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Theoretical insights into aggregation-induced emission of bis(cyanostyryl)pyrrole derivatives

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The molecular architecture and the positioning of the cyano group in cyanostilbene derivatives give rise to intriguing variations in their photophysical properties. The present study provides theoretical insights into the contrasting photoluminescence behaviors of bis(cyanostyryl)pyrrole derivatives with different cyano group positions. Using quantum mechanics/molecular mechanics (QM/MM) free energy perturbation methods, we investigated **o-DCSP** and **i-DCSP** isomers, which exhibited markedly different fluorescence quantum yields in the solution ($\Phi_f = 0.0036$ vs. 0.43) and aggregated states ($\Phi_f = 0.15$ vs. 0.12). We identified the minimum energy conical intersection (MECI) structures for both isomers, characterized by substantial rotation and pyramidalization of one ethylenic C=C bond, and determined the minimum energy path (MEP) connecting the Franck–Condon point to the MECI using the string method. By calculating the free energy profiles along this MEP, we revealed significant differences in energy barriers: **o-DCSP** showed a low barrier in solution (0.57 eV), which dramatically increased upon aggregation (2.36 eV), explaining its aggregation-induced emission behavior, whereas **i-DCSP** maintains relatively high barriers in both states (1.40 eV and 1.67 eV), resulting in efficient emission regardless of the environment. These findings establish a quantitative molecular-level understanding of the structure–property relationships in fluorescent materials and provide design principles for developing high-performance luminescent compounds with tailored emission characteristics for specific applications.

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Introduction

The exploration of organic π -conjugated materials has garnered substantial attention in recent years, owing to their ability to fine-tune their optical and electronic properties.^{1–3} This versatility has led to significant advancements in their utilization in a wide range of optoelectronic devices including organic light emitting diodes (OLEDs), organic solar cells, and chemical sensors.^{4,5} The use of these materials in various optoelectronic applications depends on their photophysical parameters, particularly fluorescence quantum yields.⁶ Fluorophores with high quantum yields emit strong fluorescence even at low concentrations and are suitable for lasers and OLEDs. Dyes with high quantum yields are also useful for the staining, labeling, and imaging of cells.

A few organic luminescent materials exhibit high quantum yields in both solid and liquid phases.^{7,8} Typically, these

materials suffer from low quantum yields in the condensed phase due to co-facial π – π stacking interactions, which lead to aggregation-caused quenching (ACQ).⁹ Another photophysical phenomenon observed in luminescent molecules is aggregation-induced emission (AIE), which is characterized by weak emission in dilute solutions but highly emissive properties upon aggregation or in the solid phase.¹⁰

Cyanostilbene derivatives are promising materials for the production of AIE-active compounds and have received significant attention because of their high emissivity, particularly in the solid phase.¹¹ While these fluorophores generally display weak emission in solution,^{12,13} some reports have documented exceptions where certain cyanostilbene molecules exhibit highly emissive behavior in solution.^{14,15} This variability illustrates that the fluorescence properties of cyanostilbene-based compounds are highly dependent on their molecular architecture and substitution patterns.

In this context, Yokoyama and Nishiaki recently synthesized two structural isomers of bis(cyanostyryl)pyrrole derivatives, named **o-DCSP** and **i-DCSP**, which differ in the positioning of the cyano groups on the olefin units (Scheme 1). These compounds exhibit structure-dependent fluorescence emission, making them ideal systems for investigating the aggregation-induced emission (AIE) phenomenon.¹⁶ **o-DCSP** exhibited AIE behavior,

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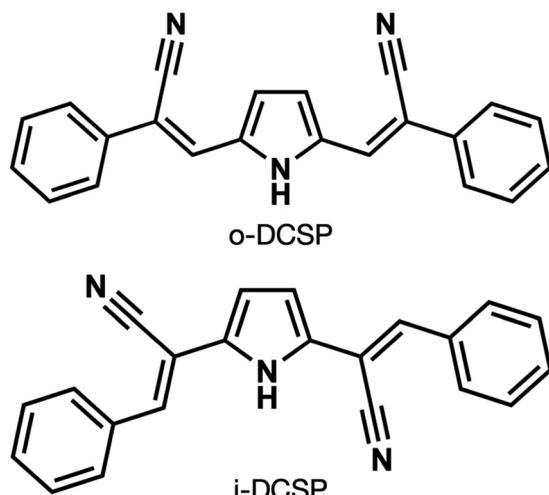
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Scheme 1 Chemical structures of bis(cyanostyryl)pyrrole derivatives **o-DCSP** and **i-DCSP**.

displaying minimal emission in solution ($\Phi_f = 0.0036$) but enhanced luminescence in the single-crystal state ($\Phi_f = 0.15$). In contrast, **i-DCSP** demonstrated different photophysical properties with a high fluorescence quantum yield in solution ($\Phi_f = 0.43$), which slightly diminished upon aggregation to the single-crystal state ($\Phi_f = 0.12$). This striking contrast in emission properties between positional isomers highlights the critical role of cyano group positioning in determining the photophysical behavior. The investigation of these structurally similar yet photophysically distinct molecules offers valuable insights into the molecular mechanisms underlying the AIE phenomenon and structure–function relationships in fluorescent materials.

