Fig. 1a). This is due to the strong electronic coupling between the porphyrin core and the peripheral TTF fragments. Meanwhile, a broad lowest energy band could be attributed to the intramolecular charge transfer transition. The core modification present in 1 affords overall redshifts of absorption bands compared to the original pyrrole-based congeners (e.g., $\lambda_{\text{Soret}}$ = 448 nm for 1 and $\lambda_{\text{Soret}}$ = approximately 430 nm for 3 and 4). The photo-induced charge transfer event occurring in 1 was also confirmed by using fluorescence spectroscopy. The overall fluorescence of 1 is largely quenched compared to that of S$_2$TPP, which is may involve the electron transfer from TTF to the excited state of porphyrin core, leading to the rapid excited intramolecular charge transfer. (Fig. 1b). In the polar media such as THF, the emission intensities of 1 and 2 were further quenched, indicating possibly acceleration of the electron transfer rate (Fig. S11). As considered that N$_2$ porphyrin derivatives 3 and 4 has no detectable luminescence under the ambient condition, the core modifications of the porphyrin skeleton would effect on the inherent molecular orbital interactions between the donor TTF moiety and acceptor porphyrin core in 1.

In the π-expanded system of TTF-rubyrin 2, due to the expansion of π-conjugation network, lower energy absorption bands are seen in the NIR region. The characteristic CT band originated from the TTF moieties to the rubyrin core unit was likewise appeared in the far NIR region (Fig. 1c). Consequently the fluorescence of 2 was quenched in comparison with S$_2$TTR as well (Fig. 1d).

The molecular orbital analysis (B3LYP/6-31G(d) levels) revealed that the electron densities of HOMO are localized on the TTF moieties whereas those of LUMOs are predominantly distributed on the whole macrocycle core for both cases, suggesting the presence of intramolecular charge transfer character (Fig. 2 and Fig. S12 and S13). The latter MO energetic trend can be interpreted by the classical Goutermann’s four orbital theory$^{16}$, and the broad CT band observed in the NIR region could thus be attributed to the HOMO-LUMO electronic transition. The time dependent (TD) DFT simulation of the electronic spectrum of 1 also supports the CT-based transition (oscillator strength, $f$: 0.0026) with the energy of 1.44 eV (Fig. S14a). These intramolecular interactions are found in those of TTF-annulated porphyrins 3 and 4.

Accordingly, the energy trend of MO diagram of the π-expanded derivative 2 demonstrated the similar MO fashion; the TTF-based HOMO and HOMO-1 as well as the macrocycle-based MOs with degenerated HOMO-2 and HOMO-3 pair and the LUMO pair were found to be considered. The forbidden transition is identified with the CT nature ($f$: 0.0048, 1.31 eV) by TD-DFT calculation (Fig. S14b).

Redox properties. In an effort to determine the driving force for the electron transfer processes, the electrochemical potentials of 1 and 2 were measured by cyclic voltammetry (CV) and differential pulse voltammetry (DPV) technique in the CH$_2$Cl$_2$ containing 0.1 M n-tetramethylammonium hexafluorophosphate (TBAPF$_6$) (Fig. 3). The compound 1 revealed two reversible one-electron oxidation waves located at −0.12 and 0.05 V (vs. ferrocene/ferrocenium couple) and a two-electron oxidation wave seen at 0.39 V in the anodic side. These peaks are corresponded to the TTF entities and the trans-symmetric orientation of the TTF moieties in 1 resulted in the separation of the first oxidation wave (for the process of TTF radical cation formation). This leads us to speculate the presence of electronic communication between the two TTF

