Investigation on fluorescein derivatives with thermally activated delayed fluorescence and their applications in imaging†

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Fluorescein derivatives with thermally activated delayed fluorescence (TADF) show much stronger competition ability and vaster prospects than traditional fluorescein dyes due to their prominent long lifetime. It will be of great significance to synthesize more fluorescein derivatives with TADF. In this work, compounds DCF-MPYA and FL with TADF properties were obtained by fine tuning the substituents' structure on the basis of fluorescein derivative DCF-MPYM. Their long-lived triplet excited states (21.78 μs, 32.0 μs) were proved by nanosecond time-resolved transient absorption spectra. The steady-state and time-resolved fluorescence spectra showed that DCF-MPYA and FL exhibited red fluorescence around 645 nm and 651 nm, respectively. The results of sensitivity to oxygen and heavy atoms further demonstrated that the time-resolved fluorescence spectra originate from the delayed fluorescence. The time correlated single-photon counting (TCSPC) data indicated that DCF-MPYA and FL showed long-lived lifetimes of 13.16 μs and 23.72 μs, respectively. The energy gap (ΔE) between the singlet (S1) and triplet (T1) states of DCF-MPYA and FL was calculated to be 3.32 meV and 9.98 meV from the decay rate of DF as a function of temperature. The small energy gap is conducive to the occurrence of efficient TADF at room temperature. Meanwhile, Gaussian calculation was employed to observe the electron density of DCF-MPYA and FL in the ground and excited states. The calculation results indicate that the shapes and energy levels of the highest occupied molecular orbitals (HOMOs), lowest unoccupied molecular orbitals (LUMOs), and LUMOs+1 for the monoanion and dianion forms are similar and thus DCF-MPYA and FL exhibit almost the same luminescence properties. Finally, DCF-MPYA and FL with low toxicity were used in confocal and time-resolved fluorescence imaging. Our construction strategy will be beneficial for developing more fluorescein derivatives with TADF in the future.

Introduction

Fluorescein dyes were first obtained from resorcinol and phthalic anhydride by von Bayer under the catalysis of zinc chloride in 1871. They have the advantages of large molar extinction coefficient,† high fluorescence quantum yield,‡ easy synthesis and purification,§ and low environmental and biological toxicity.¶ Therefore, fluorescein is one of the fluorescent dyes which is widely used in the development of fluorescent probes.† However, the traditional fluorescein dyes still have some disadvantages, such as: (1) the absorption and emission wavelengths for the designed fluorescent probes based on fluorescein dyes are mainly concentrated in the visible band, which limits the application of these fluorescent probes in the fields of clinical laboratory diagnosis and chemical biology;‡ (2) the fluorescein dyes have small Stokes shift (<25 nm), which causes fluorescence self-quenching and errors because of the Rayleigh scattering;¶ (3) the fluorescence lifetime (nanosecond scale) of fluorescein dyes is short, which brings about the interfere of background fluorescence.§ Therefore, it is of great significance to develop fluorescein dyes with long emission wavelength, large Stokes shift and long-lived fluorescence lifetime.

In 1961, Parker and Hatchard discovered that eosin has the TADF property with long fluorescence lifetime.§ Compared with traditional rare earth and transition metal complexes, eosin not only has long fluorescence lifetime but also has better biocompatibility. Thus, this excellent property has greatly attracted the scientific researchers’ attention. Following the dye

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eosin, Song group firstly introduced substituents at the position 4’ and 5’ of 2’, 7’-dichlorofluorescein (DCF) matrix through Duff and Knoevenagel reactions.4 And they obtained the dye DCF-MPYM with long wavelength emission (λem > 590 nm), large Stokes shift (>150 nm) and long-lived TADF lifetime (22.11 μs in deoxy ethanol).4 According to the characteristics of long-lived delayed fluorescence of DCF-MPYM, it was used in time-resolved fluorescence imaging to achieve the effect of no background fluorescence interference. Based on this derivation, this group has done a lot of application researches on the performance of TADF.