In this study, we focused on elucidating the molecular mechanism underlying the distinct photoluminescence behaviors of **o-DCSP** and **i-DCSP** through quantum mechanics/molecular mechanics (QM/MM) modeling. We employed QM/MM free energy perturbation methods to quantitatively analyze the excited-state relaxation processes in both solution and aggregated states. By determining the free-energy profiles along the minimum energy path for the relaxation process after photoexcitation, we reveal how subtle differences in cyano group positioning lead to dramatically different energy barriers that control the radiative and non-radiative decay pathways. This comprehensive analysis provides fundamental insights into the structure–property relationships in AIE-active materials and establishes design principles for developing high-performance luminescent compounds with tailored emission characteristics for specific applications.

Computational details

Quantum chemical calculations offer valuable insights into molecular-level photoexcitation processes by analyzing ground and excited-state potential energy surfaces (PESs).^{17,18} Conical intersection (CI), the crossing point between PESs, is a theoretical tool for describing the photoexcitation process.^{19–21}

The lowest energy point within the CI hyperspace is referred to as the minimal energy conical intersection (MECI).^{22–24} Identifying the location of the MECI between the ground and first electronic excited states (S_0/S_1) can provide valuable information on the radiative and non-radiative decay processes.^{25,26} The fluorescence quantum yield of a molecule is primarily determined by the fraction of non-radiative transitions that occur near the S_0/S_1 -MECI.^{21,27}

In this study, we employed QM/MM models to investigate the effects of molecular environments on these photophysical processes. Specifically, we determined the minimum energy path (MEP) from the Franck–Condon (FC) point to the MECI using the string method^{28,29} and then calculated the free energy changes along this MEP using QM/MM free energy perturbation (FEP) methods.^{30,31} This approach allows us to quantitatively analyze how the surrounding molecules in both solution and aggregated states influence the energetic barriers between the excited and ground states, providing a molecular-level explanation for the observed differences in fluorescence quantum yields.

Geometry optimizations

Density functional theory (DFT) calculations were performed to determine the minimum energy structures in the S_0 ground state. The optimized structure in the S_0 state obtained here is denoted as S_0 -Min. The point at which S_0 -Min is vertically excited to reach the S_1 -state potential energy hypersurface is called the Franck–Condon (FC) point. The BHHLYP functional^{32,33} was selected for its effectiveness in handling electron correlations in spin-flip (SF) time-dependent (TD) DFT scenarios,³⁴ which is particularly crucial for analyzing conical intersections in molecular systems.³⁵ For all DFT calculations, 6-31G(d) basis functions were utilized.

The minimum energy point on the CI hypersurface (MECI) is the critical point below which nonadiabatic transitions *via* CIs cannot occur. The MECI between the S_0 and S_1 states of **o-DCSP** and **i-DCSP** was determined using a branching-plane updating technique,³⁶ where the electronic structures of the molecule were determined using the spin-flip approach³⁷ within the time-dependent DFT (SF-TD-DFT) method.³⁵ The initial coordinates for the MECI searches were the rotated and pyramided conformations of one of the two ethylenic C=C bonds in each compound. The Q-Chem 6.2 program package³⁸ (Q-Chem, Inc., Pleasanton, CA) was used for the DFT calculations.

Path optimizations

An MEP can be defined as the reaction coordinate at which the potential energy is minimized for changes in the molecular geometry corresponding to the orthogonal motions. The MEPs for the excited-state relaxation process to reach the CIs were determined for both **o-DCSP** and **i-DCSP** using DFT calculations.

The MEPs connecting the FC points and MECI of both **o-DCSP** and **i-DCSP** were determined using the string method.^{28,29} The basic idea of the string method is to find an MEP by evolving a sequence of images, called a string, in the direction of the force derived from the potential energy gradient. The potential energy



gradients for the S_1 states of the molecules were calculated using SF-TD-DFT. Path optimization using the string method was performed using a program developed in-house. An initial path was prepared by linearly interpolating the molecular geometries of the FC point and the MECI in internal coordinates with ten discrete points. The update step size in path optimization was set to 1.0 Bohr. The convergence of the path optimization was examined using the average value of the potential energy change when updating the path, ΔV , and the average value of the energy gradient in the direction perpendicular to the path in each image, $\nabla_{\perp} V$. The resultant MEPs for **o**-DCSP and **i**-DCSP were linearly interpolated with 60 discrete points in the internal coordinate space to improve the accuracy of the free energy calculations. The reaction coordinate parameter $\sigma \in [0,1]$ was used to parameterize the MEPs.

