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Introduction

Photosensitization occurs when light at an appropriate wavelength interacts with a photosensitizer, from which the photon energy is transferred onto target molecules.¹ Photodynamic therapy (PDT) is an alternative, minimally invasive treatment for various human diseases,^{2,3} including cancer, rheumatoid arthritis, and psoriasis, that uses only photon energy.¹ PDT uses a photosensitizer that releases appropriate energy to generate singlet oxygen (¹O₂) from triplet oxygen (³O₂).⁴ The cellular oxygen (³O₂) absorbs the energy released from the photosensitizer to generate ¹O₂,⁵ and the toxicity of ¹O₂ leads to cell death in cancer, *e.g.*, for ³O₂ (³ \sum_{g}) \rightarrow ¹O₂ (¹ Δ_{g}), $\Delta E^{T_1 \rightarrow S_1} \sim 0.97$ eV (ref. 5) (~94 kJ mol⁻¹); in normal cells (healthy tissues), the oxygen levels are in general lower compared to cancer cells, thus resulting in less ¹O₂ generation and lower cytotoxic effects.⁶

Photoluminescence mechanisms of BF₂formazanate dye sensitizers: a theoretical study†

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Photodynamic therapy (PDT) is an alternative, minimally invasive treatment for human diseases such as cancer. PDT uses a photosensitizer to transfer photon energy directly to cellular ${}^{3}O_{2}$ to generate ${}^{1}O_{2}$ (Type II), the toxicity of which leads to cancer cell death. In this work, the photoluminescence mechanisms of a BF₂-formazanate dye sensitizer (BF₂-FORM) and its iodinated derivative (BF₂-FORM-D) were studied using complementary theoretical approaches; the photoluminescence pathways in the S_1 and T₁ states were studied using density functional theory (DFT) and time-dependent (TD)-DFT methods, the kinetic and thermodynamic properties of the pathways using the transition state theory (TST), and the time evolution and dynamics of key processes using non-adiabatic microcanonical molecular dynamics simulations with surface-hopping dynamics (NVE-MDSH). Evaluation of the potential energy surfaces (PESs) in terms of the rotations of the phenyl rings suggested a pathway for the S₁ \rightarrow S₀ transition for the perpendicular structure, whereas two pathways were anticipated for the $T_1 \rightarrow S_0$ transition, namely, $[T_1 \rightarrow S_0]_1$ occurring immediately after the S_1/T_1 intersystem crossing (ISC) and $[T_1 \rightarrow S_0]_1$ $S_0]_2$ occurring after the S_1/T_1 ISC and T_1 equilibrium structure relaxation, with the $T_1 \rightarrow S_0$ energy gap being comparable to the energy required for ${}^3\text{O}_2 \rightarrow {}^1\text{O}_2$. The PESs also showed that because of the heavy-atom effect, BF_2 -FORM-D possessed a significantly smaller S_1/T_1 energy gap than BF_2 -FORM. The TST results revealed that at room temperature, BF₂-FORM-D was thermodynamically more favorable than the parent molecule. Analysis of the NVE-MDSH results suggested that the librational motions of the phenyl rings play an important role in the internal conversion (IC) and ISC, and the S_1/T_1 ISC and T_1 \rightarrow S₀ transitions could be enhanced by varying the irradiation wavelength and controlling the temperature. These findings can be used as guidelines to improve and/or design photosensitizers for PDT.

> Effective photosensitizers for therapeutics should have high light absorption coefficients, particularly in the infrared/nearinfrared range, to allow for effective tissue penetration. They should exhibit low photobleaching quantum yields, high intersystem crossing (ISC) efficiencies, and low toxicity in the absence of light.⁷ These include the photodynamic properties, such as appropriate triplet relaxation energy ($\Delta E^{T_1 \rightarrow S_0}$), high phosphorescence quantum yield (Φ_T), and long lifetime (τ^{T_1}) of the triplet state.⁸

> The two photochemical mechanisms of PDT involving oxygen molecules are shown in Fig. 1. In Type I, radicals (*e.g.*, OH[•]) act as intermediates, transferring electron energy from the photosensitizer to the oxygen derivatives, whereas for Type II, the photosensitizer passes light energy directly to the oxygen molecules.⁴ In general, phosphorescence is not easily observed because of the interference of fluorescence,⁸ and triplet excited states are difficult to generate through direct photoexcitation because ISC is symmetrically forbidden. For example, the $S_1 \rightarrow T_1$ transition in chromophores possesses a large singlet–triplet energy gap.⁹

> Theoretical and experimental studies have proposed several strategies to improve the efficiency of ISC and promote the

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Fig. 1 Schematic diagram showing two types of reaction pathways for formation of singlet oxygen from triplet oxygen, ${}^{3}O_{2} ({}^{3}\Sigma_{g}) \rightarrow {}^{1}O_{2} ({}^{1}\Delta_{g})$. ISC = intersystem crossing; IC = internal conversion; F = fluorescence; P = phosphorescence.

generation of triplet states. These strategies include for example: Heavy atom effect;¹⁰ the introduction of heavy atoms like halogen (such as Br and I) into photosensitizers has been shown to increase ISC rates, and the introduction of chalcogen (such as S, Se, or Te) into the photosensitizers could also increase the production of ${}^{1}O_{2}$ upon irradiation due to spinorbit interactions:¹¹ Molecular design;¹² the modification of the structure of photosensitizer molecules can optimize their electronic characteristics to promote ISC and: Sensitizer-sensitizer interactions;¹³ ISC can also be enhanced through interactions among photosensitizer molecules, whether through complex formation or aggregation.

Boron difluoride (BF₂) complexed with π -conjugated organic compounds is widely used as a photosensitizer in PDT, among which boron dipyrromethene (IUPAC = 4,4-difluoro-4-bora-*3a*,4*a*-diaza-*s*-indacene, or BODIPY) and its derivatives are the most widely studied.^{14,15} Such compounds have been reported to possess visible absorption and fluorescence emission between 470 and 550 nm, with a high emission quantum yield ($\Phi_{\rm F}$ = 0.60).¹⁶ Although BODIPY has a high potential to be a functional component in PDT, attempts have been made to improve $\Phi_{\rm T}$ using phenyl and/or heavy-atom substitutions.⁸

Several experimental methods have been developed to enhance the triplet excitation of photosensitizers by incorporating halogen atoms, such as the iodine (I) or bromine (Br) atoms, into organic aromatic compounds; these heavy atoms could help increase the spin–orbit coupling (SOC) and ISC rate, as well as the ${}^{1}O_{2}$ quantum yield of photosensitizers.^{17,18} In most cases, the I atom has a higher ${}^{1}O_{2}$ quantum yield (SOQY) than the Br atom.¹⁹ The higher Φ_{T} and SOQY for the I atom than the Br atom can be attributed to several factors, such as the higher heavy atom effect, higher radiative decay and lower internal conversion (IC) rates.^{20–22} These collective factors could lead to an enhancement of Φ_{T} and SOQY in BODIPY-based PDT photosensitizers.