Among them, the group has done a lot of application researches on the performance of TADF. The diagnosis and treatment were designed and synthesized on the basis of TADF fluorescence derivative DCF-MPYM.6 They can be used for specific fluorescence diagnosis and photodynamic therapy of hypoxic tumours with overexpression of nitroreductase. In 2018,11 based on the fluorescence resonance energy transfer (FRET) mechanism, they developed the ratio photosensitizer FL-CyN by utilizing DCF-MPYM as energy donor and cyanine as energy acceptor. In 2019,12 they constructed dual targeted diagnosis and treatment integration photosensitizer FL-RGD through covalently coupling TADF fluorescence derivative FL with cyclic tripeptide RGD, which was successfully applied to the integration of tumour diagnosis and treatment. Therefore, the above results show that developing more fluorescence derivatives is significant for the future research of fluorescence diagnosis and photodynamic therapy.

Firstly, the UV-vis absorption and steady-state emission spectra of DCF-MPYA and FL were studied (Fig. S1 and S2†). Second, by using the steady-state fluorescence spectroscopy, time-resolved fluorescence spectroscopy, nanosecond transient absorption technology and fluorescence lifetime. All the results indicated that both DCF-MPYA and FL showed long-lived lifetimes and TADF. Besides, the density functional theory (DFT) was used to observe the electron density of DCF-MPYA and FL at the ground and excited state. The calculation results show that the shapes and energy levels of HOMOs, LUMOs, and LUMOs+1 for monoanion and dianion forms are similar. Thus, it explains the reason that DCF-MPYA and FL exhibit almost same luminescent properties through the analysis of electron density. The aforementioned results show that fine tuning the structures of dyes with TADF property is helpful to construct new TADF dyes. Our constructing strategy of fluorescent dyes and the research methods will be conducive to the development of more organic dyes with TADF in the future.

Materials and synthesis

Materials

Common reagents used in the experiments were all of analytical grade. Urotropine, trifluoroacetic acid, piperdine, 2-cyano-2-(2,6-dimethyl-4h-pyran-4-ylidene) ethyl acetate, and 2,6-dimethyl-4h-pyran-4-ylidene malononitrile are purchased from Aladdin. All the reactions were carried out under nitrogen (N2) atmosphere with dry, freshly distilled solvents under anhydrous conditions, unless otherwise noted. Silica gel (100-200 mesh) was used for flash column chromatography.

Synthesis

Compound DCF-MPYA. To a solution of DCF dialdehyde (0.159 g, 0.348 mmol) in dry EtOH (50 mL)/CH3OH (10 mL), ethyl 2-cyano-2-(2,6-dimethyl-4h-pyran-4-ylidene) ethyl acetate (0.147 g, 0.670 mmol) and piperidine (0.172 g, 2.00 mmol) were added under N2 atmosphere. The mixture was refluxed for 24 h under N2 atmosphere at 85 °C. After finishing reaction, the reaction mixture was allowed to cool to 25 °C, then the solvent was evaporated in vacuum. After being purified by column chromatography on silica gel (elution: methanol/dichloroethane from 0.1 : 250 to 1 : 250), compound DCF-MPYA was obtained as black red powder (0.130 g, 0.152 mmol, 22.7% yield). Rf = 0.38 (5% CH3OH in dichloromethane).

ΔH NMR (400 MHz, CD3OD) δ 8.23 (d, J = 6.7 Hz, 1H), 8.17–8.04 (m, 2H), 7.88 (d, J = 7.6 Hz, 1H), 7.81–7.68 (m, 4H), 7.68–7.54 (m, 1H), 7.46 (d, J = 7.9 Hz, 1H), 7.04 (s, 2H), 6.48 (dd, J = 21.2, 9.3 Hz, 2H), 4.32–4.15 (m, 2H), 4.11 (dd, J = 10.4, 7.1 Hz, 2H), 2.39 (d, J = 6.4 Hz, 3H), 2.27 (d, J = 7.1 Hz, 3H), 1.35 (td, J = 7.1, 2.0 Hz, 3H), 1.27 (td, J = 7.0, 3.1 Hz, 4H), 1H.