Free energy calculations

The free-energy profiles along the MEPs of **o**-DCSP and **i**-DCSP embedded in their aggregates were computed using the QM/MM-FEP method.^{30,31} In this study, a single molecule embedded in its aggregate formed a QM subsystem that changed its conformation along with the MEP, and the surrounding molecules formed an MM subsystem. The QM/MM-FEP method can be summarized as follows: the free energy difference between the two adjoining conformational states, A and B, in a QM/MM system can be defined as

$$\Delta F_{\text{qm/mm}}^{(A \rightarrow B)} = \Delta E_{\text{qm}}^{(A \rightarrow B)} + \Delta F_{\text{int}}^{(A \rightarrow B)}. \quad (1)$$

The $\Delta E_{\text{qm}}^{(A \rightarrow B)}$ term is expressed as follows:

$$\Delta E_{\text{qm}}^{(A \rightarrow B)} = \langle E_{\text{qm}}(\mathbf{r}^{(B)}, \mathbf{R}^{(B)})_{\mathbf{R}^{(B)}} \rangle - \langle E_{\text{qm}}(\mathbf{r}^{(A)}, \mathbf{R}^{(A)})_{\mathbf{R}^{(A)}} \rangle, \quad (2)$$

where $\langle \dots \rangle_{\mathbf{R}^{(B)}}$ represents an ensemble average over the MM subsystem in the Bth state. Here, the perturbation corresponds to the forward or backward movement of the QM atoms when all MM atoms are fixed. Canonical MD simulations were performed while fixing QM atoms to obtain the required ensembles. The $\Delta F_{\text{int}}^{(A \rightarrow B)}$ term is related to an average of the function of the energy difference between the two states, as determined by sampling for the Ath state:

$$\Delta F_{\text{int}}^{(A \rightarrow B)} = -\frac{1}{\beta} \ln \left\langle \exp \left[-\beta \Delta E_{\text{int}}^{(A \rightarrow B)} \right] \right\rangle_{\mathbf{R}^{(A)}}, \quad (3)$$

where β is the reciprocal temperature, and

$$\Delta E_{\text{int}}^{(A \rightarrow B)} = E_{\text{int}}(\mathbf{r}^{(B)}, \mathbf{R}^{(A)}) - E_{\text{int}}(\mathbf{r}^{(A)}, \mathbf{R}^{(A)}) \quad (4)$$

The first term represents the perturbation corresponding to the conformational change of the QM atoms from $\mathbf{r}^{(A)}$ to $\mathbf{r}^{(B)}$ when all the MM atoms are fixed at $\mathbf{R}^{(A)}$. The E_{int} term consists of additive contributions from van der Waals and electrostatic interactions between QM and MM atoms:

$$E_{\text{int}}(\mathbf{r}^{(B)}, \mathbf{R}^{(A)}) = E_{\text{vdw}}(\mathbf{r}^{(B)}, \mathbf{R}^{(A)}) + E_{\text{es}}(\mathbf{r}^{(B)}, \mathbf{R}^{(A)}). \quad (5)$$

QM/MM-FEP calculations were performed using an in-house program.

MD simulations

The QM/MM-FEP calculation requires ensemble averages over the MM subsystem, while fixing all QM atoms. Canonical MD simulations were performed to obtain the necessary ensembles. All MD simulations were performed using the AMBER 20 program package³⁹ (University of California, San Francisco, CA). The computational models for the aggregated structures of the **o**-DCSP and **i**-DCSP were taken from the crystal structures reported in the previous study.¹⁶ For the crystal structures, we extended the unit cell to construct a supercell encompassing a region with a radius of 20 Å from the central molecule. The MD simulations of the aggregated state were performed for the systems containing 192 **o**-DCSP and 180 **i**-DCSP molecules in the $8 \times 3 \times 2$ and $3 \times 3 \times 5$ supercells with sizes of 105.4 and 85.4 nm³, respectively, under periodic boundary conditions. Following the energy minimization and equilibration processes, 2 ns of production runs were performed under an isothermal-isobaric (NpT) ensemble. A time step of 2.0 fs was used in all MD simulations. Room temperature (300 K) and a constant pressure of 1 bar were maintained using the weak-coupling algorithm.⁴⁰ All bonds involving hydrogen were constrained using the SHAKE algorithm.⁴¹ The electrostatic interactions were computed using the particle mesh Ewald algorithm⁴² with a cut-off radius of 8 Å. The coordinates of the QM subsystem were frozen during the MD simulations, and the MM subsystem was modeled using the general AMBER force field (GAFF).⁴³ For the QM/MM calculations and QM/MM-FEP analysis, 50 and 1000 structures were sampled, respectively, from the 2-ns MD trajectory at each discrete point along the MEP.

