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RESEARCH ARTICLE

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2026, 10, 938High quantum yield NIR emission *via* charge transfer states in buckybowl-TPA based D–A systemsYumi Yakiyama,^{id}*^{abc} Junyi Han,^a Hayato Sakai,^{id}*^d Taku Hasobe,^{id}^d
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The development of highly efficient near-infrared (NIR) emissive materials presents tremendous opportunities across diverse applications, from biological imaging to advanced optoelectronics. While planar aromatic dyes have traditionally dominated this field, we report herein the successful design and synthesis of a novel bowl-shaped NIR dye, **TPA-TOS**, incorporating the rarely emissive trioxosumanene (**TOS**) as the acceptor and triphenylamine (**TPA**) as the donor. In polar solvents, **TPA-TOS** exhibited weak luminescence confined to the UV region; however, in nonpolar environments, it demonstrated intense NIR emission spanning the 650 to 850 nm range. Remarkably, a quantum yield exceeding 60% in the NIR range was achieved, attributed to an intriguing photo-relaxation pathway. This mechanism involves exciton relaxation from localized excitation to the charge transfer (CT) state, mediated by thermally activated delayed fluorescence (TADF) between CT singlet and triplet states, along with room-temperature phosphorescence (RTP), ultimately transitioning to a long-lived charge-separated (CS) state. The outstanding amphiphilic nature, high quantum yield of NIR dual emission enabled by TADF and RTP, and the generation of a long-lived CS state are likely driven by the pronounced spin-orbit coupling facilitated by the curved sumanene skeleton. This work establishes **TPA-TOS** as a platform for NIR materials and insights into non-planar photophysics.

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Introduction

Near-infrared (NIR) dyes have garnered significant attention due to their vast array of applications in solar cells,¹ organic light-emitting diodes (OLEDs),² organic field-effect transistors (OFETs),³ and various biochemical fields.⁴ Beyond optoelectronics, NIR dyes have proven highly effective in tissue imaging and photopharmacology, owing to the minimal absorption and

autofluorescence of biological tissues in the NIR range (700–1000 nm).^{5,6} The design strategy for organic NIR dyes typically revolves around the donor- π -acceptor (D- π -A) paradigm, which facilitates intramolecular charge transfer (ICT) by modulating the frontier orbital energies—destabilizing the HOMO of the donor and stabilizing the LUMO of the acceptor—to precisely tune the energy gap between the ground and excited states.⁷ Numerous factors influence the photophysical behavior of such systems, including the nature and arrangement of electron-rich and electron-deficient chromophores, the functionalization and conjugation style of linkers bridging donor and acceptor units, their spatial distance, and environmental conditions. The efficiency and dynamics of charge transfer are largely governed by electronic coupling and solvent polarity: stronger electronic coupling promotes orbital overlap and enhances CT character, while polar solvents stabilize the CT state *via* dielectric screening, increasing the population of CT species.

Over the years, various π -conjugated charge transfer molecules and polymers have been developed for NIR applications such as LEDs and *in vivo* imaging.⁸ These materials often employ well-known strong electron acceptors, including benzothiadiazole (BT),⁹ benzobisthiadiazole (BBT),¹⁰ thiadiazoloquinoline (TQ),¹¹ triazolobenzothiadiazole (BTT),¹² diketopyrrolopyrrole

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(DPP),¹³ thienopyrroledione (TPD),¹⁴ and boron-dipyrromethene (BODIPY).¹⁵ However, such planar systems often suffer from strong π - π stacking, aggregation-induced quenching, and limited regulation of excited-state dynamics, which restrict their emission efficiency and multifunctionality. In contrast, non-planar conjugated molecules with twisted or distorted molecular geometries have recently emerged as an effective strategy to overcome these limitations.¹⁶ Zhao *et al.* reported a hybrid planar-twisted indoline donor that balances π -conjugation and molecular distortion, enabling bright near-infrared emission in both molecular and aggregated states.¹⁷ Sun *et al.* demonstrated that pyrazine-fused trichalcogenasumanenes with a non-planar π -conjugated framework enable effective excited-state regulation through triplet-state involvement, highlighting the role of molecular curvature in tuning excited-state dynamics.¹⁸ These studies indicate the feasibility of employing non-planar molecular structures for near-infrared emissive materials.

Recently, trioxosumanene (TOS), a tri-oxidized bowl-shaped derivative of sumanene, has emerged as a promising multi-step electron acceptor due to its exceptional electron-withdrawing capability, with reduction potentials surpassing those of fullerene derivatives (Fig. 1).^{19,20} Interestingly, TOS displays such pronounced electron-accepting properties that even strongly electron-deficient groups like nitrophenyl can function as donors when paired with it.²¹ Additionally, as a typical π - π^* triplet molecule, TOS exhibits rapid intersystem crossing (ISC) from the triplet state to the ground state (S_0), facilitated by a large spin-orbit coupling (SOC) due to its curved, non-planar structure.²² These unique features make TOS highly attractive for applications in electron injection and the development of organic phosphorescent materials.