BF₂-formazanate (3,3-difluoro-2,4-diphenyl-2,3-dihydro-1,2,4 λ^4 ,5,3 λ^4 -tetrazaborinine, BF₂-FORM in this work) is a fluorophore with several applications in microscopy. The structure of BF₂-FORM, shown in Fig. 2, consists of a BF₂ group coupled to a chelating N-donor ligand, forming a stable six-membered heterocyclic ring. BF₂-FORM exhibits outstanding photophysical properties such as large molar extinction coefficients and high Φ_F , which are generally in the far-red or near-infrared region. Therefore, BF₂-FORM has significant potential in cell imaging and PDT applications.^{23,24}

The optical properties of BF₂-FORM are strongly affected by the electron-donating substituents at R_2 and R_3 (Fig. 2), leading to an increase in Φ_F and a red-shift in the emission spectra. The



Fig. 2 (a) and (b) Structures and numbering systems of BF₂-FORM and BF₂-FORM-D dye sensitizers, respectively. BF₂-FORM = 3,3-difluoro-2,4-diphenyl-2,3-dihydro-1,2,4 λ^4 ,5,3 λ^4 -tetrazaborinine; BF₂-FORM-D = 3,3-difluoro-2,4-bis(4-iodophenyl)-2,3-dihydro-1,2,4 λ^4 ,5,3 λ^4 -tetrazaborinine; Carbonitrile; ω_1 and ω_2 = dihedral angles used in the calculations of the potential energy surfaces.

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strong red-shift could be attributed to a smaller HOMO–LUMO energy gap upon the substitutions.⁶ For example, in the case of F-BODIPY, the introduction of two ethyl groups at the C₂ and C₈ positions led to a red-shifted absorption spectra compared to the parent molecule due to a decrease in the HOMO–LUMO energy gap and large charge transfer interaction within the molecule.²⁵ Because electron-withdrawing substituents at R₁ could also affect the photophysical properties, the differences in optical properties between Ph-, CN-, and NO₂-substituted BF₂-FORM are attributed primarily to the electron-withdrawing nature of the substituents (*e.g.*, NO₂ > CN \gg Ph).²⁶

In our previous study,²⁴ the photochemical properties of BF₂-FORM-based photosensitizers were experimentally and theoretically studied in the electronic ground (S₀), lowest singlet, and triplet excited states (S₁ and T₁) using density functional theory (DFT) and time-dependent density functional theory (TD-DFT) methods with the Becke, 3-Parameter, Lee–Yang–Parr (B3LYP) hybrid functional and 6-311G basis sets. A comparison of the experimental and theoretical results showed that the DFT/B3LYP/ 6-311G and TD-DFT/B3LYP/6-311G methods could provide insight into the PDT mechanisms and confirmed the effect of heavy-atom substituents (I and Br at R_2 and R_3) on the ISC rate.

In this study, complementary theoretical approaches were applied to investigate the photoluminescence mechanisms of BF₂-FORM to improve its efficiency as a photosensitizer in PDT. Because theoretical and experimental studies²⁴ revealed that substitutions of R₂ and R₃ at the phenyl rings by I and R₁ at the heterocyclic ring by CN can significantly enhance the S₁/T₁ ISC with high $\Phi_{\rm T}$, both BF₂-FORM and its iodinated derivative [3,3-difluoro-2,4-bis(4-iodophenyl)-2,3-dihydro-1,2,4 λ^4 ,5,3 λ^4 -tetrazaborinine-6-carbonitrile, BF₂-FORM-D in this work] were selected as model molecules.

Theoretical studies focusing on Type II mechanism began with calculations of the equilibrium structures and energetic and spectroscopic properties of BF2-FORM and BF2-FORM-D in the S₀, S₁, and T₁ states. The potential energy surfaces (PESs) for the $S_1 \rightarrow S_0$, S_1/T_1 , and $T_1 \rightarrow S_0$ transitions were computed using the nudged elastic band (NEB) method, from which the kinetics and thermodynamics of the photoluminescence pathways were studied using the transition state theory (TST). Emphasis was placed on the probabilities of IC and ISC, as well as on the effect of the electron-withdrawing substituents on the photophysical properties. Furthermore, to explore the possibility of increasing photoluminescence, non-radiative relaxation, and effect of molecular dynamics on the $S_1 \rightarrow S_0$ transition and S₁/T₁ ISC were studied using non-adiabatic microcanonical molecular dynamics simulations with surface-hopping dynamics (NVE-MDSH). The theoretical results are discussed in comparison with available theoretical and experimental data.

Computational methods

Quantum chemical calculations

All the DFT and TD-DFT calculations were performed using the TURBOMOLE 7.50 software package.²⁷ For the TD-DFT method, the Tamm–Dancoff approximation (TDA) was applied to avoid singlet instabilities in the lowest singlet and triplet state

calculations. The DFT and TD-DFT methods with the B3LYP functionals were chosen based on several benchmarking calculations in photochemical reactions,28,29 including BODIPY-based photosensitizers.9 For example, our benchmarks against the complete active space multiconfigurational second-order perturbation theory (CASPT2) method revealed that for the photodissociation and formation of glycine,28,29 the characteristic structures and energies on the S₀ and S₁ PESs obtained from the DFT/B3LYP and TD-DFT/B3LYP methods were in good agreement with the CASPT2 results, and DFT/B3LYP/6-311G calculations on the I2-IR783-Mpip photosensitizer⁶ revealed the equilibrium structures, energetics and absorption spectra comparable with experimental data; the 6-311G basis set was also used successfully in a theoretical study on photoinduced charge separation-charge recombination in BODIPY compounds.30 The performance of the DFT/B3LYP method with various sizes of the basis sets was discussed in detail using boron-doped triazine based covalent organic framework as a model molecule in ref. 31.