Results

Fig. 1 displays the optimized structures of **o**-DCSP and **i**-DCSP in their ground state (S_0 -Min) and at the S_0/S_1 -MECI point. Scheme 2 shows the conformational parameters of the ethylenic C=C group, highlighting the significant structural changes at the MECI point using **o**-DCSP as an example. Table 1 provides detailed conformational data for both molecules, with a particular emphasis on the ethylenic C=C group.

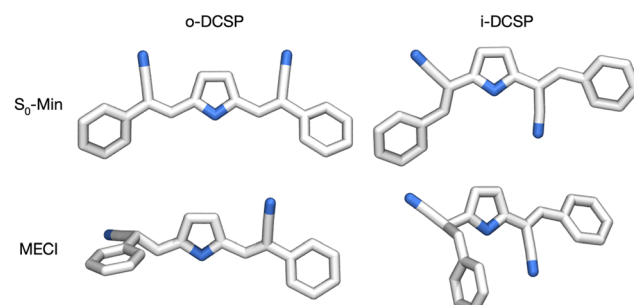
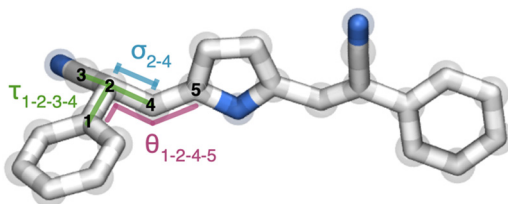


Fig. 1 Optimized local stable structures of **o**-DCSP and **i**-DCSP in the (a) ground state (S_0 -Min) and at the minimum-energy conical intersection points between the S_0 and S_1 states (MECI).





Scheme 2 Conformational parameters of the ethylenic $\text{C}=\text{C}$ moiety, as exemplified by **o-DCSP**.

Table 1 Summary of conformational details for the ethylenic $\text{C}=\text{C}$ moiety in **o-DCSP** and **i-DCSP**, comparing their ground state (S_0 -Min) structures and minimum-energy conical intersection points between S_0 and S_1 states (MECI)

Compound	σ_{2-4}		$\theta_{1-2-4-5}$		$\tau_{1-2-3-4}$	
	[\AA]	[degree]	[S_0 -Min]	[MECI]	[S_0 -Min]	[MECI]
o-DCSP	1.349	1.439	178.7	114.8	178.8	117.1
i-DCSP	1.349	1.436	178.9	113.9	178.8	113.0

The S_0/S_1 -MECI structure exhibited substantial variations from the S_0 -Min structure, as illustrated in Fig. 1 and Table 1. At the MECI point, one of the two ethylenic $\text{C}=\text{C}$ double-bond sites in each molecule rotates and forms a pyramidal shape. The ethylenic moiety parameters (including bond length, σ , and dihedral angles, θ and τ) displayed significant conformational changes from the S_0 -Min to MECI structure. These changes indicated substantial molecular distortions at the MECI point. This transition from the $S_0 \rightarrow S_1$ FC point (corresponding to the S_0 -Min state) to the MECI state is characteristic of cyanostilbene derivatives, consistent with previous studies.^{35,36}

Previous studies have demonstrated that conical intersections, which are regions of energetic overlap between the ground and excited states, play a critical and intricate role in the AIE mechanism.^{44–48} AIEgens, which are compounds that exhibit AIE, undergo spontaneous conformational changes in the excited state following photoexcitation, moving from a vertically excited FC point to CIs. At these CIs, non-radiative transitions occur from the excited state to the ground state, resulting in no emission. However, when AIEgens are in an aggregated state, the suppression of conformational changes along the relaxation pathway from the FC point to the CIs inhibits the non-radiative transition process, leading to emission.

Branching plane analysis is a helpful tool for characterizing conical intersections and is crucial for understanding photochemical reactions and non-radiative decay processes.^{49–53} This analysis, which involves calculating the energy difference gradient vector (**g**) and nonadiabatic coupling vector (**h**) in the MECI geometry, defines a plane, the branching plane. This plane is the locus of the most significant geometric distortion, which leads to non-radiative decay. In the context of AIE, branching plane analysis provides profound insights into the structural factors that govern the accessibility of conical