Compound FL. The synthesis method of intermediate and FL were according to the reported method.12

Results and discussion

Photophysical properties and nanosecond time-resolved transient difference absorption spectra of DCF-MPYA and FL

Firstly, the UV-vis absorption and steady-state emission spectra of DCF-MPYA and FL were studied (Fig. S1 and S2†) and...
absorption spectra of DCF-MPYA (2.0 × 10⁻⁵ M in deaerated CH₃CN). Black line: steady-state fluorescence spectrum under air atmosphere (delay time, 0 s). Red line: time-resolved fluorescence spectrum under N₂ atmosphere. All the delayed detection was carried out in phosphorescence mode (total decay time, 5 ms; delay time, 0.1 ms; gate time, 1.0 ms), and the excitation wavelength was 465 nm.

summarized in Tables S1 and S2. † DCF-MPYA and FL show two absorption peaks, both of them have large Stokes shifts (≥144 nm) in different organic solvents and exhibit slightly shorter in water (≥95 nm) (Tables S1 and S2). However, the Stokes shifts of both dyes are much larger than those of conventional Rhodamine, Fluorescein and Rhodol dyes. Thus, it can be observed that the photophysical properties were similar with the reported compound DCF-MPYM in different solvents. Since the structures of DCF-MPYA and FL are analogous to DCF-MPYM, it is inferred that the luminescent characteristics of DCF-MPYA and FL may be consistent with those of DCF-MPYM. Thus, the spectra of two dyes will be studied in detail.

It was reported that the fluorescein derivative DCF-MPYM emits TADF and the most essential reason is the production of long-lived triplet state. Therefore, it was expected that the DCF-MPYA and FL also can exhibit the same characteristic with DCF-MPYM. First, nanosecond time-resolved transient difference absorption spectra of DCF-MPYA and FL were carried out to prove whether the T₁ state of the fluorescein derivatives existed. As seen from Fig. 1, upon pulsed laser excitation at 532 nm, significant bleaching of the transient absorption bands was observed at about 400, 466, 561, and 650 nm in the range of 300–700 nm. The bleaching peaks at 466 nm and 561 nm were attributed to the depletion of the ground state of DCF-MPYA. Transient absorption bands at about 400 nm and 650 nm were due to the transient absorption of T₁ for DCF-MPYA. The T₁ lifetime for the transients of DCF-MPYA was determined as 21.78 μs in deaerated ethanol (Fig. S3A†). Similar transient spectra were also observed for FL in Fig. S4† and the T₁ lifetime of FL was estimated as 32.0 μs (Fig. S3B†). These results indicate that both the compounds DCF-MPYA and FL exhibit long-lived triplet excited state upon visible light excitation, therefore, the existence of long lifetime triplet state will provide prerequisite for the TADF property of DCF-MPYA and FL.

Luminescence spectra of DCF-MPYA and FL

Next, to investigate whether the luminescence spectra of DCF-MPYA and FL were related to the long-lived triplet lifetime, the steady-state and time-resolved fluorescence spectra were studied and recorded by using fluorescence spectrophotometer. As displayed in Fig. 2, a strong peak at 645 nm was observed in time-resolved fluorescence spectrum (delayed fluorescence, DF), which closely agreed with the prompt fluorescence (PF, 634 nm) in steady-state fluorescence spectrum at room temperature. It can be found that the delayed spectrum of DCF-MPYA showed a little red-shift in comparison with the steady-state fluorescence spectrum, and similar results were also observed on FL (Fig. S5†). The maximum steady-state and time-resolved emission wavelengths of FL peaked at about 649 nm and 651 nm, respectively. The red-shift phenomenon was observed in the previous report. † It can be attributed to the difference of nuclear configuration between the singlet and triplet excited states of DCF-MPYA and FL. According to the above results, it may be inferred that the prompt and delayed fluorescence spectra simultaneously contribute to the long wavelength region of DCF-MPYA and FL. Meanwhile, the time-resolved fluorescence spectra were relevant with their triplet state lifetimes.

As we all know, the triplet state is sensitive to oxygen and heavy atom. †† Thus, the time-resolved spectra will be affected by the deoxygenation and addition of substances containing heavy atoms. To clarify the origination from the delayed component, the time-resolved spectra of DCF-MPYA and FL were also measured with and without N₂ bubbling. It was found that the luminescence intensity of DCF-MPYA at 645 nm enhanced rapidly after deoxygenation (Fig. 3), and the measured photoluminescence intensity of DCF-MPYA in N₂ rich environment has 20 times enhancement than that in air atmosphere. Meanwhile, FL also shows the similar result, the luminescence intensity at 651 nm has 4 times enhancement after deoxygenation (Fig. S6†). In addition, the steady-state fluorescence spectra of DCF-MPYA and FL were also recorded after deoxygenation (Fig. S7†). It was observed that the steady-state fluorescence spectra had no significant changes whether in non-deaerated or deaerated atmosphere. Furthermore, the delayed fluorescence spectra were also obtained after the addition of ethyl iodide into the DCF-MPYA and FL nitrogen-saturated solution (Fig. 4 and S8†). As we expected, it was noted that the time-resolved fluorescence spectra showed much enhancement.
second-order exponential decay model in N\textsubscript{2} atmosphere. As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9, time scale were measured by time correlated single-photon counting (TCSPC).