In this study, we present the design and synthesis of TPA-TOS, a TOS-based donor-acceptor NIR dye with a distinctive molecular architecture (Fig. 1). Featuring the bulky triphenylamine (TPA) group as the donor and TOS as the acceptor, the system achieves efficient intramolecular charge transfer (ICT), resulting in a

narrowed bandgap and weak absorption extending into the NIR range up to 850 nm. Remarkably, TPA-TOS demonstrated an impressive quantum yield exceeding 60% in non-polar solvents such as hexane, with strong NIR emission spanning the 650–850 nm range. Femtosecond transient absorption spectroscopy (fsTA), electron spin resonance (ESR), and density functional theory (DFT) calculations suggest that the emission arises from thermally activated delayed fluorescence (TADF) and room-temperature phosphorescence (RTP), accompanied by partial charge separation. Furthermore, TPA-TOS exhibited distinct solvent dependency, showing negligible emission in polar solvents like acetonitrile (MeCN). With its unique mechanism and simplified donor-acceptor architecture, TPA-TOS paves the way for the development of efficient NIR dyes and highlights the tremendous potential of bowl-shaped molecular frameworks for advanced optoelectronic and biomedical applications.

Results and discussion

Synthesis and characterization of TPA-TOS

Fig. 2(a) illustrates the synthetic route to TPA-TOS. Bromotrioxosumanene (TOS-Br) was found to be a versatile precursor for obtaining various TOS derivatives¹⁸ via an acidic Suzuki-Miyaura coupling reaction,²⁰ which avoids the basic conditions, under which the sumanene skeleton is relatively unstable. Resulting TPA-TOS was obtained as a dark-green solid and its structure was characterized by high-resolution mass spectrometry (HRMS) and ¹H and ¹³C NMR spectroscopy (Synthetic details, NMR Spectrum S1–S3, SI). Although TPA-TOS had strong electron-accepting and electron-donating groups, resulting in a low HOMO–LUMO gap (1.809 eV at B3LYP/6-311+G(d,p)), it was stable under air and underwent only partial decomposition even at 400 °C. Interestingly, TPA-TOS was soluble not only in polar solvents such as MeCN and EtOH, but also in non-polar solvents such as hexane. This was probably attributed to the

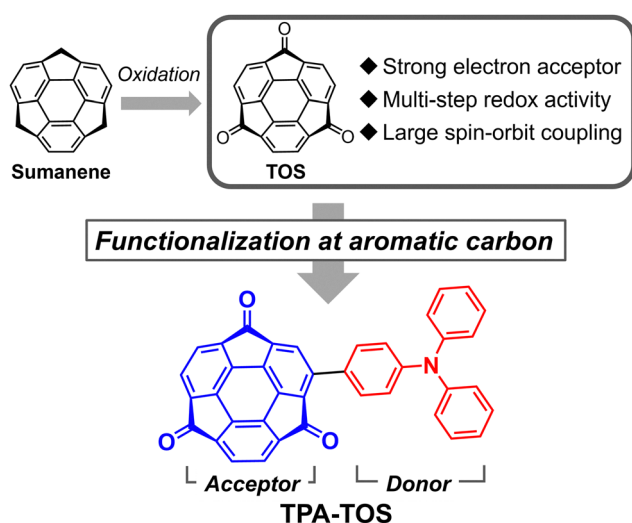


Fig. 1 Features of TOS and the schematic drawing of TPA-TOS.

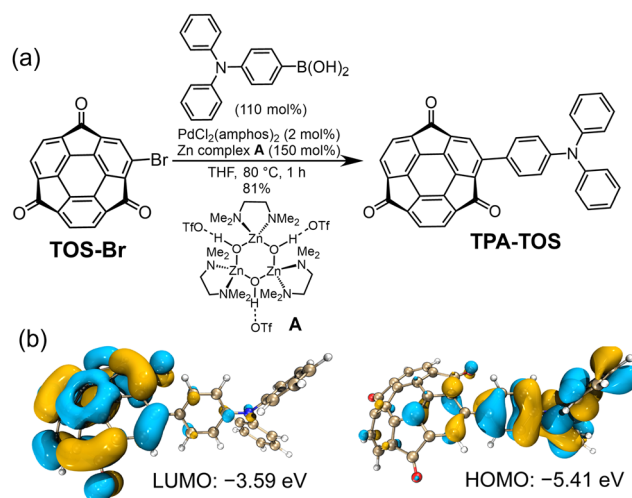


Fig. 2 Basic properties of TPA-TOS. (a) Synthetic procedure of TPA-TOS. (b) HOMO and LUMO distribution of TAP-TOS calculated at B3LYP/6-311+G(d,p) level.



curved structure of **TOS** moiety, in which partial dipole generates along the out of the plane direction of the bowl. Such carbon rich molecular skeleton with polarity realized the amphiphilic nature of **TPA-TOS**. Tremendous amounts of trials to get single crystals of **TPA-TOS** did not work to afford high quality crystal enough for single crystal X-ray diffraction analysis. Probably large **TPA** moiety disturbed the effective stacking formation of **TOS** moiety, giving only small low-quality crystals.