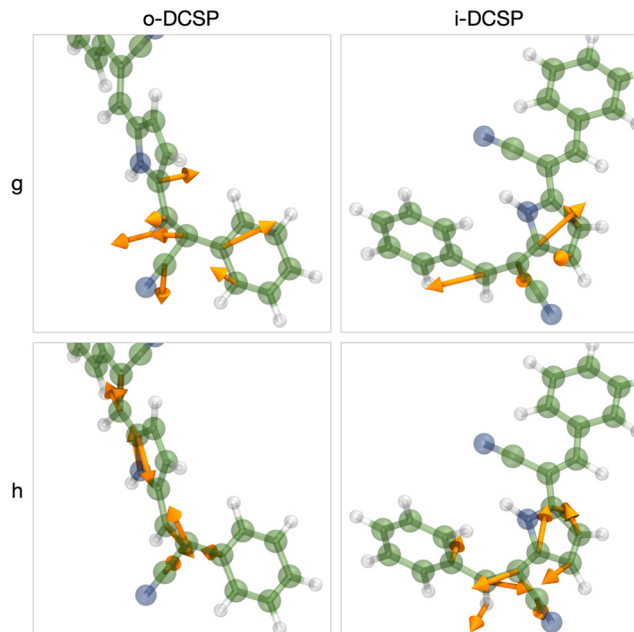


Fig. 2 Branching plane analysis of S_0/S_1 -MECIs for **o-DCSP** and **i-DCSP**. The energy difference gradient vector (**g**) and nonadiabatic coupling vector (**h**) are indicated by orange arrows.

intersections, and thus, the efficiency of non-radiative decay in the aggregated state.

Branching plane analysis, as shown in Fig. 2, revealed that the main components of both the **g** and **h** vectors for all the compounds were localized around one of the cyano groups. This concentration suggests that the relaxation pathways from S_0/S_1 -MECI primarily involve the rotational and pyramidalization motions of these groups. No significant differences in the vectors were found between **o-DCSP** and **i-DCSP**, implying that the AIE properties are influenced by other factors, such as the topology of the excited-state potential energy surface, and not solely by the MECI geometry. These findings suggest that intermolecular interactions and excited-state energy landscapes are crucial for determining the AIE behavior.

AIE is a significant photophysical phenomenon characterized by unique fluorescence properties in the aggregated molecular state. This complex phenomenon cannot be comprehensively understood through isolated gas-phase analysis alone, necessitating a detailed examination of molecular environmental effects. The two isomers, **o-DCSP** and **i-DCSP**, exhibited distinct fluorescence characteristics in THF solution *versus* aggregated states, suggesting differential relaxation processes following photoexcitation in each environment. To elucidate these differences at the molecular level, we constructed QM/MM models and conducted theoretical analyses, where we treated the central molecule as the QM region and the surrounding molecules as the MM region to examine the fluorescence properties in the aggregated state in detail.

Fig. 3 illustrates the three-dimensional structures of **o-DCSP** and **i-DCSP** in THF solution and their aggregated states. To quantitatively analyze the excited-state relaxation processes,



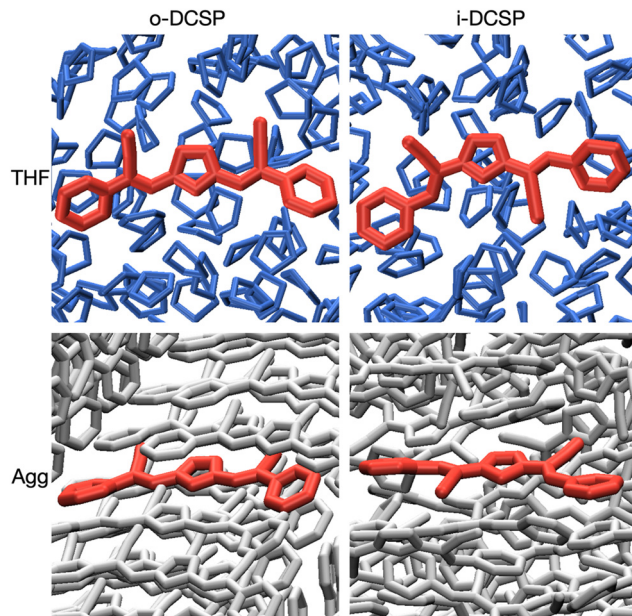


Fig. 3 Three-dimensional structures of **o**-DCSP and **i**-DCSP in THF solution and aggregated states.

we employed the QM/MM FEP method. This comprehensive analysis consisted of three sequential steps. First, the MEP from the FC point to the MECI was determined for **o**-DCSP and **i**-DCSP in the isolated gas phase using the string method.^{28,29} Second, NpT ensemble MD simulations were performed for 4 ns at 60 discrete points along the MEP between the FC and MECI points under thermodynamic conditions of constant temperature ($T = 300$ K) and pressure ($p = 1$ atm), where only the central QM molecule was fixed, while the surrounding MM molecules were allowed to move. Finally, from the latter 2 ns of the MD trajectories, we extracted 50 structures for QM/MM calculations and 1000 structures for QM/MM FEP analysis to determine free energy changes along the MEP in both the THF solution and molecular aggregated states.