Equilibrium structures. To study photoluminescence pathways for BF₂-FORM and BF₂-FORM-D, the theoretical strategy and methods shown in Fig. 3 were used. Calculations of the equilibrium structures, energetics, and spectroscopic properties of the S₀, S₁, and T₁ states were performed using the DFT/ B3LYP/6-311G and TD-DFT/B3LYP/6-311G methods. The calculated equilibrium structures and energies of BF₂-FORM and BF₂-FORM-D in the S₀, S₁, and T₁ states are listed in Tables S1 and S2,[†] respectively. The absorption spectra and fluorescence lifetimes of BF₂-FORM and BF₂-FORM-D were computed using the NEWTON-X software package³²⁻³⁴ interfaced with TURBO-MOLE 7.50, for which 200 Wigner-sampled structures were used as initial conditions. The results were used as guidelines in the photoluminescence pathway analysis.

Reaction pathway optimizations. The photoluminescence pathways were hypothesized in this work to start with an $S_0 \rightarrow S_1$ vertically excited structure (I)*, as shown in Fig. 4, followed by (I)* \rightarrow (II)* structural relaxation and $S_1 \rightarrow S_0$ transition ([$S_1 \rightarrow S_0$]). Two pathways were hypothesized for $T_1 \rightarrow S_0$ transition, namely, (I)* \rightarrow (II)* \rightleftharpoons (III)^{‡,§} and S_1/T_1 ISC followed by $T_1 \rightarrow S_0$ transition ([$(T_1 \rightarrow S_0)$]₁) or (I)* \rightarrow (II)* \rightleftharpoons (III)^{‡,§} and S_1/T_1 ISC followed by (III)[§] \rightarrow (IV) structural relaxation in the T_1 state and $T_1 \rightarrow S_0$ transition ([$(T_1 \rightarrow S_0)$]₂). These results led to the following three deactivation pathways:

$$(I)^* \to (II)^* \to [S_1 \to S_0] \tag{1}$$

$$(I)^* \to (II)^* \rightleftharpoons (III)^{\ddagger, \$} \to S_1/T_1 \text{ ISC} \to [T_1 \to S_0]_1 \qquad (2)$$

$$(I)^* \to (II)^* \rightleftharpoons (III)^{\ddagger, \$} \to S_1/T_1 \text{ ISC } \to (IV) \to [T_1 \to S_0]_2 (3)$$

The equilibrium structures of BF_2 -FORM and BF_2 -FORM-D in the S_0 , S_1 , and T_1 states obtained from DFT/B3LYP/6-311G and TD-DFT/B3LYP/6-311G geometry optimizations were used in the PES calculations.

Theoretical studies have shown that ISC in aromatic organic compounds can be mediated by intramolecular motions³⁵ such as molecular rotation or twist;³⁶ thus, reaction pathway

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Fig. 3 The theoretical strategy and methods used to study photoluminescence mechanisms of the BF₂-formazanate dye sensitizers. PES = potential energy surface; ΔE^{Ex} = excitation energy; IC and ISC = internal conversion and intersystem crossing; TST = transition state theory; NVE-MDSH = non-adiabatic microcanonical molecular dynamics simulations with surface hopping dynamics.

optimization began with the $S_0 \rightarrow S_1$ vertically excited precursor (I)*, from which the PESs for the rotations of the torsional angles ω_1 and ω_2 (Fig. 2) were constructed. Reaction pathway optimization was performed using the NEB method³⁷ with limited-memory Broyden-Fletcher-Goldfarb-Shanno (LBFGS)

optimizers included in the ChemShell software package.³⁸ In this work, to search for minimum energy pathways connecting the initial and final structures on the PESs, approximately 10 intermediate images (including the saddle points) along the pathways were optimized. In the NEB calculations,³⁷ the



Reaction path

Fig. 4 The hypothesized photochemical deactivation pathways ((I)* \rightarrow (II)* \rightleftharpoons (III)^{\ddagger , §} \rightarrow (IV)) used in the present study.

gradients on the reaction pathway were calculated based on the spring forces acting on local tangents between each image and on the true forces acting perpendicular to the local tangents.

Because SOC plays an important role in ISC and phosphorescence, the singlet-triplet energy gaps along the S_1 PES were computed using the TD-DFT/B3LYP/6-311G method and the geometries obtained from the NEB calculations. The energy gaps were refined using the RICC2/aug-cc-pVDZ method, from which the phosphorescence lifetimes were computed. SOC was computed using TURBOMOLE 7.50 based on the effective spinorbital mean field approximation,³⁹ in which the mean field two-electron contribution was computed from the Hartree–Fock density.

Kinetics and thermodynamics of reaction pathways

To study the kinetics and thermodynamics of the proposed photoluminescence pathways, quantized vibrational rate constants ($k^{\text{Q-vib}}$) were calculated over a temperature range of 300–550 K. This range of temperatures includes the standard human body temperature of 310 K. The $k^{\text{Q-vib}}$ values were computed using eqn (4),⁴⁰ in which $\Delta E^{\ddagger,\text{ZPC}}$ is the zero-point energy-corrected barrier, obtained by including the zero-point correction energy (ΔE^{ZPE}) to the energy barriers obtained from the NEB method (ΔE^{\ddagger}).

$$k^{\text{Q-vib}}(T) = \frac{k_{\text{B}}T}{\hbar} \frac{Q^{\ddagger,\text{ZPC}}}{Q^{\text{R},\text{ZPC}}} e^{-\Delta E^{\ddagger,\text{ZPC}}/k_{\text{B}}T}$$
(4)

where $Q^{\text{R,ZPC}}$ and $Q^{\ddagger,\text{ZPC}}$ are the partition functions of the precursor and transition structures, respectively, and k_{B} and \hbar are the Boltzmann and Planck constants, respectively. The activation free energies (ΔG^{\ddagger}) were derived from $k^{\text{Q-vib}}$ using $k^{\text{Q-vib}}(T) = (k_{\text{B}}T/\hbar)e^{-\Delta G^{\ddagger}/RT}$, and the activation enthalpies (ΔH^{\ddagger}) were computed using eqn (5).

$$\ln k^{\text{Q-vib}}(T) = \ln A + \frac{\Delta S^{\ddagger}}{R} - \frac{\Delta H^{\ddagger}}{RT}$$
(5)

where ΔS^{\ddagger} is the activation entropy and *R* is the gas constant. The value of ΔH^{\ddagger} was obtained from the linear relationship between $\ln k^{\text{Q-vib}}$ and 1000/T.