DCF-MPYA and FL were excited at 645 nm (nanosecond pulsed OPO laser synchronized with spectrophotometer) and monitored at 645 and 651 nm, respectively.

Table 1. Fluorescent lifetime compositions of delayed components for DCF-MPYA and FL in deaerated CH\textsubscript{3}CN. Excited at 485 nm (nanosecond pulsed OPO laser synchronized with spectrophotometer) and monitored at 645 and 651 nm, respectively.

| Compound       | \(\tau_i^a\) (\(\mu\)s) | \(n_i^b\) % | \(\tau_m^b\) (\(\mu\)s) | \(n_m^b\) % | \(\tau_a^b\) (\(\mu\)s) |
|----------------|-----------------|-------------|-----------------|-------------|-----------------|
| DCF-MPYA       | 0.92            | 40.73       | 21.58           | 59.27       | 13.16           |
| FL             | 0.61            | 56.69       | 53.97           | 43.31       | 23.72           |

\(^a\) Obtained from the double-exponential fitting of transient decay curves on an 800 \(\mu\)s scale. \(^b\) The contribution of each component to average lifetime.

DCF-MPYA and FL exhibit no obvious enhancement (Fig. S7†). Thus, the above results further proved that the delayed emission at 645 nm and 651 nm results from the T\(_1\) state of the derivatives and the time-resolved fluorescence can be attributed to TADF.

**Luminescence lifetime measurement of DCF-MPYA and FL**

It is known that the lifetime of delayed fluorescence can extend microsecond scale.\(^{15–17}\) Here, the photoluminescence lifetime (PL) of DCF-MPYA and FL for nanosecond and microsecond time scale were measured by time correlated single-photon counting (TCSPC). As shown in Fig. S9–S12,† both the nanosecond and microsecond scale lifetimes of DCF-MPYA exhibit second-order exponential decay modal in N\textsubscript{2} atmosphere (Tables S3† and 1). The nanosecond scale lifetime of fast decay component at 634 nm was 4.20 ns in oxygen-free environment (Table S3†) and the delayed component decay of DCF-MPYA after deoxyanation complete within 13.16 \(\mu\)s (Table 1). After deaeration, the fluorescent lifetimes of FL for nanosecond and microsecond scale were 3.67 ns and 23.72 \(\mu\)s (Tables 1 and S4†), respectively. According to the results of lifetime, it can be concluded that the delayed lifetimes of DCF-MPYA and FL enhanced at least 1000 times in comparison with traditional fluorescent dyes. Thus, the long wavelength emission can be attributed to the spectral overlap of PF and DF. In other words, the DF at long wavelength region exhibits almost same spectrum as the PF based on the time-resolved and steady-state fluorescence.\(^8\) Therefore, there is a fluorescence intensity ratio \((I_{DF}/I_{PF})\) between PF and DF. According to the reported calculation strategy,\(^{8,19}\) the intensity ratio \(I_{DF}/I_{PF}\) of DCF-MPYA and FL can be approximately calculated to be 3.07 at 645 nm and 6.97 at 651 nm, respectively. This result suggests that the long wavelength region predominantly originates from the slow decay of long-lived DF component.