Fig. 4 displays the free energy changes along the MEP from the FC to MECI points for both **o**-DCSP and **i**-DCSP in the THF solution. The graph shows the average potential energy values (E_{qm}) obtained from QM/MM calculations performed on 50 structures extracted from 2 ns NpT MD simulations at 60 discrete points along the MEP. The E_{qm} values reflect the influence of electrostatic interactions between the QM molecule and surrounding molecules, fluctuating with the thermal motion of the surrounding molecules. Additionally, we present values with free energy changes calculated using the QM/MM FEP method ($F_{qm/mm}$), which we define as the free energy profile along the MEP. For **o**-DCSP, the free-energy profile reveals that the MECI point is energetically more stable than the FC point. However, an energy barrier of approximately 0.57 eV (13 kcal mol⁻¹) exists along the pathway from the FC to the MECI points. Experimentally, **o**-DCSP exhibited a notably low fluorescence quantum yield in THF solution.¹⁶ This suggests that excess energy during photoexcitation readily overcomes this energy barrier, enabling an efficient radiationless transition through rapid access to CIs.

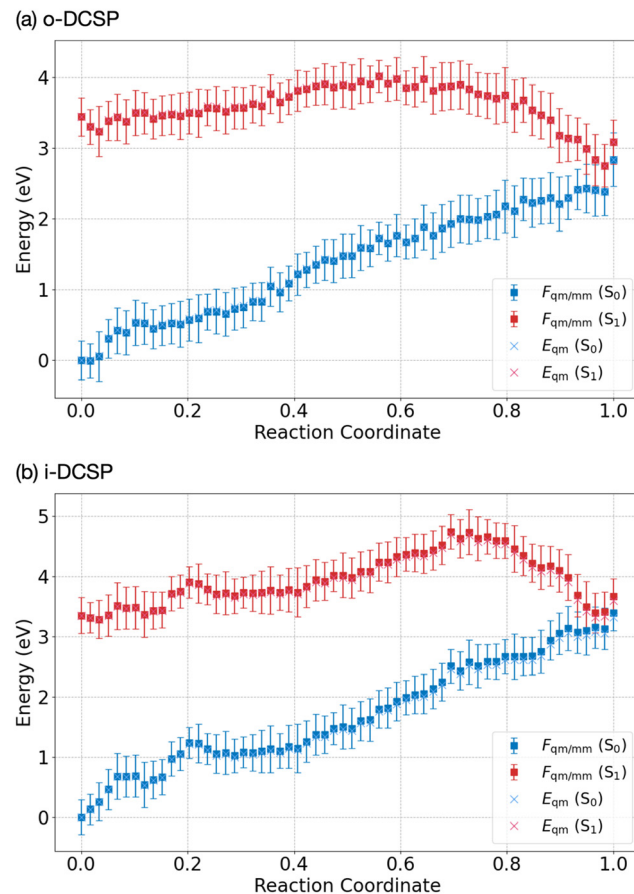


Fig. 4 Free energy profiles along the minimum energy path (MEP) from the Franck-Condon (FC) point to the minimum energy conical intersection (MECI) point for (a) **o**-DCSP and (b) **i**-DCSP in THF. The graphs show the average potential energy values (E_{qm}) from the QM/MM calculations, as well as the free energy profiles ($F_{qm/mm}$) calculated using the QM/MM FEP method.

In contrast, for **i**-DCSP, while the FC and MECI points are energetically comparable, a significant energy barrier of approximately 1.40 eV (32 kcal mol⁻¹) exists between them. Experimentally, **i**-DCSP demonstrated a relatively high fluorescence quantum yield even in a THF solution.¹⁶ This indicates that because of the substantial energy barrier, structural changes from the FC point to CIs after photoexcitation are energetically unfavorable, suppressing radiationless transitions and resulting in stronger emission compared to **o**-DCSP.

For both isomers, the contribution of the free energy changes obtained from FEP was relatively small, suggesting a limited impact of steric hindrance from the THF solvent molecules.

Fig. 5 illustrates the free energy changes along the MEP from the FC to the MECI points for **o**-DCSP and **i**-DCSP in the aggregated state. The free-energy profile of **o**-DCSP differed markedly from that of the THF solution. Specifically, the pathway from the FC to the MECI points becomes a steep uphill, with a maximum value of approximately 2.36 eV (54 kcal mol⁻¹) higher than the FC point. This reaction barrier is 1.79 eV (41 kcal mol⁻¹) greater than that in THF solution. Experimentally,