Based on the photoluminescence pathways shown in Fig. 4, the thermodynamics of the consecutive reaction pathway $(I)^* \rightarrow (II)^* \rightleftharpoons (III)^{\ddagger,\$} \rightarrow (IV)$ were studied, in which $(II)^* \rightleftharpoons (III)^{\ddagger,\$} \rightarrow (IV)$ was assumed to be in a quasi-equilibrium. The thermodynamic property of interest was the total Gibbs free energy $(\Delta G^{\circ,tot})$ of the reactions, computed using ΔG^{\ddagger} obtained from the TST method. $\Delta G^{\circ,tot}$ was computed by dividing the consecutive reaction pathway into two single steps, namely, $(I)^* \rightarrow (II)^*$ and $(II)^* \rightleftharpoons (III)^{\ddagger,\$} \rightarrow (IV)$. For $(I)^* \rightarrow (II)^*$, $\Delta G^{\circ,(I)^* \rightarrow (II)^*} = -\Delta G^{\ddagger,(I) \rightarrow (II)}_r$, whereas $\Delta G^{\circ,(II)^* \rightarrow (IV)} = \Delta G^{\ddagger,(II)^{\pm,(II)} \rightarrow (II)^{\ddagger,\$} \rightarrow (III)^{\ddagger,\$} \rightarrow (II)^{\ddagger,\$} \rightarrow (IV)$ and $\Delta G^{\circ,tot} = \Delta G^{\circ,(II)^* \rightarrow (II)^*} + \Delta G^{\circ,(II)^* \rightarrow (IV)}$. All the kinetic and thermodynamic properties were computed using the DL-FIND program⁴¹ included in the ChemShell software package.³⁸

Because the (I)* \rightarrow (II)* relaxation on the S₁ PES is exothermic ($\Delta H^{\circ} < 0$) and thermal energy is required for (II)* \rightleftharpoons (III)^{‡,§} \rightarrow (IV), the thermodynamic spontaneity could be studied. The spontaneous temperature ($T_{\rm s}$), below which the transition structure (III)^{‡,§} at the S₁/T₁ intersection is spontaneously formed from (I)*, was obtained from the plot of $\Delta G_r^{\ddagger,(I)} \leftarrow (II)^{\ddagger}$ and $\Delta G_f^{\ddagger,(II)} \rightarrow (III)^{\ddagger \$}$ versus temperature; (I)* \rightarrow (II)* \rightleftharpoons (III)^{‡,§} is spontaneous when $\Delta G_f^{\ddagger,(II)} \rightarrow (III)^{\ddagger,§} - \Delta G_r^{\ddagger,(I)} \leftarrow (II)^{\ddagger,§} = 0$. In other words, the formation of (III)^{‡,§} is spontaneous when $T \leq T_s$.

Surface-hopping molecular dynamics simulations

Because non-radiative relaxation reduces the emission quantum efficiency, to study the non-radiative $S_1 \rightarrow S_0$ relaxation in BF₂-FORM-D, NVE-MDSH simulations were performed. Because the calculations were computationally intensive, NVE-MDSH simulations were conducted using DFT and TD-DFT methods with a smaller (DZP) basis set, except for the iodine atoms, for which the larger TZVPall basis set⁴² was used to consider the heavy-atom effect. Fifty initial configurations were generated based on the Wigner distribution, from which NVE-MDSH simulations were performed over a time span of ~4 ps using the TURBOMOLE 7.50 software package.

The integration of Newton's equations of motion was conducted using the Verlet algorithm with a timestep of 0.5 fs, which was confirmed in our previous studies to be sufficient to study photochemical processes.²⁸ The characteristic dynamics in the S₁ states were categorized, and representative reactions were chosen and investigated in detail. To study the possibility of increasing the S₁/T₁ ISC, irradiation wavelengths, intramolecular motions, and temperatures, as well as the probabilities for the S₁/T₁ ISC and T₁ \rightarrow S₀ transition, were analyzed in detail for BF₂-FORM-D.

Results and discussion

To discuss characteristic structures of BF2-FORM and BF2-FORM-D, a three-character code is used, e.g., $G[k]^{eq}$, ${}^{1}E[k]^{\ddagger}$, or ${}^{3}E[k]^{\$}$, where G indicates the structure in the S_0 state and ${}^{1}E$ and ${}^{3}E$ indicate the structures in the S₁ and T₁ states, respectively. The terms [...]^{eq} and [...]* denote the equilibrium and vertically excited structures, respectively, whereas [...][‡] and [...][§] represent the transition structure and structure at the S_1/T_1 intersection, respectively. Different structures on the same PES are labeled as [k]. For example, ${}^{1}E$ -[1]* and ${}^{1}E$ -[2]^{eq} represent two structures on the S₁ PES. Additional symbols are used to represent the characteristic energies on the PES. In the discussion, for example, $\Delta E^{S_0 \rightarrow S_1}$ and ΔE^{\ddagger} represent the S₀ \rightarrow S₁ vertical excitation energy and energy barrier on PES, respectively, whereas $\Delta E^{S_1/T_1}$ represents the energy gap between the S_1 and T_1 states at or in the vicinity of the S_1/T_1 intersection. The terms $(...)^{S_0 \rightarrow S_1}$, $(...)^{\ddagger}$ and $(...)^{S_1/T_1}$ are used to represent the corresponding energies in the figures.

Static properties

Equilibrium structures. The equilibrium structures and energies of BF₂-FORM and BF₂-FORM-D in the S₀, S₁, and T₁ states obtained from DFT/B3LYP/6-311G and TD-DFT/B3LYP/6-311G geometry optimizations are presented in Tables S1 and S2,† respectively. Because the equilibrium structures and highest occupied molecular orbital-lowest unoccupied molecular orbital (HOMO–LUMO) of BF_2 -FORM and BF_2 -FORM-D were not significantly different, only the results for BF_2 -FORM-D are shown in Fig. 5.

The equilibrium structures of BF₂-FORM and BF₂-FORM-D in the S₀ and T₁ states were virtually identical, as shown by the bent and perfect planar structures G-[1]^{eq} and ³E-[4]^{eq}, respectively, in Fig. 5a. The HOMOs of G-[1]^{eq} and ³E-[4]^{eq} were characterized by a strong π character in the formazanate heterocyclic and phenyl rings, whereas the electron density of the LUMO was highly localized, resulting in a significantly lower degree of conjugation in the S₁ and T₁ states.

The equilibrium structures of BF₂-FORM and BF₂-FORM-D in the S₁ state were quite different. BF₂-FORM was represented by a propeller structure (Fig. S1[†]) or a twisted bent structure with the same HOMO–LUMO as G-[1]^{eq}, whereas BF₂-FORM-D was represented by a perpendicular structure, ¹E-[2]^{eq} (Fig. 5a). For ¹E-[2]^{eq}, the electron density distributions on the phenyl rings were not symmetrical because of the positive charge on the N(2) atom of the formazanate heterocyclic ring. Comparison of the vertical excitation energies of BF₂-FORM and BF₂-FORM-D in Tables S₁ and S₂[†] shows that the iodine substitutions at the phenyl rings directly affected $\Delta E^{S_0 \rightarrow S_1}$, namely, a strong red-shift is observed for BF₂-FORM-D because of a more extensive electron density distribution in HOMO $(G-[1]^{eq})$ and large charge transfer interaction within the molecule.

