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Theoretical research on excited-state intramolecular proton coupled charge transfer modulated by molecular structure

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At the TD-B3LYP/TZVP/IEFPCM theory level, we have theoretically studied the excited-state intramolecular proton coupled charge transfer (ESIPCCT) process for both 4'-N,N-diethylamino-3-hydroxyflavone (3HFN) and 2-([2-(2-hydroxyphenyl)benzo[d]oxazol-6-yl)methylene]malononitrile (diCN-HBO) molecules. Our calculated hydrogen bond lengths and angles sufficiently confirm that the intramolecular hydrogen bonds $O_1-H_1\cdots O_2$ and $O_1-H_1\cdots N_1$ formed at the S_0 states of 3HFN and diCN-HBO should be significantly strengthened in the S_1 state, which is further supported by the results obtained based on the analyses of infrared spectra shifts, molecular orbitals and charge density differences maps. The significant strengthening of intramolecular hydrogen bonds $O_1-H_1\cdots O_2$ and $O_1-H_1\cdots N_1$ upon photoexcitation should facilitate the ESIPCCT process of the two title molecules. The scanned potential energy curves and confirmed excited-state transition states for both 3HFN and diCN-HBO show that the proton can be easily transferred from O_1 to O_2 (N_1 for diCN-HBO) through the strengthened intramolecular hydrogen bonds upon photoexcitation to the S_1 state.

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1. Introduction

Proton transfer (PT), as one of the most fundamental processes involved in chemical and biochemical reactions, deserves the attention of vast numbers of scientists and leads to thousands of publications and numerous books on related topics.^{1–24} From the molecular point of view, proton transfer can be generally categorized into intermolecular^{9–17} and intramolecular^{18–24} ones. The former, which are frequently encountered in biological systems, involve transfer of protons from the donor molecule to the acceptor molecule and commonly require uniquely arranged proton relay systems. While the latter cases involve proton transfer within the same molecular framework. Systems invoking excited state intramolecular proton transfer (ESIPT) are the focus of this research work.

ESIPT and the subsequent reverse proton transfer in the ground state achieve a fully reversible reaction cycle. Such cycles have been ubiquitously observed in aromatic molecules possessing proton donor and acceptor groups bridged with an intramolecular hydrogen bond. Hydroxyl or amino groups generally serve as the proton donors, and carbonyl oxygen or azo nitrogen serve as the proton acceptors. Upon electronic

excitation, the redistribution of electronic charge makes the proton donor more acidic and the acceptor more basic. Thermodynamically, the enhancement in the basicity/acidity factor, which is several orders of magnitude, makes ESIPT energetically favorable and provides the strong driving force for its occurrence, which explains why the same reaction does not occur in the ground state.

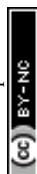
The redistribution of electron density does not only provide the driving force of ESIPT. The charge relocation can be reinforced by the integration of weak intermolecular interactions (such as solute/solvent dipole–dipole interactions), which dramatically influence both reaction kinetics and thermodynamics. Electronically excited to the Franck–Condon state, a molecule with an enhanced distribution of electronic charge and thus a large change of dipole moment strongly polarizes the surrounding medium. This leads to a dielectrically stabilized charge-transfer (CT*) excited state. The reaction leading to this state can be referred to as the excited state intramolecular charge transfer (ESICT) state. Such stabilization can also occur from the product side of ESIPT if the reaction generates a substantial charge transfer character. In a word, upon electronic excitation, the redistribution of electronic charge from electronic charge donor to acceptor obviously enhances the dipole moment of the molecule, which makes the proton donor more acidic and the acceptor more basic and finally facilitates the excited-state proton transfer.

The prerequisite of ESIPT lies in the formation of an intramolecular hydrogen bond. Recent advances have shown that

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hydrogen bonds play an important role in the excited-state behavior of molecular dyes. For instance, several theoretical approaches have explored the site-specific intermolecular excited state hydrogen bonding (H-bonding) property and found interesting correlations for fluorescence quenching, the dynamics of intermolecular charge transfer, and the spectral shifts *versus* the intermolecular H-bonding strength in the excited state.^{7,25–28} Thus, one would expect that the H-bonding properties, such as strength, distance and orientation, should affect ESIPT, ESICT, and consequently the ESIPT–ESICT coupled reaction.