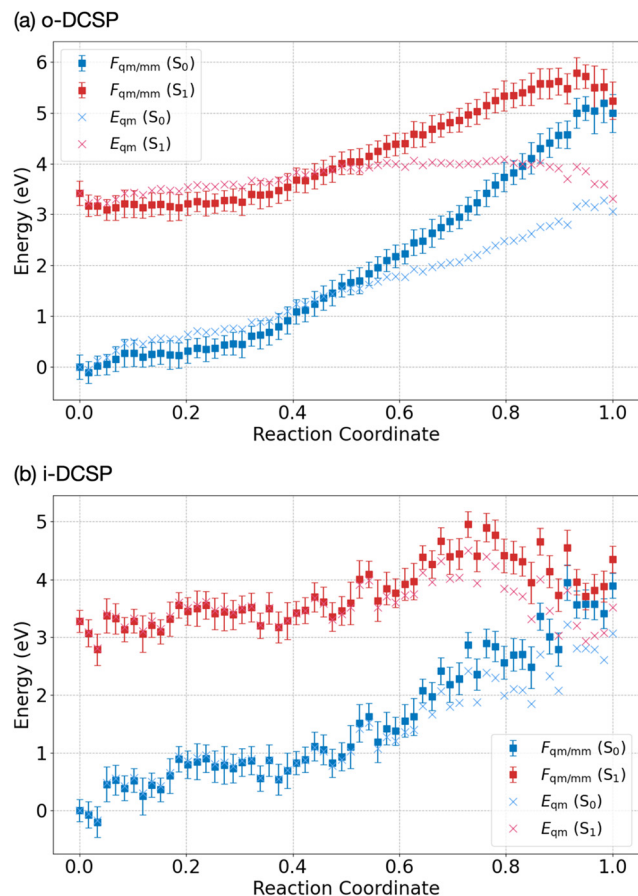


Fig. 5 Free energy profiles along the minimum energy path (MEP) from the Franck-Condon (FC) point to the minimum energy conical intersection (MECI) point for (a) **o**-DCSP and (b) **i**-DCSP in the aggregated state. The graphs show the average potential energy values (E_{qm}) from the QM/MM calculations, as well as the free energy profiles ($F_{qm/mm}$) calculated using the QM/MM FEP method.

the fluorescence quantum yield of **o**-DCSP in the aggregated state is significantly higher than that in a dilute solution.¹⁶ This suggests that in the aggregated state, steric hindrance from the surrounding molecules strongly inhibits structural changes from the FC point to CIs, making the radiationless transition pathway energetically highly unfavorable, thereby enhancing the emission intensity. This was proposed as the molecular mechanism by which **o**-DCSP induces AIE.

Conversely, the free energy profile for **i**-DCSP in the aggregated state resembles that in THF solution, with a reaction barrier of approximately 1.67 eV (39 kcal mol⁻¹) along the pathway from the FC to the MECI points. This barrier height is only 0.27 eV (6 kcal mol⁻¹) greater than that in THF solution. Experimentally, the fluorescence quantum yield of **i**-DCSP in the aggregated state was slightly lower than that in the dilute solution.¹⁶ These results suggest that for **i**-DCSP, similar to that in THF solution, radiationless transitions are suppressed owing to relatively high energy barriers, making the pathway from the FC point to CIs energetically unfavorable.

Compared to **o**-DCSP in the aggregated state, the free energy barrier for **i**-DCSP in the aggregated state is 0.69 eV

(16 kcal mol⁻¹) lower. Experimentally, the fluorescence quantum yield of **i**-DCSP in the aggregated state is lower than that of **o**-DCSP in the aggregated state.¹⁶ This indicates that steric hindrance in the aggregated state is less pronounced for **i**-DCSP than that for **o**-DCSP, resulting in weaker suppression of radiationless transitions. Consequently, some molecules undergo radiation-less transitions *via* the CI point, thereby reducing their emission intensity.

Discussion

The fluorescence quantum yield (Φ_f) depends on the balance between the radiative (k_r) and non-radiative (k_{nr}) decay rates, according to $\Phi_f = k_r/(k_r + k_{nr})$. Although our calculations do not directly yield k_r or k_{nr} values, the computed energy barriers offer insights into k_{nr} through the Arrhenius relationship ($k_{nr} \propto \exp(-\Delta E/kT)$). In solution, **i**-DCSP displays a higher quantum yield ($\Phi_f = 0.43$) than **o**-DCSP ($\Phi_f = 0.0036$),¹⁶ consistent with a larger computed barrier (1.40 eV vs. 0.57 eV) for the conformational change leading to non-radiative decay. This significant difference in barrier heights explains why **i**-DCSP maintains efficient emission in solution, whereas **o**-DCSP exhibits very weak fluorescence. The high energy barrier effectively suppresses the C=C torsion that governs the main non-radiative decay pathway in **i**-DCSP, despite both compounds having a similar potential for rotation in isolated molecules.