For BF₂-FORM-D, $\Delta E^{S_0 \to S_1} = 2.42 \text{ eV} (\lambda^{S_0 \to S_1} = 513 \text{ nm})$ was in good agreement with the absorption spectra obtained based on $\Delta E_{\rm NX}^{\rm S_0 \rightarrow S_1} = 2.32 \ \rm eV$ Wigner-sampled structures, 200 $(\lambda_{NX}^{S_0 \rightarrow S_1} = 534 \text{ nm}, \text{ shown in Fig. 5b})$ with the fluorescence lifetime, $\tau_{\rm NX}^{{\bf S}_1 \rightarrow {\bf S}_0} = 3.94 \times 10^{-9}$ s (Table S2†). The RICC2/augcc-pVDZ results confirmed the bent structure (G-[1]^{eq}) to possess $\Delta E_{\text{RICC2}}^{S_0 \rightarrow S_1} = 2.45 \text{ eV} (\lambda_{\text{RICC2}}^{S_0 \rightarrow S_1} = 506 \text{ nm})$. The calculated excitation energies/wavelengths were in good agreement with the experimental absorption spectra (e.g., in CHCl₃, $\lambda_{exp}^{S_0 \rightarrow S_1, max} = 531 \text{ nm}$).²⁴ For BF₂-FORM, the vertical excitation energies were higher, where $\Delta E^{S_0 \rightarrow S_1} = 2.88$ eV ($\lambda^{S_0 \rightarrow S_1} = 430$ nm) and $\Delta E^{S_0 \rightarrow S_1}_{NX} = 2.71$ eV ($\lambda^{S_0 \rightarrow S_1}_{NX} = 457$ nm, shown in Fig. 5b) with $\tau^{S_1 \rightarrow S_0}_{NX} = 2.95 \times 10^{-9}$ s (Table S1†). The energy values were compatible with the RICC2/aug-cc-pVDZ results, $\Delta E_{\text{RICC2}}^{S_0 \rightarrow S_1} = 2.67 \text{ eV} (\lambda_{\text{RICC2}}^{S_0 \rightarrow S_1} = 464 \text{ nm}).$ The trend of $\tau_{\text{NX}}^{S_1 \rightarrow S_0}$ was in good agreement with the experiment,24 e.g., in toluene, $\tau_{exp}^{S_1 \rightarrow S_0} = 2.26~\times~10^{-9}$ and 1.43 $~\times~10^{-9}$ s for BF2-FORM-D and BF₂-FORM, respectively.

Potential energy surfaces. The PESs for the rotations of the dihedral angles ω_1 and ω_2 in BF₂-FORM and BF₂-FORM-D, which were obtained using the DFT/B3LYP/6-311G, TD-DFT/B3LYP/6-311G, and NEB methods, are shown in Fig. S1 and S2,†



Fig. 5 (a) Structures of BF_2 -FORM-D in the S_0 , S_1 and T_1 states, obtained based on DFT/B3LYP/6-311G and TD-DFT/B3LYP/6-311G calculations. The code symbols are explained in the text. (b) Absorption spectra obtained based on 200 Wigner sampled structures.

respectively. Analysis of the PESs, shown in Fig. 6, revealed three potential deactivation pathways in which a possibility of the S₁ \rightarrow S₀ transition ([S₁ \rightarrow S₀]) is seen for the propeller structure of BF₂-FORM with $\Delta E^{S_1 \rightarrow S_0} = -2.13 \text{ eV} (\lambda^{S_1 \rightarrow S_0} = 581 \text{ nm})$, whereas $\Delta E^{S_1 \rightarrow S_0} = -1.57 \text{ eV} (\lambda^{S_1 \rightarrow S_0} = 788 \text{ nm})$ is for the perpendicular structure (¹E-[2]^{eq}) of BF₂-FORM-D. The former is the same as the experimental emission spectra of BF₂-FORM, $\lambda_{exp}^{S_1 \rightarrow S_0,max} = 581 \text{ nm}$ in CHCl₃, whereas the latter is within the range observed experimentally, $650 < \lambda_{exp}^{S_1 \rightarrow S_0,max} < 800 \text{ nm}$ for BF₂-FORM-D.²⁴

The photoluminescence pathways for BF₂-FORM and BF₂-FORM-D shown in Fig. 6 further suggest a possibility of the S₁/T₁ ISC at ¹E-[3]^{‡,§}, with the energy barriers for ¹E-[2]^{eq} \rightarrow ¹E-[3]^{‡,§}, $\Delta E^{\ddagger} = 24.5$ and 23.2 kJ mol⁻¹, and S₁/T₁ energy gaps, $\Delta E^{S_1/T_1} = 0.30$ and 0.04 eV, respectively. Because $\Delta E^{S_0 \rightarrow S_1}$ of BF₂-FORM-D is closer to the center of the visible light spectrum (550 nm) with a smaller $\Delta E^{S_1/T_1}$, BF₂-FORM-D is anticipated to be a more effective luminophore, mainly because of the heavy-atom effect. Therefore, subsequent discussions focus on

 $\mathsf{BF}_2\text{-}\mathsf{FORM}\text{-}\mathsf{D},$ with the results for $\mathsf{BF}_2\text{-}\mathsf{FORM}$ included in parentheses.

For BF₂-FORM-D, two deactivation pathways for $T_1 \rightarrow S_0$ transition, $[T_1 \rightarrow S_0]_1$ and $[T_1 \rightarrow S_0]_2$ in eqn (2) and (3), were observed after the S_1/T_1 ISC (Fig. 6). $[T_1 \rightarrow S_0]_1$ occurred immediately after the S_1/T_1 ISC, ${}^{3}E^{-}[3]^{\$} \rightarrow G^{-}[3]$ with $\Delta E^{T_1 \rightarrow S_0} = -1.44$ (-1.52) eV ($\lambda^{T_1 \rightarrow S_0} = 864$ (814) nm), whereas $[T_1 \rightarrow S_0]_2$ occurred after the ${}^{3}E^{-}[3]^{\$} \rightarrow {}^{3}E^{-}[4]^{eq}$ structural relaxation in the T_1 state, ${}^{3}E^{-}[4]^{eq} \rightarrow G^{-}[4]$ with $\Delta E^{T_1 \rightarrow S_0} = -0.87$ (-0.91) eV ($\lambda^{T_1 \rightarrow S_0} = 1430$ (1368) nm). To confirm $\Delta E^{T_1 \rightarrow S_0}$ obtained from the TD-DFT/B3LYP/6-311G method and to study the phosphorescence lifetimes, RIC-C2/aug-cc-pVDZ calculations based on the spin–orbit coupling with perturbation theory (SOC-PT-CC2) were made on ${}^{3}E^{-}[3]^{\$}$ for $[T_1 \rightarrow S_0]_1$ and on ${}^{3}E^{-}[4]^{eq}$ for $[T_1 \rightarrow S_0]_2$. The values obtained for $[T_1 \rightarrow S_0]_1$ are $\Delta E_{SOC}^{T_1 \rightarrow S_0} = -1.93(-1.94)$ eV [$\lambda = 642$ (639) nm] and $\tau_{SOC}^{T_1 \rightarrow S_0} = -0.98$ (-1.05) eV [$\lambda_{SOC}^{T_1 \rightarrow S_0} = 1265$ (1181) nm] and $\tau_{SOC}^{T_1 \rightarrow S_0} = 194$ (131) s for $[T_1 \rightarrow S_0]_2$.