The electronic properties of **TPA-TOS** were next investigated. Density functional theory (DFT) calculations revealed that the HOMO was primarily localized on the **TPA** moiety, while the LUMO resided on the **TOS** skeleton, with slight orbital overlaps within each segment (Fig. 2(b)). Cyclic voltammetry (CV) and differential pulse voltammetry (DPV) measurements in MeCN containing 0.1 M of Bu_4NClO_4 as an electrolyte under a nitrogen atmosphere exhibited two distinct reversible reduction peaks at -0.79 V and -1.27 V at a scan rate of 10 mV s^{-1} (Fig. S1). This implied a strong electron-accepting ability of **TPA-TOS** reaching C_{60} as confirmed in **TOS** (-0.92 V and -1.43 V in THF).^{19,23} Meanwhile, the observed oxidation peak of **TPA-TOS** at $+0.55$ V indicated the slightly lower donating ability of the **TPA** part on **TPA-TOS** than the unmodified **TPA** ($+0.58$ V in MeCN)²⁴ due to the conjugation with electron-accepting **TOS**. Both HOMO and LUMO energy levels were calculated from the onset potentials (E_{onset}) to be -3.72 and -5.25 eV, respectively.²⁵ These values were in good agreement with the DFT-calculated values.

Spectroscopic analyses

To gain insight into the fundamental optical properties of **TPA-TOS**, both absorption and fluorescence spectra were measured (Fig. 3(a) and Table 1 and Table S1). The acceptor unit, **TOS**, exhibited short-wavelength absorption in the UV region, characterized by an absorption coefficient (ϵ) of $1.0 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ at 300 nm (Fig. S2). In contrast, **TPA-TOS** displayed distinct and intense absorption bands between 300–450 nm ($\epsilon = 7.6 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ at 300 nm in acetonitrile), along with a prominent absorption band spanning 600–800 nm ($\epsilon = 1.2 \times 10^4 \text{ M}^{-1} \text{ cm}^{-1}$ at 685 nm in MeCN). Computational results showed excellent agreement with the experimental data, indicating that the 600–800 nm absorption arises from charge transfer (CT) transitions between the donor (**TPA**) and the acceptor (**TOS**), while the 300–450 nm band corresponds to hybridized local excitation ($\pi-\pi^*$ transition) (Fig. S3 and S4). The photoluminescence (PL) spectrum of **TPA-TOS** in polar solvents closely resembled that of **TOS**, with a principal emission peak at 350–450 nm attributable to local excitation (LE) from the **TOS** moiety and a weak, broad peak around 590 nm (Fig. 3(b)). However, in nonpolar hexane, the emission intensity in the 350–400 nm region is markedly reduced, whereas a pronounced near-infrared (NIR) emission band centered at 703 nm emerges, accompanied by a shoulder at approximately 780 nm and an extended long-wavelength tail in the NIR region upon excitation at 300 nm (Fig. S5). Same NIR emission was observed under 650 nm excitation, confirming that this emission originates from the CT state, specifically $\text{TPA}^+-\text{TOS}^-$,

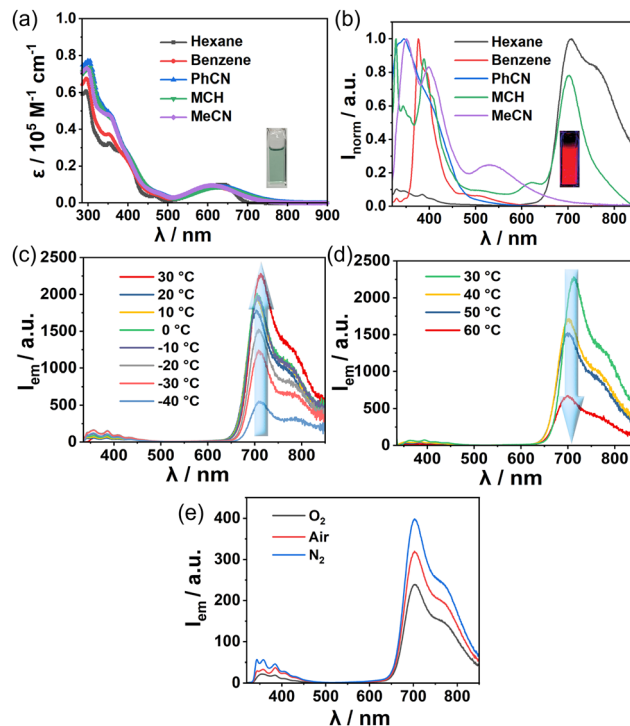


Fig. 3 Optical properties of **TPA-TOS**. (a) UV-vis-NIR absorption spectra of **TPA-TOS** in various solvents ($c = 1.0 \times 10^{-5} \text{ M}$). The inset photo was taken under general room light. (b) Photoluminescence spectra ($\lambda_{\text{ex}} = 300 \text{ nm}$) of **TPA-TOS** in diluted solutions ($c = 1.0 \times 10^{-5} \text{ M}$). The inset photo was taken under a UV lamp (365 nm) in hexane at dark. Intensity is normalized at the wave length with the highest peak intensity. (c and d) Temperature-dependent photoluminescence spectra ($\lambda_{\text{ex}} = 300 \text{ nm}$) of **TPA-TOS** ($c = 1.0 \times 10^{-5} \text{ M}$) in hexane (c) from -40 °C to 30 °C and (d) from 30 °C to 60 °C. (e) Effect of outer atmosphere on long wavelength emission of **TPA-TOS** at 300 K. PhCN: benzonitril, MCH: methylcyclohexane.