The proton is a positively charged particle and its transfer leads to a new redistribution of electron density in the molecule in solution. When the ESIPT reaction is observed in condensed dielectric media, the dielectric polarization of the environment can have a substantial impact on photophysical events, stabilizing the charge-separated states. Therefore, the ESIPT process can be coupled with ESICT reaction both kinetically and energetically.^{29–31} This coupling depends strongly on the molecular structure. There are systems in which the interplay between ESICT and ESIPT is apparent, rendering drastically different photophysical phenomena such as dual emissions simultaneously associated with both ESIPT and ESICT. For example, as in the case of **3HFN**^{22,24} (shown in Scheme 1), when ESICT takes place upon Franck–Condon excitation, *i.e.*, N–N* (N: normal ground populated state, N*: electronically excited state), N* may thus have large charge separation. The subsequent N*–T* ESIPT (T* stands for excited tautomer), leading to strong redistribution of the electronic density, may be subject to drastic change of the dipole moment. For the other possibility, as in the case of **diCN-HBO**^{23,24} (shown in Scheme 1), the ESICT state is not formed upon excitation, whereas ESIPT is accompanied by significant charge separation. Thus, the (T*) form attains the property of the ESICT state and then relaxes to the (N) state *via* the (T) state, closing a reaction cycle.

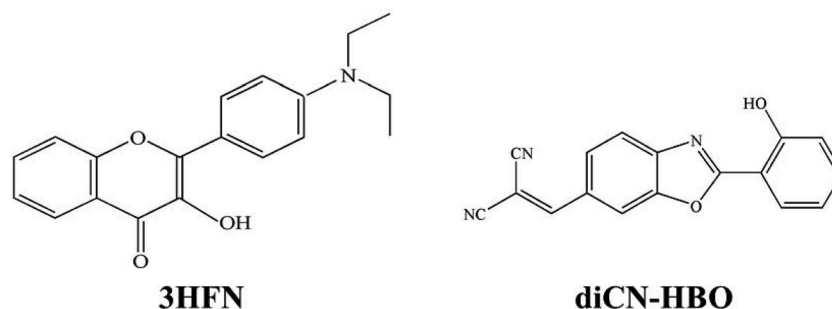
The **3HFN** molecule is strategically designed so that it undergoes ESICT prior to the ESIPT reaction.^{22,24} As depicted in Scheme 1, the electron donor (D_e, the dialkylamino group) and proton donor (D_p, O₁–H₁) occupy different positions but the same group serves as both proton acceptor and electron acceptor (A_{e,p}, O₂). Upon photoexcitation, ultrafast ESICT takes place, generating the CT* state on time scale (a few hundred femtoseconds). Subsequently, the solvent relaxation process from CT* to CT*_{eq} (subscript eq: equilibrium) and ESIPT occur

competitively. After reaching the solvent equilibrium, proton transfer reaction from CT*_{eq} to PCT* takes place. The **diCN-HBO** molecule is strategically designed so that it undergoes ESIPT prior to the ESICT reaction.^{23,24} As depicted in Scheme 1, the proton acceptor (A_p, N₁) and electron acceptor (A_e, the di-cyano group) are at different positions but the proton donor and electron donor are the same (D_{e,p}, O₁–H₁). The D_{e,p} group of **diCN-HBO** molecule is not a strong electron donating group in the excited state until it forms an anion *via* transferring a proton (H⁺) to the proton acceptor A_p. Once the proton is transferred, the resulting oxide anion –O₁[–] becomes a powerful electron donating group. Since D_{e,p} is π -conjugated with the electron acceptor A_e, a strong coupling matrix between D_e[–] and A_e is then expected, which should execute charge transfer toward the electron accepting cyano-group *via* π -conjugation. However, the overall perspective mechanism of the ESIPCCT process for the two molecules are not fully understood, especially in the computational way. Therefore, in this work, we are motivated to elucidate the overall perspective mechanism of the ESIPCCT process for the two molecules **3HFN** and **diCN-HBO** in a computational way by optimizing geometrical structures, analyzing molecular orbitals and charge density difference maps as well as the infrared vibrational spectra, especially the potential energy curves.

In this work, we aim to present a detailed and overall perspective of the ESIPCCT process for both **3HFN** and **diCN-HBO** molecules at the TD-B3LYP/TZVP/IEFPCM theory level since time dependent density functional theory (TDDFT) method has been demonstrated to be a reliable tool to investigate the excited state dynamics^{7,25–28,32–36} and provide a clear mechanism for previous experiment. The work is organized as follows: Section 2 introduces the computational methods adopted in this work; Section 3 presents the results and discussions, including functional test, optimized geometrical structures, molecular orbitals and charge density difference maps, infrared vibrational spectra analyses as well as potential energy curves and transfer mechanism; Section 4 gives the conclusion of this research work.