The energy difference estimation between \(S_1\) and \(T_1\) of DCF-MPYA and FL

Literature reported that the small energy difference between the \(S_1\) and \(T_1\) plays a significant role in the realization of reverse intersystem conversion (RISC) in pure organic fluorescent dyes.\(^8\) It is the prerequisite to realize efficient TADF. The energy difference can be quantitatively calculated by the equation:

\[
k_{ISC}^{T \rightarrow S} = k_{ISC}^{S \rightarrow T} \exp(-\Delta E_{ST}/k_B T)
\]  

where \(k_B\) is Boltzman’s constant, \(T\) is temperature, and \(k_{ISC}^{T \rightarrow S}\) is the average rate constant for ISC. The lifetime of DF is mainly determined by RISC \((k_{ISC}^{T \rightarrow S})\) and thus the decay rate of DF can be used to represent \(k_{ISC}^{T \rightarrow S}\) for \(\Delta E_{ST}\) estimation. Here, the temperature-dependent long lifetimes for DCF-MPYA and FL were measured at 640 nm from 77 K to 300 K (Tables S5 and S6†). Then the logarithm of the fluorescence decay rate vs. the reciprocal of temperature for DCF-MPYA and FL was obtained in Fig. 5 and S13,† the red fitting curves were obtained on the basis of eqn (1). The energy gap between the \(S_1\) and \(T_1\) \((\Delta E_{ST})\) was calculated to be 3.32 meV for DCF-MPYA and 9.98 meV for FL, respectively. The small energy gap is profitable for efficient TADF.\(^{20}\) The results were consistent with our previously reported results of DCF-MPYM, which fully proves that the luminescence mechanism of DCF-MPYA and FL is the same as the reported DCF-MPYM.\(^8\)
emission. DMSO; they are involved in vertical excitation (UV-vis absorption) and energy levels were calculated and presented in Fig. 6. All the electron density of HOMO for them mainly locates on the appendant part, xanthene, and benzene ring moiety. HOMO and S18, LUMO, and LUMO+1 energy levels were calculated and presented in Fig. 6. Meanwhile, the HOMO, LUMO, and LUMO+1 energy levels were calculated and presented in Fig. 6 and S18–S20. At the ground and excited state, the electron densities on the HOMO, LUMO, and LUMO+1 of [DCF-MPYA-H]−, [DCF-MPYA-2H]2−, [FL-H]−, and [FL-2H]2− distribute between appendant part, xanthene, and benzene ring moiety. All the electron density of HOMO for them mainly locates on the xanthene unit, the LUMO+1 electron density is positioned at the partial xanthene moiety as well as appendant part. The LUMO electron density for [DCF-MPYA-H]− and [FL-H]− mainly locates at xanthene unit and benzene ring, but electron density of LUMO for [DCF-MPYA-2H]2− and [FL-2H]2− distributes at xanthene unit and unilateral substituent. Despite the slightly structural difference between DCF-MPYA and FL, the shapes and energy levels of the HOMOs, LUMOs, and LUMOs+1 for monoanion and dianion forms are similar, which determines that the calculated absorption and emission wavelengths for DCF-MPYA and FL exhibit almost same (Tables S7–S10†). This also fully explains the reason that the two dyes actually have the same luminescence characteristics.

DFT calculations of DCF-MPYA and FL

To gain insight into the optical properties of DCF-MPYA and FL, their DFT calculations were performed using Gaussian 09 program. According to the reported literature,23–25 the luminescence contribution of DCF derivatives may originate from the monoanion and dianion forms. Thus, the thermodynamically stable configurations for [DCF-MPYA-H]− (Fig. S14†), [DCF-MPYA-2H]2− (Fig. S15†), [FL-H]− (Fig. S16†), and [FL-2H]2− (Fig. S17†) were initially performed with empirical dispersion corrected PBE0 functional (D3BJ damping) and the 6-31G(d) basis set. The optimized conformation shows that the two appendant parts at the position of 4' and 5' exhibit a remarkable twist from xanthene moiety and appendant parts are parallel to each other. Meanwhile, the HOMO, LUMO, and LUMO+1 energy levels were calculated and presented in Fig. 6 and S18–S20. At the ground and excited state, the electron densities on the HOMO, LUMO, and LUMO+1 of [DCF-MPYA-H]−, [DCF-MPYA-2H]2−, [FL-H]−, and [FL-2H]2− distribute between appendant part, xanthene, and benzene ring moiety. All the electron density of HOMO for them mainly locates on the xanthene unit, the LUMO+1 electron density is positioned at the partial xanthene moiety as well as appendant part. The LUMO electron density for [DCF-MPYA-H]− and [FL-H]− mainly locates at xanthene unit and benzene ring, but electron density of LUMO for [DCF-MPYA-2H]2− and [FL-2H]2− distributes at xanthene unit and unilateral substituent. Despite the slightly structural difference between DCF-MPYA and FL, the shapes and energy levels of the HOMOs, LUMOs, and LUMOs+1 for monoanion and dianion forms are similar, which determines that the calculated absorption and emission wavelengths for DCF-MPYA and FL exhibit almost same (Tables S7–S10†). This also fully explains the reason that the two dyes actually have the same luminescence characteristics.