Table 1 Photophysical properties of **TPA-TOS**

Solvent	λ_{abs} [nm] ($\epsilon \times 10^4 [\text{M}^{-1} \text{ cm}^{-1}]$)	λ_{em} [nm] ^a (Φ_{em})
Hexane	296 (6.1)	350 (0.01 ^b)
	643 (1.2)	703 (0.66 ^b , 0.34 ^c)
MeCN	300 (7.6)	355 (0.01 ^b)
	662 (1.2)	

^a $\lambda_{\text{ex}} = 320 \text{ nm}$ at 300 K. ^b Under N_2 atmosphere. ^c Under O_2 atmosphere.

formed following photoexcitation (Fig. S6). The excitation spectrum demonstrated that **TPA-TOS** absorbs across nearly the entire visible range, enabling efficient NIR emission (Fig. S7). Relative to the weak emissive nature of **TOS** alone (Φ_{em} : 0.02 at 330 nm),²² **TPA-TOS** exhibits a significantly enhanced fluorescence quantum yield of 0.66 at 703 nm in hexane under an N_2 atmosphere. Additionally, its NIR emission capability is robust in PMMA and Zeonex films, achieving Φ_{em} values of 0.32 and 0.74, respectively (Fig. S8). The possibility of aggregation-induced effects was denied by the concentration dependent PL spectra ranging from 10^{-6} to 10^{-3} M . As shown in the experimental spectra (Fig. S9(b)), the emission intensity



decreases with increasing concentration, indicating that the molecule exhibits general aggregation-induced quenching behaviour. No Tyndall scattering was neither observed in non-polar solvent systems.

To elucidate the origin of this strong NIR emission, various external factors influencing emission properties were investigated. Temperature-dependent PL measurements revealed a rise in emission intensity as the temperature increased from $-40\text{ }^{\circ}\text{C}$ to $30\text{ }^{\circ}\text{C}$, followed by a noticeable decline at temperatures exceeding $30\text{ }^{\circ}\text{C}$ (Fig. 3(c) and (d)). This temperature-dependent behaviour is characteristic of systems where triplet states play a significant role in the emission process.^{26–28} At relatively low temperatures, a moderate increase in temperature facilitates thermally activated processes, including enhanced reversible intersystem crossing, resulting in increased fluorescence intensity. Upon further increasing the temperature into this range, enhanced molecular vibrational and rotational motions promote non-radiative decay pathways, leading to increased energy dissipation and a subsequent decrease in fluorescence intensity. Atmospheric effects further verified the contribution of triplet states to the NIR emission of **TPA-TOS** (Fig. 3(e)). Phosphorescence at 749 nm was observed both in 2-methyltetrahydrofuran glass at 77 K and in Zeonex film, which overlapped with the lower-energy portion of the emission spectrum (Fig. S9 and S10). Collectively, these findings suggest that the observed NIR emission arises from both singlet ($^1[\text{TPA}^+\text{TOS}^-]$, ^1CT) and triplet ($^3[\text{TPA}^+\text{TOS}^-]$, ^3CT) components. Importantly, the energy gap between ^1CT and ^3CT (ΔE_{st}) was experimentally estimated to be $<0.01\text{ eV}$ based on the fluorescence and phosphorescence emission spectra (Fig. S10a), implying efficient thermally activated delayed fluorescence (TADF) coupled with room-temperature phosphorescence (RTP), which is mainly reported in the solid state systems^{29–31} and the solution-based ones are still rare.^{32,33} This small ΔE_{ST} results from the effective spatial separation of the HOMO and LUMO (Fig. 2(b)) within the **TPA-TOS** system, underscoring the unique photophysical properties of TADF incorporating RTP in nonpolar environments such as hexane. As above discussion revealed the solvent dependency in the emission property of **TPA-TOS**, we measured the fluorescence spectra of **TPA-TOS** using a series of hexane–acetonitrile mixtures with varying polarity. It showed significant positive solvatochromism as a function of the solvent permittivity (Fig. S11). A linear correlation between Stokes shift and Lippert–Mataga solvent polarizability revealed that the dipole moment at the excited state (μ_e) reached 54 Debye (33 Debye by DFT calculation at B3LYP/6-311G**), which is significantly larger than the ground-state dipole moment μ_g (3.41 Debye). This finding supported the formation of CT structure at the excited state with a large dipole moment. Moreover, the fluorescence intensity of **TPA-TOS** decreased with increasing solvent polarity, indicating that polar solvents accelerate non-radiative deactivation of the CT species involved in delayed fluorescence.