![Fig. 2. Selected frontier molecular orbitals of (a) 1 and (b) 2 obtained by using B3LYP/6-31G(d) level calculations](image-url)
units as considering the electrochemical behaviours of 3 and 4.
In the reduction side, two reversible waves corresponded to the radical anion and dianion of the porphyrin core are seen, which are significantly anodic shifted in comparison with the pyrrolic reference, 3 ($E_{\text{red}} = -1.57$ and $-1.81$ V vs. Fe/Fe$^+$), whereas the oxidation potentials were identical. Accordingly, the narrower HOMO-LUMO electrochemical energy gap of 1.21 V in 1 reflected to the redshift of the optical bands. In the TTF-rubyrin system 2, the TTF-based oxidation waves peaked at $-0.05$ and $+0.33$ V are both two-electron processes. This implies that the longer spatial separation of TTF arrangement causes less electronic interactions (diameter between inter TTF units is approx.17 Å for 2 vs. 14 Å for 1). Further two oxidation waves originated from the radical cation and dication formations of the rubyrin core were observed at 0.59 and 0.89 V. The macroscopic reductions at $-1.18$ and $-1.34$ V (vs. Fe/Fe$^+$) were determined on the basis of the comparative electrochemical study of S$_2$TTR ($E_{\text{red}} = -0.86$, $-1.05$ vs. SCE). The further anodic shifts of the reduction potentials of 2 owing to the core expansion resulted in the narrow HOMO-LUMO gap of 1.13 V.

**Photoinduced charge transfer.** By using the electrochemical, optical and computational data, the relevant thermodynamic driving forces (dG$^{\text{CS}}$) for intramolecular charge separation were calculated using the Rehm-Weller equation. By taking all the parameters into account, the estimated values of 1 and 2 are $-0.48$ eV and $-0.31$ eV, respectively. This result ensures that the electron transfer from the TTF units to the photoexcited porphyrin core is thermodynamically feasible reaction.

The supports for the formation of the proposed photoinduced charge separated species (i.e., [(TTF)$^+$*(P)$^-$*(TTF)]) of 1 and 2 were identified by EPR spectroscopy (Fig. S15). The EPR spectra of 1 and 2 recorded in CH$_2$Cl$_2$ exhibited a signal at g = 2.0058 and 2.0062 under Xe light irradiations, respectively, while the spectra taken under dark conditions were silent. The formation of TTF radical cation species in the chemical oxidations of 1 and 2 with tris(4-bromophenyl)aminium hexachloridoantimonate (known as Magic Blue) revealing EPR active signals at g tensor values of 2.0063 and 2.0065, respectively, are in consistent with those of the photoinduced electron transfer species in the oxidized species for 1 and 2 also support this conclusion (Fig. S16).

**Protonation-induced charge transfer.** Protonation of the pyrrolic nitrogen sites in the porphyrin ring could enhance the electron accepting ability due to the cationic electron-withdrawing effect in the core. Therefore we thought that the driving force energies for intramolecular electron transfer of 1 and 2 could be fine-tuned upon protonation of the core-modified porphyrins. The electronic and redox properties of the dicatonic species of 1 (i.e., H$_2$TTP$^{2+}$) and 2 (H$_2$2TTP$^{2+}$) were significantly altered with those of the neutral species. The UV-vis-NIR absorption spectrum of H$_2$TTP$^{2+}$ exhibited a further redshifted Soret band along with an unprecedented broad CT band reaching >1700 nm (Fig. 4a). Likewise, the significantly broad Soret band appeared at 600 nm and featureless Q-like bands were observed in the rubyrin system 2 (Fig. 4b). The different choice of acids (e.g., methanesulfonic acid) also yielded the similar spectral features in the absorption spectroscopy (Fig. S17). The fluorescence emission are redshifted and furthermore quenched, in contrast to the fact that the unsubstituted dications (H$_2$S$_2$TTP$^{2+}$ and H$_2$S$_2$TTR$^{2+}$) exhibited the enhanced fluorescent emissions (Fig. S18). This imply the enhancement of the internal CT nature occurring in the protonated forms.

Upon addition of TFA, the 1H NMR spectral signals of protonated species, H$_2$TTP$^{2+}$ and H$_2$2TTP$^{2+}$ were not changed initially, however, further addition of TFA afforded broadening resonance peaks (Fig. S19-20). It is noteworthy that diprotonation of 1 or 2 with excess amount of TFA afforded the thermoeffect electron transfer products partially judging from active EPR signals at g = 2.0066 at room temperature (Fig.4c and 4d). In this process, the formation of charge-separated species, i.e., [(TTF)$^+$H$^+$P$^-$*(TTF)]$^2$TFA$^-$ is thus assumed, since the cationic porphyrin core is a better electron transfer acceptor. Upon neutralization with base (e.g., triethylamine) yielded the recovery of the original spectra of 1 and 2. The dual roles of triethylamine acting as a proton scavenger and a reducing agent may be accounted for the reversibility.