Fig. 6 (a) and (b) Proposed photoluminescence pathways for BF₂-FORM and BF₂-FORM-D dye sensitizers obtained from the DFT/B3LYP/6-311G, TD-DFT/B3LYP/6-311G and NEB methods. Energies are in kJ mol⁻¹ unless specified otherwise. $(...)^{S_0 \rightarrow S_1} =$ vertical excitation energy in eV; $\Delta E^{\text{Rel}} =$ relative energy with respect to the total energy in the S₀ state; $(...)^{\text{Rel}} =$ relative energy with respect to the transition structure; $(...)^{\ddagger} =$ structure at the S₁/T₁ intersection.

Because $\Delta E^{T_1 \rightarrow S_0}$ and $\Delta E^{T_1 \rightarrow S_0}_{SOC}$ for $[T_1 \rightarrow S_0]_2$ are close to the absorption energy for ${}^{3}O_2 ({}^{3}\sum_g) \rightarrow {}^{1}O_2 ({}^{1}\Delta_g), \Delta E^{T_1 \rightarrow S_1} \approx 0.97$ eV,⁵ and $\tau^{T_1 \rightarrow S_0}_{SOC}$ is considerably longer than $[T_1 \rightarrow S_0]_1, [T_1 \rightarrow S_0]_2$ is confirmed to be a key process to drive ${}^{3}O_2 ({}^{3}\sum_g) \rightarrow {}^{1}O_2$ $({}^{1}\Delta_g)$, and BF₂-FORM-D is thus a better photosensitizer in PDT; experiments have shown that long-lived triplet excited state could promote the formation of ${}^{1}O_2$ and increases the efficiency of the PDT.¹¹ Therefore, only the kinetics and thermodynamics of $[T_1 \rightarrow S_0]_2$ are discussed further.

Thermodynamics and kinetics of the $T_1 \rightarrow S_0$ transition. The kinetic and thermodynamic results for $[T_1 \rightarrow S_0]_2$ are shown in Tables S3 and S4[†] for BF₂-FORM and BF₂-FORM-D, respectively. Because ¹E-[1]^{*} \rightarrow ¹E-[2]^{eq} are barrierless, ¹E-[2]^{eq} \rightarrow ¹E-[3]^{‡,§} could be considered the rate-determining step of the S₁/T₁ ISC. The TST results showed that BF₂-FORM was kinetically more favorable than BF₂-FORM-D; for example, at 300 K, $k_f^{Q-vib} = 5.85 \times 10^8$ and 2.45×10^4 s⁻¹, respectively.

Analysis of $\Delta G^{\circ,\text{tot}}$ in Fig. 7c shows that although $[T_1 \rightarrow S_0]_2$ was thermodynamically favorable for both BF₂-FORM-D and BF₂-FORM, *e.g.*, at 300 K, $\Delta G^{\circ,\text{tot}} = -119.5$ and -122.5 kJ mol⁻¹, respectively, the rate-determining process (II)* \rightleftharpoons (III)^{‡,§} (¹E-[2]^{eq} \rightarrow ¹E-[3]^{‡,§}) was spontaneous at room temperature only for BF₂-FORM-D; for BF₂-FORM-D, the plot of $\Delta G_r^{\ddagger,(II)^+ \rightarrow (III)^{\ddagger,§}}$ and $\Delta G_r^{\ddagger,(II)^+ \leftarrow (III)^{\ddagger,§}}$ could be spontaneous at $T \leq T_s = 320$ K, whereas the same plot did not show T_s for BF₂-FORM (Fig. 7a). These results indicate that for $[T_1 \rightarrow S_0]_2$, although BF₂-FORM was kinetically more favorable, BF₂-FORM-D was thermodynamically more favorable because the ¹E-[2]^{eq} \rightarrow ¹E-[3]^{‡,§} could be spontaneous the substantiant of the spontaneous below T = 320 K.

Surface-hopping dynamics

In order to enhance the photoluminescence quantum efficiency, it is important to study the non-radiative $S_1 \rightarrow S_0$ relaxation process in BF₂-FORM-D and the possibility of increasing the S_1/T_1 ISC. Because the NVE-MDSH simulations applied in this work considered only the non-radiative $S_1 \rightarrow S_0$ relaxation ([$S_1 \rightarrow S_0$] in Fig. 6), to explore the possibility of the S_1/T_1 ISC (1E -[3]^{‡,§} \rightarrow 3E -[3][§] in Fig. 6), the time evolutions of the total energies in the S_0 , S_1 , and T_1 states, temperatures, and dihedral angles ω_1 and ω_2 were extracted from the NVE-MDSH simulations and considered in the dynamic analysis. Characteristic results were selected as examples and are shown in Fig. 8.

The time evolutions of ω_1 and ω_2 suggest two characteristic motions of the phenyl rings; large- (L-ALM) and small-amplitude librational motions (S-ALM). L-ALM is characterized by ω_1 and ω_2 varying over a wide range, whereas S-ALM can be considered fine structures of L-ALM. To study the effect of L-ALM on the nonradiative $S_1 \rightarrow S_0$ relaxation and on the S_1/T_1 ISC, two vectors were defined on the phenyl rings. Newman projections of these two vectors and $\Delta\omega_{\text{MDSH}} = |\omega_{1,\text{MDSH}} - \omega_{2,\text{MDSH}}|$ acquired from the NVE-MDSH simulations are shown in Fig. 8.