2. Computational methods

Using density functional theory (DFT) and time dependent density functional theory (TDDFT) methods with Becke's three-parameter hybrid exchange function with the Lee–Yang–Parr



Scheme 1 Geometrical structures of **3HFN** and **diCN-HBO**.



gradient-corrected correlation functional (B3LYP)^{37–42} as well as the TZVP basis set, we have theoretically studied the excited-state overall perspective of proton coupled charge transfer process modulated by molecular structure within **3HFN** and **diCN-HBO** systems by Gaussian 09 program.⁴³ Since previous experimental works were carried out in dichloromethane and chloroform solvents for **3HFN** and **diCN-HBO** respectively,^{22–24} we took these two solvents into consideration in all calculations *via* Polarizable Continuum Model (PCM) using the integral equation formalism variant (IEF-PCM)^{44–46} to be consistent with former experiments.^{22–24} For the ground-state part, all the corresponding structures (see Fig. 1 and 2) have been optimized without constraint using DFT methodology with vibrational frequencies analyses to ensure that all these configurations correspond to the local minima on the S_0 potential energy

surface (PES). The calculations about vertical excitation process were carried out from the S_0 -state optimized structures using TDDFT method with six low-lying absorbing transitions.

Furthermore, all the S_1 -state calculations were also based on the S_0 -state optimized structures with vibrational frequencies analyses. Even though the process for scanning potential energy curves of proton transfer is very time-consuming, especially for the excited state, we have theoretically scanned the S_0 - and S_1 -state potential energy curves to provide the corresponding dynamical overall perspectives of proton coupled charge transfer qualitatively and quantitatively. All the stationary points along the proton transfer coordinate were scanned by constraining optimizations and frequency analyses (no imaginary frequency) to obtain the thermodynamic corrections in the corresponding electronic state. The excited-state transition

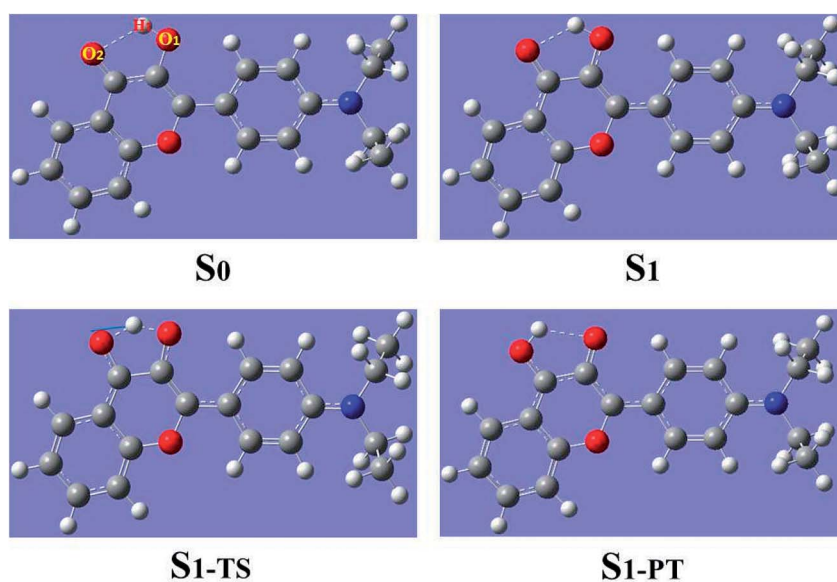


Fig. 1 Optimized geometrical structures of **3HFN** in dichloromethane at ground state S_0 , first singlet excited state S_1 , transition state S_1 -TS and after proton transfer S_1 -PT at (TD)B3LYP/TZVP/IEFPCM theory level.

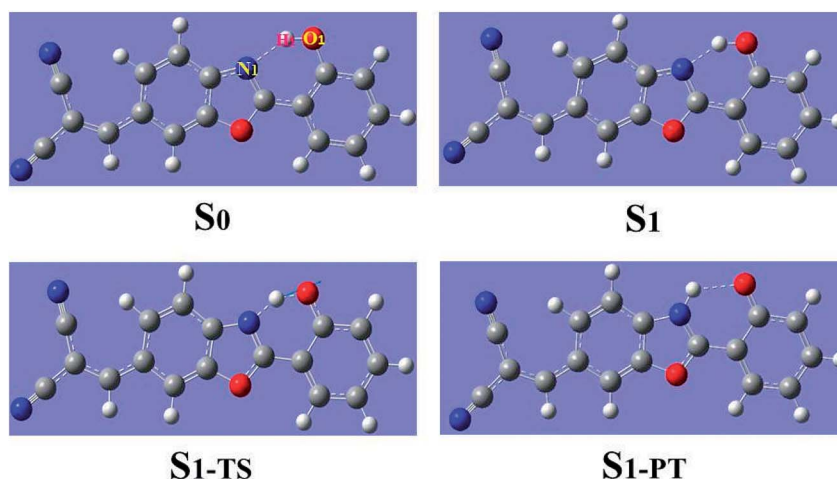


Fig. 2 Optimized geometrical structures of **diCN-HBO** in chloroform at ground state S_0 , first singlet excited state S_1 , transition state S_1 -TS and after proton transfer S_1 -PT at (TD)B3LYP/TZVP/IEFPCM theory level.