Transient absorption and time-resolved spectroscopy

To investigate the solvent-dependent photo-relaxation dynamics of **TPA-TOS**, femtosecond transient absorption (fsTA)

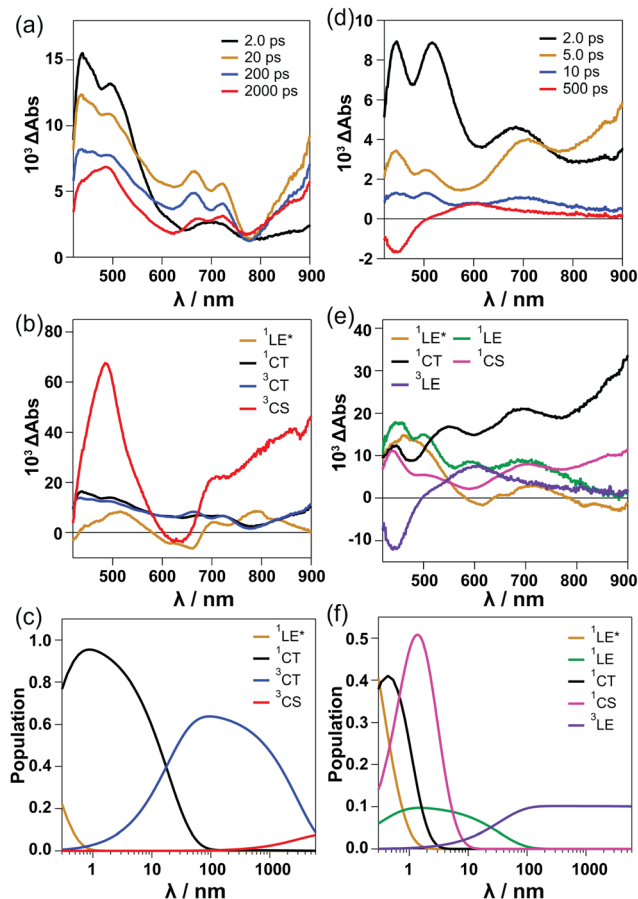


Fig. 4 Dynamic Optical properties of **TPA-TOS**. (a) and (d) fsTA spectra of **TPA-TOS**. (b) and (e) The species-associated spectra. (c) and (f) Time profiles of population change of each excited species. (a)–(c): obtained in hexane and (d)–(f): obtained in MeCN.

Table 2 Lifetime of each excited species

Solvent	$^1\text{LE}^*$	^1LE	^1CT	^3CT	^1CS	^3CS	^3LE
Hexane	$<410\text{ fs}$	—	19 ps	2.9 ns	—	$>6\text{ ns}$	—
MeCN	$<410\text{ fs}$	550 fs	1.9 ps	—	34 ps	—	$>6\text{ ns}$

measurements were performed in nonpolar hexane and polar acetonitrile (MeCN) (Fig. 4 and Table 2). Upon excitation at 343 nm, the fsTA maps during the initial 2 ps appeared similar in both solvents. Specifically, ultrafast spectral changes were observed, which are attributable to the localized excited (LE) state absorption originating from the **TPA** and **TOS** skeletons (Fig. 4(a) and (d)). This assignment aligns well with previously reported transient absorption (TA) spectra of **TOS**²² and photo-excited **TPA**³⁴ (Fig. S12). Meanwhile, a broad absorption component around 700 nm, corresponding to charge transfer (CT) absorption associated with the formation of TPA^+TOS^- , emerged in both solvents, indicating that fast CT processes occur even in nonpolar hexane. At longer timescales—after 5 ps in MeCN and 20 ps in hexane—a new absorption band emerged in the near-infrared (NIR) region beyond 900 nm, which



extended past 1000 nm as confirmed by picosecond transient absorption (psTA) measurements (Fig. S13). This long-wavelength absorption exceeding 1000 nm closely resembles the characteristic absorption of the TOS radical anion ($\text{TOS}^{\bullet-}$, 1040 nm).¹⁹ Simultaneously, the characteristic dual absorption band of the TPA radical cation ($\text{TPA}^{\bullet+}$) appeared at 650–750 nm, corroborating previous reports on $\text{TPA}^{\bullet+}$ absorption spectra.³⁵ Taken together, these findings strongly support the generation of a charge-separated (CS) state, specifically $\text{TPA}^{\bullet+}\text{-TOS}^{\bullet-}$, in both hexane and MeCN. These solvent-independent CS state dynamics and their distinct spectral fingerprints provide direct insight into the photophysical behaviours underlying TPA-TOS, establishing its ability to undergo efficient charge separation and radical formation across a wide range of solvent polarities.