Our QM/MM-FEP analysis demonstrated that aggregation influences excited-state behavior by reshaping the overall energy landscape through specific intermolecular interactions. In **o**-DCSP, aggregation dramatically increases the barrier to the non-radiative pathway (from 0.57 eV in solution to 2.36 eV in the solid state), leading to a more pronounced solid-state emission. X-ray crystallographic analysis revealed that **o**-DCSP forms a hydrogen-bonding network between stacking columns with a π - π stacking distance of 3.395 Å,¹⁶ indicating strong intermolecular interactions that effectively restrict molecular motion. In contrast, **i**-DCSP experienced only a modest increase in barrier height (0.27 eV) upon aggregation, resulting in a smaller enhancement; indeed, its solid-state quantum yield decreased slightly compared to its solution value. The molecular packing of **i**-DCSP, characterized by dimeric structures with hydrogen bonding (2.230 Å) and weaker π - π interactions (a stacking distance of 3.611 Å),¹⁶ provides less restriction to the critical motion compared to **o**-DCSP. These observations reinforce the conclusion that subtle differences in molecular packing can significantly modulate excited-state processes rather than uniformly impeding all possible rotational motions.

While this study focused primarily on intramolecular photo-physics, intermolecular processes may also contribute to non-radiative decay in the condensed phase. Recent reports have indicated the possibility of dimeric conical intersections in closely stacked aromatic systems,⁵⁴ which could provide alternative non-radiative pathways. In particular, for **i**-DCSP, such intermolecular channels may partially offset the high intramolecular barrier, contributing to its reduced solid-state

quantum yield compared with that of the solution. This mechanism could explain why **i-DCSP**, despite its inherently high barrier to intramolecular conical intersection, shows a decrease in fluorescence efficiency upon aggregation. The present study provides a clear theoretical explanation of the impact of cyano group positioning on the photophysical properties in the solution state, along with the various factors that emerge in the aggregated state to modulate these properties.⁵⁵ Future investigations employing multi-chromophore QM regions within QM/MM frameworks could offer greater insight into these intermolecular mechanisms and their potential roles in fine-tuning the aggregation-induced emission phenomenon.

Conclusion

Through systematic QM/MM free energy perturbation analysis, we established a quantitative molecular-level understanding of the contrasting photoluminescence behaviors observed in **o-DCSP** and **i-DCSP** isomers. Our computational findings correlate with the experimental observations and provide mechanistic explanations for their divergent fluorescence properties.

The AIE phenomenon exhibited by **o-DCSP** can be directly attributed to the dramatic increase in the energy barrier from the FC to the MECI point when transitioning from the solution (0.57 eV) to the aggregated state (2.36 eV). This 1.79 eV increase results from steric hindrance imposed by the surrounding molecules in the aggregated state, effectively blocking the pathway to radiationless decay. As reported by Yokoyama *et al.*,¹⁶ this leads to a significant enhancement in emission efficiency, with the fluorescence quantum yield increasing from $\Phi_f = 0.0036$ in solution to $\Phi_f = 0.15$ in the aggregated state.

Conversely, the lack of significant AIE behavior of **i-DCSP** can be attributed to its substantial energy barrier in solution (1.40 eV), which increases only marginally (by 0.27 eV) upon aggregation. This molecular mechanism aligns with the experimental observations of Yokoyama *et al.*,¹⁶ who reported a high fluorescence quantum yield in solution ($\Phi_f = 0.43$) and a slight decrease in the aggregated state ($\Phi_f = 0.12$). The relatively lower barrier in the **i-DCSP** aggregated state compared to **o-DCSP** (by 0.69 eV) suggests weaker steric constraints, allowing some molecules to access radiation-less decay pathways.

These findings highlight the profound impact of cyano group positioning on the molecular conformation, excited-state dynamics, and photophysical properties. By employing QM/MM (and particularly QM/MM-FEP) calculations, we demonstrated a powerful approach for probing environment-dependent phenomena in complex molecular systems. Our results offer valuable design principles for next-generation AIE-active materials, indicating that molecular architectures that significantly increase the energy barriers upon aggregation hold the greatest promise for enhanced solid-state emission.

Moreover, by focusing our study on isomers with experimentally determined crystal structures, we established a reliable computational framework for quantifying and explaining intricate photophysical processes. This methodology not only

captures the subtle interplay between molecular structure, electronic properties, and environmental effects but also provides predictive capabilities for designing luminescent compounds with tailored emission profiles. These insights will serve as a foundation for the rational design of innovative AIE-active materials for specific technological applications.

Data availability

The quantum chemical calculations in this study were performed using Q-Chem 6.2 software. The optimized molecular geometries and energies for the S₀-Min and S₀/S₁-MECI structures of **o-DCSP** and **i-DCSP** are available from the corresponding author upon reasonable request. No new experimental data were generated as part of this theoretical study.

Conflicts of interest

The authors declare no conflicts of interest.

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