state was confirmed based on the stationary point with the highest potential energy in the excited-state potential energy curve. The transition state should have only one imaginary frequency whose vibrational mode points to the proton transfer product. The optimization and frequency analysis of the transition state are carried out with the Gaussian 16 program.⁴⁷

3. Results and discussions

3.1 Functional test

Based on experimental absorption spectra of **3HFN** in dichloromethane and **diCN-HBO** in chloroform, we have carried out the functional test with TZVP basis set on the TD/TZVP/IEFPCM theory level and the calculated results are listed in Table 1. From Table 1 we can find that the Becke's three-parameter hybrid exchange function with the Lee–Yang–Parr gradient-corrected correlation functional B3LYP should be more suitable for describing the photoexcitation of both **3HFN** in dichloromethane and **diCN-HBO** in chloroform. Therefore, all the calculations of the paper are at the (TD)B3LYP/TZVP/IEFPCM theory level.

3.2 Optimized geometrical structures

The geometrical structures of both **3HFN** in dichloromethane and **diCN-HBO** in chloroform have been optimized at the (TD) B3LYP/TZVP/IEFPCM theory level. For convenience, we named the intramolecular hydrogen bonds within **3HFN** and **diCN-HBO** as $O_1-H_1\cdots O_2$ and $O_1-H_1\cdots N_1$ respectively. The optimized ground-state (S_0) and first singlet excited-state (S_1) geometrical structures of **3HFN** in dichloromethane and **diCN-HBO** in chloroform are provided in Fig. 1 and 2 respectively and the important hydrogen bond parameters are listed in Table 2.

Table 1 Electronic excitation energy (nm), corresponding oscillator strengths (in parenthesis) and the corresponding compositions of the S_0-S_1 electronic transition for the **3HFN** (in dichloromethane) and **diCN-HBO** (in chloroform) chemosensor based on the TZVP basis set and IEFPCM combined with different functionals

	B3LYP	CAM-B3LYP	WB97XD	Exp.
3HFN	428 (0.806)	366 (1.05)	358 (1.07)	410
	H $\rightarrow$ L (99.1%)	H $\rightarrow$ L (91.1%)	H $\rightarrow$ L (87.1%)	
diCN-HBO	417 (0.777)	354 (1.48)	348 (1.52)	380
	H $\rightarrow$ L (98.7%)	H $\rightarrow$ L (82.8%)	H $\rightarrow$ L (78.9%)	

Table 2 The calculated primary bond lengths (Å) and angles (°) of **3HFN** (in dichloromethane) and **diCN-HBO** (in chloroform) in the S_0 , S_1 -TS and S_1 -PT states at (TD)B3LYP/TZVP/IEFPCM theory level

	3HFN			diCN-HBO		
	O_1-H_1	$H_1\cdots O_2$	δ ($O_1-H_1\cdots O_2$)	O_1-H_1	$H_1\cdots N_1$	δ ($O_1-H_1\cdots N_1$)
S_0	0.98	2.00	119.19	0.99	1.80	145.22
S_1	0.99	1.85	124.71	1.02	1.65	147.85
S_1 -TS	1.24	1.25	139.42	1.18	1.32	150.91
S_1 -PT	1.97	0.98	119.42	1.94	1.02	124.97

From Table 2, it can be found that the bond lengths of intramolecular hydrogen bonds $O_1-H_1\cdots O_2$ in **3HFN** and $O_1-H_1\cdots N_1$ in **diCN-HBO** are both decreased from ground state S_0 to first singlet excited state S_1 . At the same time, the bond angles of the two intramolecular hydrogen bonds are both increased from ground state S_0 to first singlet excited state S_1 .

For **3HFN** in dichloromethane, the hydrogen bond length is decreased from 2.00 Å in ground state S_0 to 1.85 Å in first singlet excited state S_1 and the hydrogen bond angle is increased from 119.19° in ground state S_0 to 124.71° in first singlet excited state S_1 . For **diCN-HBO** in chloroform, the hydrogen bond length is decreased from 1.80 Å in ground state S_0 to 1.65 Å in first singlet excited state S_1 and the hydrogen bond angle is increased from 145.22° in ground state S_0 to 147.85° in first singlet excited state S_1 . Both the decreases of the hydrogen bond lengths and the increases of the hydrogen bond angles indicate that the intramolecular hydrogen bonds formed in the ground state S_0 of both **3HFN** and **diCN-HBO** systems should be significantly strengthened upon photoexcitation to the first singlet excited state S_1 , which should result in the excited-state proton transfer from atom O_1 to atom O_2 (N_1) for **3HFN** (**diCN-HBO**).