For BF₂-FORM-D, the Newman projections and $\Delta \omega_{\text{MDSH}}$ reveal two types of L-ALM, namely, anti- and non-synchronous L-ALM, abbreviated anti-L-ALM and non-L-ALM, respectively. Because the librational motions of ω_1 and ω_2 are coupled in the S₁ state, anti-L-ALM is characterized by $\Delta \omega_{\text{MDSH}}^{S_1}$ varying uniformly across a narrow range, *e.g.*, 0° < $\Delta \omega_{\text{MDSH}}^{S_1}$ < 68°, shown in Fig. 8a and b, whereas $\Delta \omega_{\text{MDSH}}^{S_1}$ for non-L-ALM varies



Fig. 7 (a) and (b) Plots of $\Delta G_r^{\ddagger,(1)^* \leftarrow (11)^*}$ and $\Delta G_f^{\ddagger,(11)^* \leftarrow (111)^{\ddagger, \$}}$ as a function of *T* for BF₂-FORM and BF₂-FORM-D, respectively. T_s = spontaneous temperature below which the reaction is spontaneous. (c) Gibbs free energies for the photochemical deactivation processes (Fig. 4) for BF₂-FORM at 300 K. [...] = values for BF₂-FORM-D.



Fig. 8 Examples of time evolutions of energies in the S₀, S₁ and T₁ states, temperatures, and dihedral angles ω_1 and ω_2 , obtained from NVE-MDSH simulations on BF₂-FORM-D. The Newman projections at the bottom of the figures are used to differentiate the anti- and non-synchronous large-amplitude librational motions, anti-L-ALM and non-L-ALM, respectively. (a) and (b) anti-L-ALM. (c) and (d) non-L-ALM. $\tau_{S_1 \to S_0}^{S_1 \to S_0} = S_1 \to S_0$ surface hopping time; T_{av} = the average temperature; ω_1 and ω_2 = dihedral angles; T_s = spontaneous temperature; (S₁/S₀) = structure at $\tau_{MDSH}^{S_1 \to S_0}$.

across a wider range, *e.g.*, $0^{\circ} < \Delta \omega_{\text{MDSH}}^{S_1} < 85^{\circ}$, shown in Fig. 8c and d. Analysis of the NVE-MDSH results shows that short $S_1 \rightarrow S_0$ surface-hopping times $(\tau_{\text{MDSH}}^{S_1 \rightarrow S_0})$ are associated with anti-L-ALM (*e.g.*, $\tau_{\text{MDSH}}^{S_1 \rightarrow S_0} < 630$ fs, shown in Fig. 8a and b), whereas non-L-ALM dominates for long $\tau_{\text{MDSH}}^{S_1 \rightarrow S_0}$ (*e.g.*, $\tau_{\text{MDSH}}^{S_1 \rightarrow S_0} > 1100$ fs, shown in Fig. 8c and d).

Because the NVE-MDSH simulations do not directly account for $S_1 \rightarrow S_0$ fluorescence and $T_1 \rightarrow S_0$ phosphorescence, further structural, energetic, and dynamic analyses must be performed for BF₂-FORM-D. Based on the hypothesis that the non-radiative $S_1 \rightarrow S_0$ relaxation occurs in an ultrashort $\tau_{MDSH}^{S_1 \rightarrow S_0}$, the probability of the $S_1 \rightarrow S_0$ fluorescence ($\tau_{NX}^{S_1 \rightarrow S_0} = 3.94 \times 10^{-9}$ s), S_1/T_1 ISC and $T_1 \rightarrow S_0$ phosphorescence ($\tau_{SOC}^{T_1 \rightarrow S_0} = 194$ s) could increase when $\tau_{MDSH}^{S_1 \rightarrow S_0}$ is sufficiently long. Therefore, the factors that could affect the length of $\tau_{MDSH}^{S_1 \rightarrow S_0}$ were studied during NVE-MDSH simulations. Likewise, because the S_1/T_1 energy degeneration is one of the preconditions for the S_1/T_1 ISC and $T_1 \rightarrow S_0$ phosphorescence, the factors affecting the duration of the S_1/T_1 energy degeneration were monitored during NVE-MDSH simulations. The S_1/T_1 energy degenerations are clearly seen in Fig. 8c and d (yellow stripes), for which long S_1/T_1 degeneration times $(\tau_{MDSH}^{S_1/T_1})$ are associated with non-L-ALM. The time evolutions of the S_1 and T_1 total energies also suggest that for non-L-ALM, each $\tau_{MDSH}^{S_1/T_1}$ could span between $704 < \tau_{MDSH}^{S_1/T_1} < 1153$ fs, compared with $135 < \tau_{MDSH}^{S_1/T_1} < 195$ fs for anti-L-ALM. In other words, the higher the probability of the S_1/T_1 ISC and $T_1 \rightarrow S_0$ phosphorescence.

To study the effect of the absorbed radiation energy (the $S_0 \rightarrow S_1$ excitation energy) on the non- and anti-L-ALM modes, the correlations among $\Delta E^{S_0 \rightarrow S_1}$ of the Wigner-sampled structures, average temperatures in the S_1 state $(T_{Av,MDSH}^{S_1})$, $S_1 \rightarrow S_0$ surface hopping times $(\tau_{MDSH}^{S_1 \rightarrow S_0})$, and $\Delta \omega_{MDSH}$ of the structures at $\tau_{MDSH}^{S_1 \rightarrow S_0}$ ($\Delta \omega_{MDSH}^{S_1/S_0}$) obtained from all NVE-MDSH simulations are plotted in Fig. 9. The results show that for anti-L-ALM, $\tau_{MDSH}^{S_1 \rightarrow S_0}$ can be categorized into two groups, namely, $\tau_{MDSH}^{S_1 \rightarrow S_0} < 500$ fs and $500 < \tau_{MDSH}^{S_1 \rightarrow S_0} < 1000$ fs, whereas $\tau_{MDSH}^{S_1 \rightarrow S_0} > 1000$ fs is for non-L-ALM; these are indicated as (a), (b), and (c) in Fig. 9a, respectively.





Fig. 9 (a) Correlation between $\tau_{\text{MDSH}}^{S_1 \to S_0}$ and $\Delta E^{S_0 \to S_1}$. (b) Correlation between $\tau_{\text{MDSH}}^{S_1 \to S_0}$ and $T_{\text{AV,MDSH}}^{S_1}$. (c) Correlation between $\Delta \omega_{\text{MDSH}}^{S_1/S_0}$ and $\Delta E^{S_0 \to S_1}$. (b) Correlation between $\tau_{\text{MDSH}}^{S_1 \to S_0}$ and $T_{\text{AV,MDSH}}^{S_1}$. (c) Correlation between $\Delta \omega_{\text{MDSH}}^{S_1/S_0}$ and $\Delta E^{S_0 \to S_1}$. (c) Correlation between $\Delta \omega_{\text{MDSH}}^{S_1/S_0}$ and $\Delta E^{S_0 \to S_1} = S_0 \to S_1$ excitation energy; $T_{\text{av,MDSH}}^{S_1}$ = average temperature in the S_1 state; $\Delta \omega_{\text{MDSH}}^{S_1/S_0}$ = different between ω_1 and ω_2 at $\tau_{\text{MDSH}}^{S_1 \to S_0}$.