To further confirm the formation of the charge-separated (CS) state, electron spin resonance (ESR) measurements were conducted, which provided evidence for the generation of triplet radical species [$^3(\text{TPA}^{\bullet+}\text{-TOS}^{\bullet-})$] (^3CS) (Fig. 5(a)). Upon photoirradiation of a hexane solution of TPA-TOS at 123 K with 365 nm UV light, an ESR signal was detected at $g = 2.0049$, along with a distinct zero-field splitting pattern—a hallmark of triplet state formation. This observation unequivocally confirmed the presence of the ^3CS species. The zero-field splitting parameter (D) was determined from the fine structure of the ESR spectrum to be 116 G (11.6 mT), which was much smaller than the reported D value of the ^3CT species of carbazole-based small D-A system, reflecting the longer spin-spin distance than the case of ^3CT state.³⁶ From this parameter, the distance (r) between the two unpaired electrons was calculated to be 6.2 Å,³⁷ consistent with previously reported spin-spin distances in similar triplet radical ion pairs. This calculated distance aligns well with the structural characteristics of the $\text{TPA}^{\bullet+}\text{-TOS}^{\bullet-}$ ^3CS species, further supporting its assignment as a triplet charge-separated state (Fig. 5(b)). These ESR findings provide direct evidence for the involvement of the triplet CS state in the photophysical processes of TPA-TOS.

A more detailed analysis revealed a striking difference in the fate of the photo-relaxation process depending on the solvent polarity. In nonpolar hexane, the charge-separated (CS) states

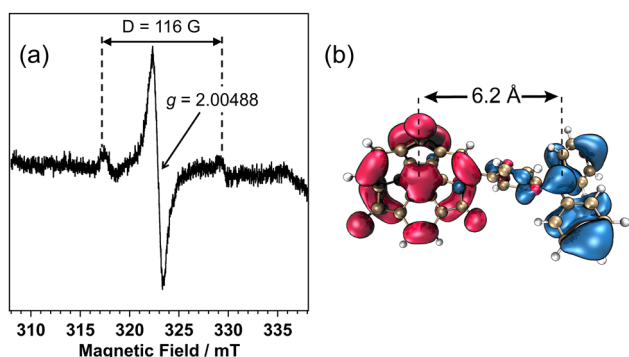


Fig. 5 Radical ion pair of TPA-TOS. (a) ESR spectrum of TPA-TOS (5×10^{-4} M) in deaerated hexane after 365 nm photoirradiation at 123 K. (b) Scheme of radical ion pair distance. Spin density plot of was calculated as [$^3(\text{TPA}^{\bullet+}\text{-TOS}^{\bullet-})$] at UB3LYP/6-311+(d,p).

exhibited lifetimes ranging from several thousand picoseconds to hundreds of microseconds (Fig. 4(a) and Fig. S13(c) and (d)), indicating that the relaxation pathway involved intersystem crossing (ISC) and reverse intersystem crossing (RISC) processes between the ^1CT and ^3CT states. In contrast, in polar MeCN, all excited states of TPA-TOS rapidly relaxed within 10 ps, ultimately producing a weak broad absorption band around 600 nm after 500 ps, likely attributable to the previously reported triplet state of TOS (Fig. S12(b)).²² This rapid decay process in MeCN, lacking ^1CT emission, suggests a competition between two relaxation pathways: one leading to the ^3LE state via the ^1LE state, and the other proceeding to the ^1CS state through fast electron transfer from the TPA to the TOS moiety of the ^1CT species. Based on these assumptions, the best-fitting results revealed the involvement of ^1LE , ^3LE , ^1CT , and ^1CS states in the MeCN solution, while additional states— ^3CT and ^3CS —were found to contribute to the hexane solution (Fig. 4(b), (c), (e) and (f)). The proposed emission mechanisms, depending on the solvent system, are summarized in Fig. 6. The results further clarified key rate dynamics. In hexane, the ISC rate from the ^1CT to ^3CT state ($k_{3\text{CT}}$: $3.53 \times 10^{10} \text{ s}^{-1}$) was higher than the fluorescence rate from the ^1CT state (k_{LUM^1} : $1.81 \times 10^{10} \text{ s}^{-1}$),