3.3 Molecular orbitals and charge density difference maps

To visually illustrate the changes of charge distribution upon photoexcitation, the molecular orbitals responsible for the electronic transition of S_0-S_1 of **3HFN** and **diCN-HBO**, namely the lowest unoccupied molecular orbital (LUMO) and the highest occupied molecular orbital (HOMO) as shown in Table 1, are provided in Fig. 3 and 4 respectively. From Fig. 3, it can be found that the charge density localized on atom O_1 in HOMO of **3HFN** is totally disappeared in LUMO whereas that localized on atom O_2 of **3HFN** is greatly increased from HOMO to LUMO. The charge density difference map (CCDM) of **3HFN** shows the same results as that of the molecular orbitals from HOMO to LUMO. The results obtained from molecular orbitals and charge density difference map analyses indicate that due to the intramolecular charge transfer from O_1 and the dialkylamino group to O_2 , the intramolecular hydrogen bond $O_1-H_1\cdots O_2$ formed in the ground state S_0 of **3HFN** should be significantly strengthened upon photoexcitation to the first singlet excited state S_1 , which should result in the excited-state intramolecular proton transfer from atom O_1 to atom O_2 of **3HFN**.

For **diCN-HBO**, we can find from Fig. 4 that the charge densities formerly localized on both atoms O_1 and N_1 are almost completely disappeared from HOMO to LUMO. And the charge density difference map (CCDM) of **diCN-HBO** shows the same results. The charge density difference map of S_1 -PT-state **diCN-HBO** (CCDM*) shows that after excited-state proton transfer of H_1 from O_1 to N_1 , the charge density formerly localized on O_1 is greatly decreased and intramolecular charge transfer takes place from O_1 to the di-cyano group. That is to say, the excited-state intramolecular proton transfer of H_1 from O_1 to N_1 in **diCN-HBO** results in the intramolecular charge transfer from O_1 to the di-cyano group.



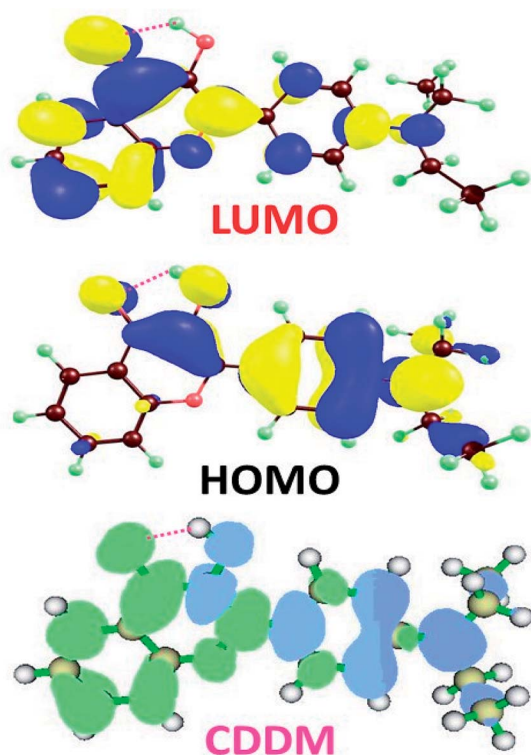


Fig. 3 Lowest unoccupied molecular orbital (LUMO), highest occupied molecular orbital (HOMO) and charge density difference map (CDDM) of **3HFN** in dichloromethane at TD-B3LYP/TZVP/IEFPCM theory level. The green and blue regions in CDDM represent increase and decrease of charge density upon photoexcitation respectively.

3.4 Infrared vibrational spectra analyses

We all know that detecting the infrared (IR) vibrational spectral shift is an effective way to explore the changes of hydrogen bonds upon photoexcitation and it has been demonstrated that the changes of hydrogen bonding can be monitored by the

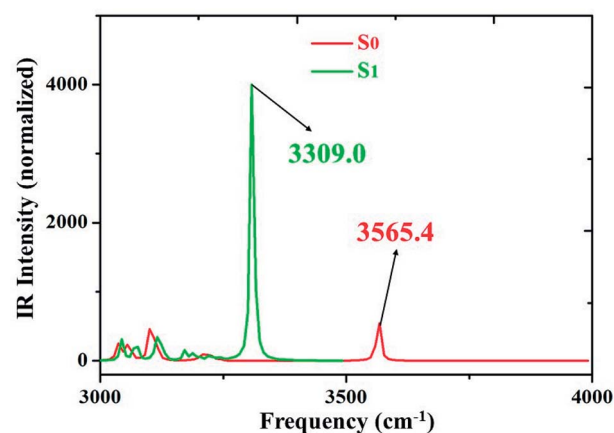


Fig. 5 IR spectra of **3HFN** in dichloromethane at the spectral region of O_1-H_1 stretching band in both S_0 and S_1 states at (TD)B3LYP/TZVP/IEFPCM theory level.