The confocal and time-resolved fluorescence imaging of DCF-MPYA and FL

Based on the excellent photophysical properties of DCF-MPYA and FL, they can be expanded to the biological application. First, low cytotoxicity of organic fluorophores is one of the important factors for live cells imaging.24 MTT assays were performed in the MCF-7 cell lines by monitoring the absorbance of formazan at 570 nm to confirm the low cytotoxicity of DCF-MPYA and FL. As shown in Fig. 7, the low cytotoxicity of them in vitro was observed after different incubation time. Even the incubation time was extended to 10 h, the cytotoxicity was moderately low for cells. Then the fluorescent images of DCF-MPYA and FL were performed on confocal fluorescence microscope. The results showed that both of them have good biocompatibility and they can well stain the MCF-7 cell lines (Fig. 8), thus they exhibit promising potential in biological application. To utilize the advantages of long-lived lifetime of DCF-MPYA and FL, the time-resolved fluorescence images of DCF-MPYA and FL were also taken from the laboratory-used time-domain luminescence microscope. As shown in Fig. 9, DCF-MPYA and FL-loaded MCF-7 cells emitted clearly red luminescence after 15 μs delay time. Thus, the aforementioned results manifested that DCF-MPYA and FL with TADF characteristic will be good candidates for fluorescent imaging.

**Fig. 5** Logarithm of the fluorescence decay rate vs. the reciprocal of temperature for DCF-MPYA; red line represents the fitting curves that obtained on the basis of eqn (1).

**Fig. 6** The frontier molecular orbital plots of [DCF-MPYA-H]− in DMSO; they are involved in vertical excitation (UV–vis absorption) and emission.

**Fig. 7** Biological toxicity of MCF-7 cells in the presence of DCF-MPYA (20 μM), and FL (20 μM) was determined after different incubation time. Mean values and standard deviations are obtained from three independent experimental determinations.
In summary, two red-emission fluorescein derivatives (DCF-MPYA and FL) with TADF property were synthesized on the basis of fine tuning the substituents’ structure of DCF-MPYM. Luminescence characteristics of them were investigated by the nanosecond time-resolved transient difference absorption spectra, steady-state and time-resolved fluorescence spectra. The results indicated that DCF-MPYA and FL exhibited long-lived T1 lifetime and delayed fluorescence. Then the lifetimes of delayed fluorescence for them were measured to be microsecond scale by TCSPC. Besides, the low energy gap between S1 and T1 was estimated to prove the reason for the occurrence of TADF. The DFT calculation results show that the shapes and energy levels of HOMOs, LUMOs, and LUMOs+1 for DCF-MPYA-H, DCF-MPYA-2H, FL-H, and FL-2H are similar, which explains the reason that DCF-MPYA and FL exhibit almost same luminescent properties. DCF-MPYA and FL were finally used in the confocal and time-resolved fluorescence imaging of MCF-7 cells, which proves that DCF-MPYA and FL will have great potentials in biological applications. This work manifests that fine tuning the structures of dyes with TADF property is helpful and provides a good platform for the development of more organic dyes with TADF in the future.

Author contributions

Z. Lv and X. Xiong contributed equally to this work. Zhaoye Lv: data curation, analytical measurements, original draft preparation and revision. Jun Hou: compounds’ synthesis, data curation, original draft preparation. Junjie Yao: analytical measurements, data curation. Ye Yuan: analytical measurements, data curation. Yuanlan Qian: analytical measurements, data curation. Junyang Zhu: analytical measurements, data curation. Hongjuan Zhao: writing-reviewing. Xiaoning Xiong: supervision and validation, methodology, conceptualization, writing-reviewing, and editing. Chengqi Jiao: DFT calculations.

Conflicts of interest

The authors declare no competing financial interest.

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