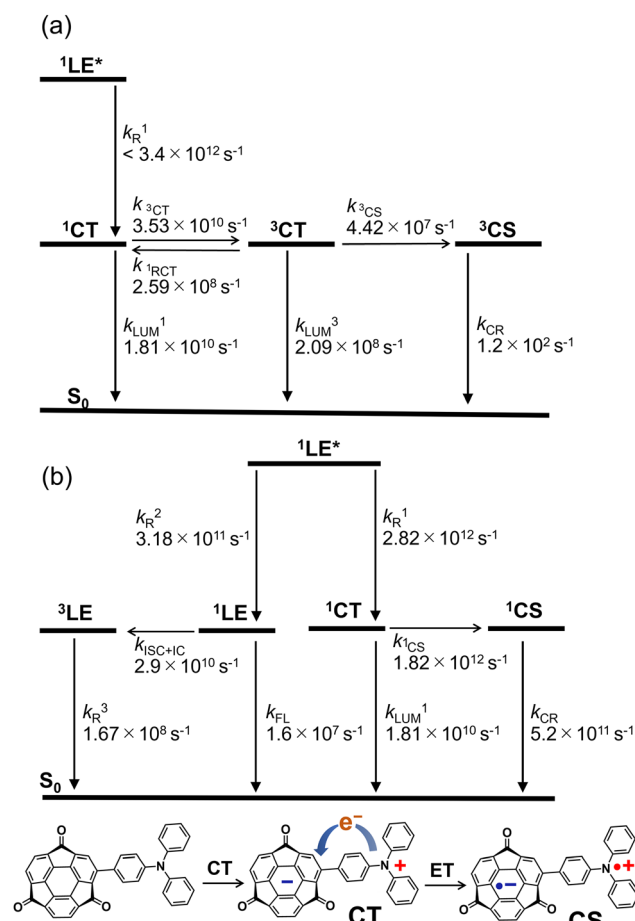


Fig. 6 Energy-level diagram of TPA-TOS. (a) Schematic illustration of the excited-state energy relationships of TPA-TOS in hexane. (b) Schematic illustration of the excited-state energy relationships of TPA-TOS in MeCN.



explaining the negligible emission intensity from the ^1LE state. Another noteworthy observation is that the RISC rate from the ^3CT to ^1CT state ($k_{1\text{RCT}}: 2.59 \times 10^8 \text{ s}^{-1}$) was nearly identical to the phosphorescence rate ($k_{\text{LUM}}^3: 2.09 \times 10^8 \text{ s}^{-1}$), facilitating efficient thermally activated delayed fluorescence (TADF) alongside phosphorescence emission to give high Φ_{em} reaching more than 0.60. Such coexistence of TADF and room-temperature phosphorescence (RTP) is relatively rare, particularly in solution systems.³⁸ Considering the case in MCH and benzene, probably the population of TADF and RTP path is likely controlled by the solvent systems with relatively low polarity, which will be our future work (Fig. 3(b)). The relaxation pathway from the ^3CT state to the ^3CS state was slower by an order of magnitude compared to the RISC process, resulting in only a small fraction of the excitons reaching the CS state as their final destination. These CS-state excitons transitioned slowly to the ground state *via* charge recombination ($k_{\text{CR}}: 1.2 \times 10^2 \text{ s}^{-1}$). Conversely in MeCN, two competitive fast relaxation pathways were observed: one leading to the ^1CT state ($k_{\text{R}}^1: 2.82 \times 10^{12} \text{ s}^{-1}$) and the other to the ^1LE state ($k_{\text{R}}^2: 3.18 \times 10^{11} \text{ s}^{-1}$). The faster charge transfer process quickly transitioned to the ^1CS state ($k_{1\text{CS}}: 1.82 \times 10^{12} \text{ s}^{-1}$), followed by charge recombination ($k_{\text{CR}}: 5.2 \times 10^{11} \text{ s}^{-1}$) due to a significant stabilization of CT and CS states in polar solvent. On the other hand, the slower relaxation path to the ^1LE state eventually reached the ^3LE state *via* ISC ($k_{\text{ISC+IC}}: 2.9 \times 10^{10} \text{ s}^{-1}$), exhibiting negligible fluorescence emission ($k_{\text{FL}}: 1.6 \times 10^7 \text{ s}^{-1}$).

Plausible discussion about the effect of the curved- π structure

Considering that previously reported materials exhibiting long-CS lifetimes often feature a C_{60} skeleton, and that the curved- π systems typically display large spin-orbital coupling which accelerates ISC, it is reasonable to have interest in the contribution of the sumanene skeleton for the relevant phenomena. To tackle this mystery, we conducted density functional theory (DFT) calculation. While several benchmarking studies have suggested recommended computational methods for the excited-state properties of intramolecular CT systems, it was found that the methodological dependence of **TPA-TOS** is rather complex due to its extended π -conjugation system. Indeed, as already applied in the evaluation of UV-vis absorption measurement results, TD-RB3LYP/6-311G(d,p) level of theory well reproduced the experimental CT band around 600–800 nm, although the excitation energy appears to be underestimated to some extent. This method also predicted the dihedral angle θ -dependent S_1 - T_1 gap (ΔE_{ST}) and $\Delta E_{\text{ST}} \sim 0$ around $\theta = 90^\circ$. The S_1 potential energy curve was nearly flat in the range of $30^\circ \leq \theta \leq 120^\circ$, and the geometry optimization for the S_1 state converged to a structure with $\theta \sim 90^\circ$ (Fig. 7). This result supports experimental observation. However, further theoretical investigation revealed significant dependence on the function used, which prevents us from drawing more detailed conclusions how bowl shape of sumanene contributed the effective ISC and the high stability of CS state with μs -ordered lifetime at this stage (see SI). Nevertheless, the effect of curved- π structure of sumanene cannot be overlooked, as it at