infrared spectra of some characteristic vibrational modes involved in the hydrogen bond formation.^{7,25–27} Therefore, the vibrational spectra involved in intramolecular hydrogen bonding moiety of **3HFN** and **diCN-HBO** in the conjunct vibrational O–H stretching modes have been displayed in Fig. 5 and 6, respectively. From Fig. 5, it should be noticed that our calculated O_1-H_1 stretching vibrational frequency of **3HFN** is redshifted from 3565.4 cm^{-1} in S_0 state to 3309.0 cm^{-1} in S_1 state. That is to say, the 256.4 cm^{-1} redshift is arisen from the photoexcitation. Similarly, the O_1-H_1 stretching vibrational frequency of **diCN-HBO** is redshifted from 3350.4 cm^{-1} in S_0 state to 2733.4 cm^{-1} in S_1 state and the 617 cm^{-1} redshift is also arisen from the photoexcitation. Han *et al.* have concluded that redshift of the infrared vibrational spectra of O–H upon photoexcitation should be resulted from the strengthening of hydrogen bond with O–H as hydrogen donor.^{7,25–27} Thus, we once again confirmed that the intramolecular hydrogen bonds formed in the ground state S_0 of both **3HFN** and **diCN-HBO**

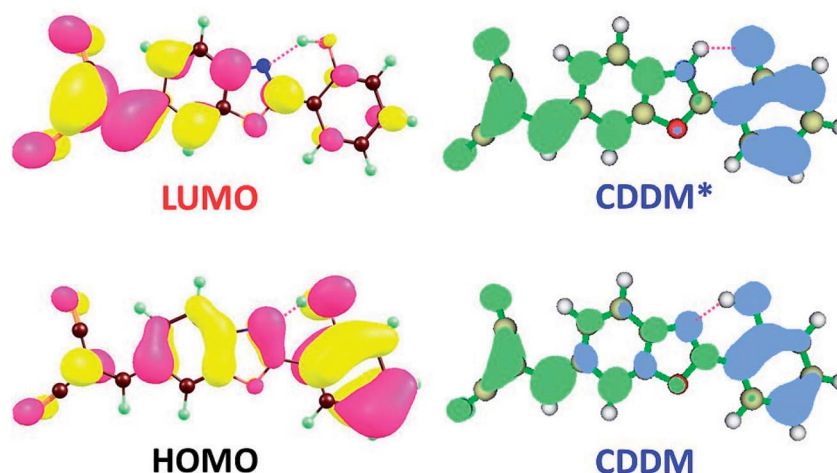
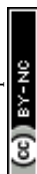


Fig. 4 Lowest unoccupied molecular orbital (LUMO), highest occupied molecular orbital (HOMO) and charge density difference map (CDDM) of **diCN-HBO** in chloroform at TD-B3LYP/TZVP/IEFPCM theory level. CDDM* is for **diCN-HBO** after proton transfer. The green and blue regions in CDDM represent increase and decrease of charge density upon photoexcitation respectively.



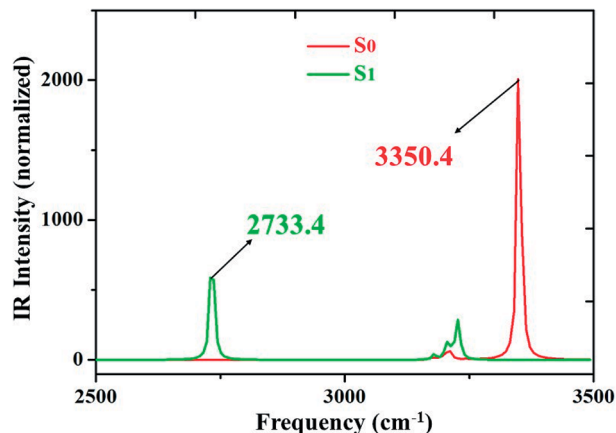


Fig. 6 IR spectra of diCN-HBO in chloroform at the spectral region of O₁–H₁ stretching band in both S₀ and S₁ states at (TD)B3LYP/TZVP/IEFPCM theory level.

systems should be significantly strengthened upon photoexcitation to the first singlet excited state S₁, which are consistent with the results obtained by both geometrical structure analyses and molecular orbital as well as charge density difference analyses.