It appears in Fig. 9a that anti-L-ALM (a) occurs exclusively in the absorbed radiation energy range of $2.30 < \Delta E^{S_0 \rightarrow S_1} < 2.40 \text{ eV}$, whereas anti-L-ALM (a) and (b), and non-L-ALM (c) could be concurrently found when $\Delta E^{S_0 \rightarrow S_1} > 2.40 \text{ eV}$. The asymptotic behaviors of $\Delta E^{S_0 \rightarrow S_1}$, $T_{Av,MDSH}^{S_1}$, and $\Delta \omega_{MDSH}^{S_1/S_0}$ in Fig. 9 suggest that to increase the probability of long $S_1 \rightarrow S_0$ surface-hopping time, the absorbed radiation energy should be $\Delta E^{S_0 \rightarrow S_1} \approx 2.67 \text{ eV}$ or $\lambda^{abs} \approx 464 \text{ nm}$, corresponding to blue light source (Fig. 5b) with $T_{Av,MDSH}^{S_1} \approx 600 \text{ K}$ and $\Delta \omega_{MDSH}^{S_1/S_0} \approx 130^\circ$.

These results suggest that the absorbed radiation energy and intramolecular librational motions govern the non-radiative $S_1 \rightarrow S_0$ relaxation process and time, and non-L-ALM could increase the probability of fluorescence, S_1/T_1 ISC and phosphorescence. Because the TST results suggest that the S_1/T_1 ISC is thermodynamically favorable at $T_s < 320$ K and because the asymptotic average temperature for long $S_1 \rightarrow S_0$ surface-hopping time is $T_{Av,MDSH}^{S_1} \approx 600$ K, to thermodynamically enhance the probability of photoluminescence, a temperature control mechanism is required.

Conclusion

Photodynamic therapy (PDT) is a promising medical treatment for a range of human diseases, in which one of the key factors in its effectiveness is how well the photosensitizer can transfer photon energy to the target molecules. In this study, to improve the efficiency of photosensitizers for PDT, theoretical methods were used to study the photoluminescence mechanisms of BF₂formazanate dye (BF₂-FORM) and its iodinated derivative (BF₂-FORM-D) to investigate the heavy-atom effect. To complete this mechanistic study, complementary theoretical approaches were applied to investigate three important issues; (1) luminescence pathways in the S₁ and T₁ states, studied using DFT and TD-DFT methods, (2) kinetic and thermodynamic properties of the proposed pathways, studied using the TST method, and (3) time evolution and dynamics of the key processes, studied using NVE-MDSH simulations.

The DFT/B3LYP/6-311G and TD-DFT/B3LYP/6-311G results showed that in the S₀ and T₁ states, the equilibrium structures of BF₂-FORM and BF₂-FORM-D were similar, represented by bent and perfect planar structures, respectively, whereas in the S₁ state, the equilibrium structure of BF₂-FORM took a propeller structure and that of BF₂-FORM-D a perpendicular structure. The HOMOs of the equilibrium structures in the S₀ state were characterized by strong π character at the formazanate heterocyclic and phenyl rings, whereas electron density distribution in LUMOs was localized, and iodine substitutions at the phenyl rings led to a strong red-shift of $\Delta E^{S_0 \rightarrow S_1}$ close to the center of the visible light spectrum (green light with maximum intensity at 520 < $\lambda^{S_0 \rightarrow S_1} < 532$ nm).

The PESs for rotations of the dihedral angles ω_1 and ω_2 suggested two mechanisms for $T_1 \rightarrow S_0$ transition: $[T_1 \rightarrow S_0]_1$

occurring immediately after S_1/T_1 ISC and $[T_1 \rightarrow S_0]_2$ occurring after S_1/T_1 ISC and T_1 equilibrium structure relaxation. $[T_1 \rightarrow S_0]_2$ transition ($\Delta E^{T_1 \rightarrow S_0}$) is in the near IR range and close to the absorption energy for ${}^{3}O_2 ({}^{3}\sum_g) \rightarrow {}^{1}O_2 ({}^{1}\Delta_g)$. Because $\Delta E^{S_0 \rightarrow S_1}$ of BF₂-FORM-D is closer to the center of the visible light spectrum and $\Delta E^{S_1 \rightarrow T_1}$ is significantly smaller than BF₂-FORM, the iodinated derivative is anticipated to be a more effective luminophore, mainly owing to the heavy-atom effect. The RICC2/ aug-cc-pVDZ (SOC-PT-CC2) calculations confirmed these findings and further suggested that the phosphorescence lifetime of $[T_1 \rightarrow S_0]_2$ of BF₂-FORM-D is significantly longer than that of BF₂-FORM.

Based on the PESs obtained using the DFT/B3LYP/6-311G, TD-DFT/B3LYP/6-311G, and NEB methods, TST calculations confirmed that $[T_1 \rightarrow S_0]_2$ of BF₂-FORM-D is thermodynamically favorable below the spontaneous temperature ($T_s = 320$ K). The time evolutions of the dihedral angles ω_1 and ω_2 obtained from the analysis of NVE-MDSH simulations suggested that anti-L-ALM underlies ultrafast non-radiative $S_1 \rightarrow S_0$ relaxation, whereas non-L-ALM could enhance the probability of S₁/T₁ ISC and photoluminescence. Therefore, to delay non-radiative $S_1 \rightarrow$ S₀ relaxation and increase the probability of photoluminescence, anti-L-ALM of the phenyl rings should be promoted. Analysis of the NVE-MDSH results suggested that the photoluminescence quantum yield could also be enhanced by varying the irradiation wavelength, for example, by using a blue light source to promote phosphorescence. Because efficient delivery of the photosensitizer to target cells is also a critical factor in PDT, our upcoming theoretical study will examine the photodynamic properties of BF2-FORM-D when incorporated into a specific nanostructure. These results will serve as a foundation for future theoretical and experimental research, to optimize and/or design effective PDT photosensitizers.

Data availability

The data that support the findings of this study are available in the ESI† of this article.

Conflicts of interest

There are no conflicts to declare.

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