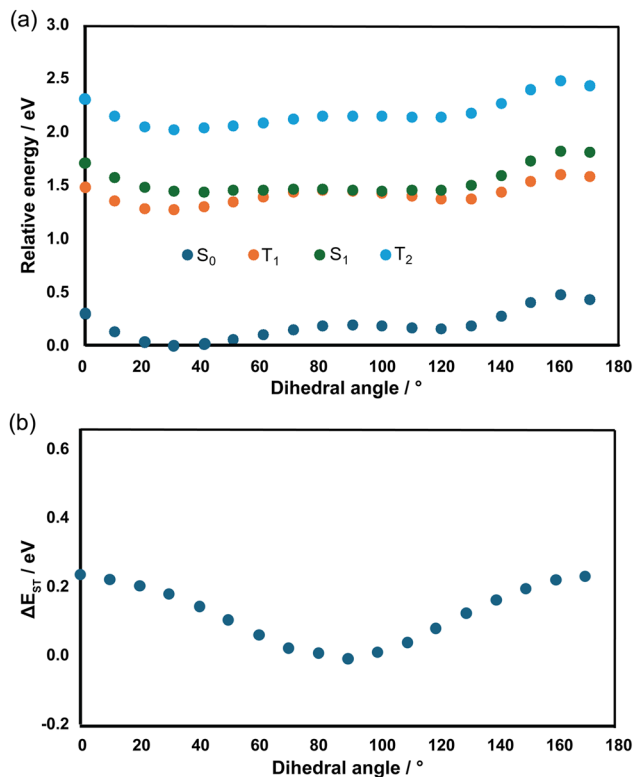


Fig. 7 Preliminary energy evaluation depending on the dihedral angle between **TPA** and **TOS** moieties. (a) The dihedral angle-dependent relative energy of S_0 , S_1 , T_1 , T_2 states. (b) The S_1 - T_1 gap (ΔE_{ST}) of **TPA-TOS** was calculated at TD-RB3LYP/6-311G(d,p) level of theory.

least contributed to; (1) high solubility of **TPA-TOS** even in non-polar hexane; (2) possessed a large |SOC| to accelerate ISC between ^1CT and ^3CT , supporting effective TADF and RTP to give high quantum yield, as well as long life time of the CS state. Although additional theoretical refinement is required to fully understand the impact of the sumanene skeleton and its bowl-shaped geometry, the experimental and theoretical findings suggest that this unique curved- π architecture plays a pivotal role in enhancing the photophysical performance of **TPA-TOS**. Indeed, such novel effect of the curved structure has also been confirmed in pristine **TOS**²² and recently reported *N*-pyrrolic polycyclic aromatic hydrocarbons.³⁹ In the present case, the curved- π skeleton supports not only efficient ISC and TADF/RTP processes with high quantum yield, but also the observed long-lived CS state, which could have significant implications for applications in advanced photonic and optoelectronic devices.

Conclusions

In this study, we have developed a novel donor-acceptor (D-A) near-infrared (NIR) dye, **TPA-TOS**, featuring a rigid bowl-shaped acceptor, **TOS**, and a strong donor, **TPA**, without relying on long-range conjugation or complex multi-donor/acceptor systems. This unique molecular design successfully establishes an effective TADF and RTP system, enabling high-efficiency NIR



emission with an impressive quantum yield. Remarkably, the exciton in nonpolar hexane proceeds through efficient charge separation, reaching a long-lived CS state despite the short D–A distance. Conversely, in polar MeCN, the CS state becomes short-lived due to faster relaxation dynamics. Although further theoretical studies are required for a deeper understanding, the curved TOS framework clearly contributes to the observed photophysics, providing a high solubility in non-polar hexane, large |SOC| to accelerate ISC between ¹CT and ³CT states, giving rise to TADF and RTP. This unique design and the remarkable photophysical characteristics of TPA-TOS open new avenues for the development of efficient NIR dyes with innovative molecular architectures. By overcoming conventional limitations of D–A systems, this work provides a promising platform for future advancements in optoelectronic applications and deepens understanding of charge separation dynamics in tailored curved- π systems.

Author contributions

Han Junyi, Yumi Yakiyama and Hidehiro Sakurai: conceptualization. Han Junyi, Yumi Yakiyama and Hayato Sakai: validation. Han Junyi, Yumi Yakiyama, Hayato Sakai, Youhei Takeda, Ryohei Kishi and Kei Ohkubo: investigation. Han Junyi and Yumi Yakiyama: writing – original draft. Han Junyi, Yumi Yakiyama, Hayato Sakai, Taku Hasobe, Youhei Takeda, Ryohei Kishi, Kei Ohkubo and Hidehiro Sakurai: writing – review & editing. Yumi Yakiyama, Ryohei Kishi, Kei Ohkubo and Hidehiro Sakurai: funding acquisition. Yumi Yakiyama, Hayato Sakai and Hidehiro Sakurai: supervision.

Conflicts of interest

There are no conflicts to declare.

Data availability

The authors confirm that the data supporting the findings of this study are available within the article and its SI. Experimental raw data were generated at The university of Osaka and Keio University. Derived data supporting the findings of this study are available from the corresponding author Y. Y. on request.

Supplementary information (SI) including synthetic details, NMR, MS, PL, TA spectra and DFT calculation detail is available. See DOI: <https://doi.org/10.1039/d5qm00818b>.

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