3.5 Potential energy curves and transfer mechanism

To further reveal the detailed ESIPCCT mechanism, we have constructed the potential energy curves in both S₀ and S₁ states with fixing H₁–O₂ of **3HFN** (H₁–N₁ for **diCN-HBO**) bond lengths at a series of certain values in step of -0.05 Å. For **3HFN** system, the H₁–O₂ bond lengths have been fixed from 2.002 Å to 1.002 Å in step of -0.05 Å to scan the ground state potential energy curve, as shown in Fig. 7(a). From Fig. 7(a), we can find that the potential energy is gradually increased as the distance between atoms H₁ and O₂ decreases from 2.002 Å to 1.152 Å, with a high energy barrier 12.93 kcal mol⁻¹. Therefore, the proton transfer cannot happen in the ground state of **3HFN** due to the rather high potential energy barrier. Furthermore, to scan the S₁-state potential energy curve, the H₁–O₂ bond lengths have been fixed from 1.851 Å to 0.951 Å in step of -0.05 Å. Quite different from the situation in ground state, there only exist a rather small potential barrier (around 5.05 kcal mol⁻¹) separating the S₁-state **3HFN** and the S₁-PT-state **3HFN** (shown in Fig. 1, S₁-PT) configurations, as shown in Fig. 7(b). Furthermore, the transition-state geometrical structure (shown in Fig. 1, S₁-TS) corresponding to the point with the highest potential energy in Fig. 7(b) has been confirmed to only have one imaginary frequency that points to the direction of proton transfer. More important, the energy of the S₁-PT-state **3HFN** configuration is lower than that of the S₁-state **3HFN** by 2.15 kcal mol⁻¹. Thus, the intramolecular proton transfer from atom O₁ to atom O₂ of **3HFN** can be easily realized in the first singlet excited state. After the formation of S₁-PT-state **3HFN**, the radiation fluorescent process occurs with emission peak 585.59 nm (experimental value located at about 570 nm (ref. 22)) back to ground state, which is redshifted by more than 100 nm from that

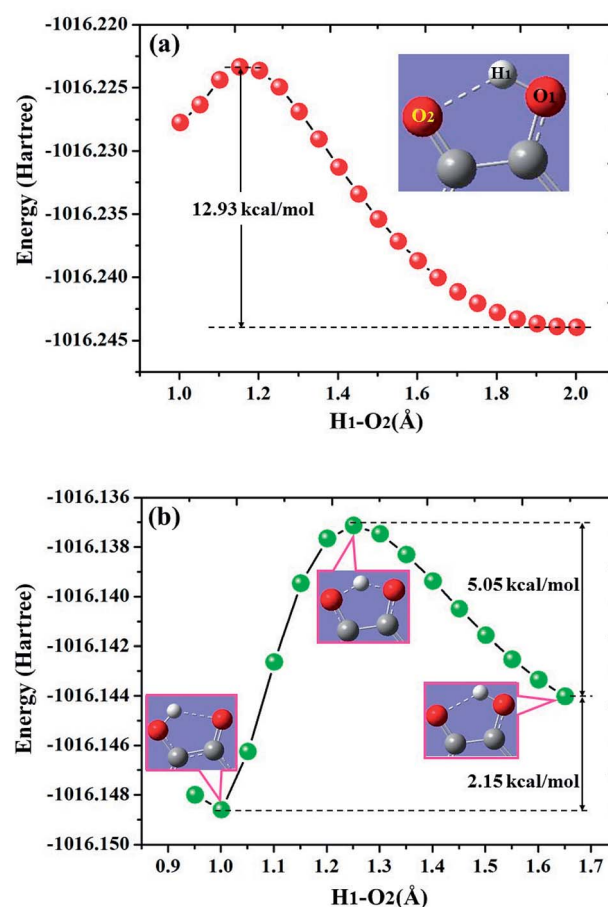


Fig. 7 Calculated potential energy curves of both S₀ (a) and S₁ (b) states for **3HFN** in dichloromethane along with the H₁–O₂ bond distance at (TD)B3LYP/TZVP/IEFPCM theory level.

(482.34 nm) of the S₁-state **3HFN** (experimental value located at about 495 nm (ref. 22)).