In the electrochemical properties on the effect of protonation, both the oxidation and reduction potentials of H$_2$TTP$^{2+}$ and H$_2$2TTP$^{2+}$ were anodically shifted (Fig. 3c and 3d). Particularly, the extent of the shifts of reduction waves was remarkable. On this basis, protonation of the macrocyclic cores of 1 and 2 facilitates the intramolecular charge transfer as we expected. The resulting HOMO-LUMO gaps for H$_2$TTP$^{2+}$ and H$_2$2TTP$^{2+}$ are estimated to be 0.89 and 0.47 V, which are well consistent with the observations of the NIR lower energy band (> 2000 nm) appeared in the absorption spectra.
The DFT (B3LYP) calculations of the diprotonated species, H$_2$1$^{2+}$ and H$_2$2$^{2+}$ represent the deformation of the core structures and the extremely decreased energy gap between HOMO and LUMO (i.e., $AE = 1.13$ V for H$_2$1$^{2+}$ and 0.66 V for H$_2$2$^{2+}$) resulted from the remarkable destabilization of the HOMOs (Figs. S21-S22) in comparison with those of H$_2$S$_2$TTP$^{2+}$ and H$_2$S$_2$TTR$^{2+}$.\textsuperscript{22} Such MO behaviour in the diagrams is not seen for the unsubstituted derivatives. As the matter of the fact, the TD-DFT simulations of H$_2$1$^{2+}$ and H$_2$2$^{2+}$ also support the presence of optically allowed electronic transitions in the NIR absorptions (Figs S23). The attachment of two protons therefore triggers the drastic perturbation in their electronic structures via effective expansion of the protonated π framework through appropriate orbital admixing at the appended sulfur atoms of TTF peripheries.

Conclusions

In summary, we have synthesised and characterized the first core-modified families of TTF-annulated porphyrin 1 and rubyrin 2. In comparison with their photophysical properties of pyrolic counterparts (i.e., 3 or 4), the core modification in 1 was reflected in the difference of Gibbs driving force energies for electron transfer processes. The energetics of 1 were accounted for the facile reduction properties compared to the TTF-tetrapyrolic porphyrin systems. Similarly, the negative driving force is seen in the expanded 2. Furthermore, upon protonation of the macrocyclic cores, the energies for internal electron transfer could be enhanced due to the fact that protonation leads to the enhanced electron accepting ability of the macrocyclic core. Their HOMO-LUMO interactions were significantly changed as inferred by DFT calculations. Although the further studies on time-resolved laser spectroscopy is necessary to characterize the excited state dynamics of charge separation process, synthetic chemistry of these TTF-porphyrin ensembles would contribute the deep understanding of the photoinduced or thermal charge separation states. Furthermore, novel designs for organic materials tailored for electrochemically and optically controllable proton-electron coupled systems (e.g., artificial light harvesting antenna models) will be established.

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Notes and references


18. The typical g-tensor values for the radical cation species of TTF-annulated pyrrole has been reported to be at approximately 2.008. See; ref 2c.


20. The EPR spectra apparently favour to the formation of [(TTF)+H2P(TTF)]3+, which was presumed that the in situ oxidation of the product by molecular dioxygen.


22. In order to explain the magnetic properties, we have examined the open shell electronic structures of H1T1 and H2T2 by using unrestricted B3LYP methods. The contribution of the open shell electronic species, such as dication biradicals were found to be thermodynamically unfavorable; the singlet and triplet energy gap were ΔE = 2.31 kcal/mol and 3.61 kcal/mol, respectively.
The effect of the core modification and protonation on the intramolecular charge transfer phenomenon was studied in the core-modified families of TTF-annulated porphyrinoids.