For **diCN-HBO** system, the H₁–N₁ bond lengths have been fixed from 1.797 Å to 0.947 Å in step of -0.05 Å to scan the ground state potential energy curve, as shown in Fig. 8(a). From Fig. 8(a), we can find that the potential energy is gradually increased as the distance between atoms H₁ and N₁ decreases from 1.797 Å to 1.097 Å, with a high energy barrier 11.46 kcal mol⁻¹. Therefore, the proton transfer cannot happen in the ground state of **diCN-HBO** due to the rather high potential energy barrier. Furthermore, to scan the S₁-state potential energy curve, the H₁–N₁ bond lengths have been fixed from 1.653 Å to 0.903 Å in step of -0.05 Å. Quite different from the situation in ground state, there only exist a rather small potential barrier (around 1.26 kcal mol⁻¹) separating the S₁-state **diCN-HBO** and the S₁-PT-state **diCN-HBO** (shown in Fig. 2, S₁-PT) configurations, as shown in Fig. 8(b). Furthermore, the transition-state geometrical structure (shown in Fig. 2, S₁-TS) corresponding to the point with the highest potential energy in Fig. 8(b) has been confirmed to only have one imaginary frequency that points to the direction of proton transfer. More important, the energy of the S₁-PT-state **diCN-HBO** configuration is rather lower than that of the S₁-state **diCN-HBO**



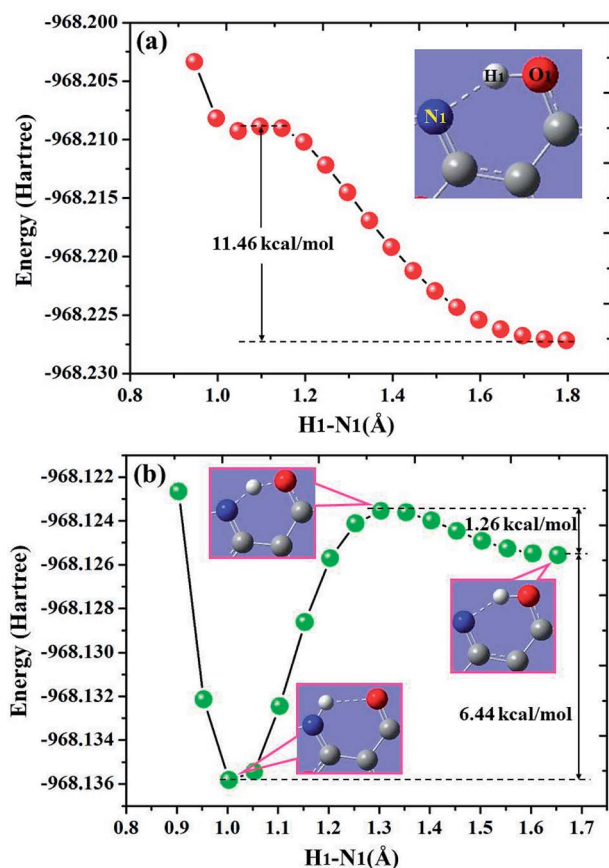


Fig. 8 Calculated potential energy curves of both S_0 (a) and S_1 (b) states for diCN-HBO in chloroform along with the H_1-N_1 bond distance at (TD)B3LYP/TZVP/IEFPCM theory level.

by $6.44 \text{ kcal mol}^{-1}$. Thus, the intramolecular proton transfer from atom O_1 to atom N_1 of diCN-HBO can be easily realized in the first singlet excited state. After the formation of S_1 -PT-state diCN-HBO, the radiation fluorescent process occurs with emission peak 729.85 nm (experimental value located at about 650 nm (ref. 23)) back to ground state, which is significantly redshifted by about 250 nm from that (480.7 nm) of the S_1 -state diCN-HBO (experimental value located at about 435 nm (ref. 23)).

4. Conclusion

In summary, we have theoretically studied the ESIPCCT process for both 3HFN and diCN-HBO molecules. By carrying out the functional test and comparing with previous experimental data, we confirm that the theoretical level (TDDFT/B3LYP/TZVP/IEFPCM) is more suitable for studying these two molecules. Our calculated results sufficiently confirm that intramolecular hydrogen bonds $O_1-H_1\cdots O_2$ and $O_1-H_1\cdots N_1$ are formed in the S_0 state of 3HFN and diCN-HBO molecules. Analyses of hydrogen bond lengths and bond angles indicate that the intramolecular hydrogen bonds formed within 3HFN and diCN-HBO molecules in ground state should be strengthened in the S_1 state, which is consistent with the results obtained based on

infrared spectra shifts, molecular orbitals and charge density differences maps. The significant strengthening of intramolecular hydrogen bonds $O_1-H_1\cdots O_2$ and $O_1-H_1\cdots N_1$ upon photoexcitation should facilitate the ESIPCCT process of the two title molecules. To obtain the detailed excited-state proton transfer mechanism, we have scanned the potential energy curves at S_0 and S_1 states for both 3HFN and diCN-HBO, which show the overall perspective of the ESIPCCT dynamics and provides a clear mechanism for previous experiment.

Conflicts of interest

There are no conflicts to declare.

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