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QM/MM studies on luminescence mechanism of dinuclear copper iodide complexes with thermally activated delayed fluorescence†

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The QM/MM method is employed to investigate the photophysical mechanism of two dinuclear copper iodide complexes with thermally activated delayed fluorescence (TADF). The S_1 – T_1 energy differences (ΔE_{ST}) in these two complexes are small enough so that repopulating the S_1 state from T_1 becomes energetically allowed. Both forward and reverse intersystem crossing (ISC and rISC) processes are much faster than the corresponding radiative fluorescence and phosphorescence processes [k_{ISC} (10^8 s⁻¹) > k_f^F (10^6 s⁻¹), k_{rISC} (10^5 s⁻¹) > k_p^P (10^3 s⁻¹)]. The faster rISC process than the phosphorescence emission enables TADF. Moreover, the diphosphine ligands are found to play an important role in regulating the electronic structures and thereto the radiative and nonradiative rate constants. The present work rationalizes experimental phenomena and helps understand the intrinsic luminescence properties. The obtained insights could be useful for tuning the luminescence performance of dicopper-based luminescence materials.

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1. Introduction

As third generation emitting materials, organic and inorganic compounds with thermally activated delayed fluorescence (TADF) have been extensively studied in recent decades.^{1–9} Due to the use of inexpensive Cu metal, a large variety of luminescent Cu(I) compounds have been investigated since TADF was observed in Cu(I) complexes by Blasse and co-workers in 1980.¹⁰ In addition, TADF Cu(I) materials exhibit certain superior luminescence properties, for example, they harvest all singlets and triplets for the generation of light in organic light emitting diodes (OLEDs). The first successful application as emitting materials in OLEDs was reported in 1999.¹¹ Nowadays several classes of important structures that include three- and four-coordinated mononuclear, and four-coordinated dinuclear Cu(I) complexes have been applied in commercial OLEDs.¹² These complexes usually have distorted tetrahedral geometries around central Cu(I) atoms with diimine and diphosphine ligands. These quasi-tetrahedral structures often lead to low quantum yields due to efficient non-radiative decays. To

overcome this weakness a proposal of adopting binuclear Cu(I) complex structure was put forward, because similar structures between ground and emissive excited states in binuclear Cu(I) complexes result in an efficient emission.¹³

As a result, dinuclear Cu(I) complexes with monodentate or chelating phosphines have gained high experimental attention in the past decade.^{14,15} In 2007 the first device using iodo-bridged complexes was reported.^{16,17} In 2010 Deaton *et al.* reported a diamond core dinuclear Cu(I) complex with high TADF emission, which is comparable to high-performance phosphorescent emitting complexes with rare metals.¹⁸ Yersin *et al.* synthesized another dinuclear complex, which has a butterfly-shaped metal-halide core that leads to a small ΔE_{ST} of 0.05 eV for TADF emission.¹⁹ Later, they further prepared some attractive shorter-lived halide-bridged dicopper TADF compounds with chelating aminophosphane ligands.²⁰ The latest dinuclear Cu(I) TADF materials exhibit suitably short decay times and realize highly efficient OLEDs.²⁰ These bridges are often unstable especially in fluid environments, which however can be enhanced by bridging two Cu(I) centers with additional bidentate ligands.²¹

On the other hand, it is commonly known that TADF is highly sensitive to structural details of emitting materials. To achieve better performance it is usually needed to effectively restrict certain vibration modes of emitting molecules, for example, recently reported efficient TADF of mononuclear tetradentate Cu(I) complexes.²² However, the corresponding studies on dinuclear copper complexes are less studied. It is not clear whether the introduced bidentate ligands restrict certain

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vibrational modes of dinuclear Cu(I) complexes leading to efficient TADF.

Recently, Kato *et al.* have synthesized solution-stable luminescent dinuclear Cu(I) complexes *i.e.* $\text{Cu}_2(\mu\text{-I})_2(\text{dpppy})_2$ [dpppy = 2,3-bis(diphenylphosphino)pyridine] (Cu-py) using chelating diphosphine ligands with heteroaromatic rings.⁹ These compounds were found to exhibit interesting TADF phenomena from mixed (M + X)LCT excited singlet and triplet states. Further analysis clearly shows the molecular structure is almost identical to the parent $\text{Cu}_2(\mu\text{-I})_2(\text{dppb})_2$ [dppb = 1,2-bis(diphenylphosphino)benzene] (Cu-bz).¹⁶ The emission quantum yields of Cu-bz and Cu-py are measured to be 0.99 and 0.48 in the solid state at 298 K, respectively. The 548 nm emission wavelength of Cu-py is also redshifted compared to the 497 nm one of Cu-bz. In previous studies, these emission properties are simply discussed, but, detailed excited-state properties and underlying photophysical processes related to TADF, for example, forward and reverse intersystem crossing rates, competition between radiative and nonradiative transitions, are elusive and not explored computationally.

In this work we have carried out a comprehensive computational study on two dinuclear Cu(I) complexes [$\text{Cu}_2(\mu\text{-I})_2(\text{dppb})_2$] (**1**) and [$\text{Cu}_2(\mu\text{-I})_2(\text{dpppy})_2$] (**2**) with chelating diphosphine ligands (see Fig. 1). Their excited-state geometric and electronic structures, forward and reverse intersystem crossing rates (ISC and rISC), fluorescence and phosphorescence emission rates are explored using both density functional theory (DFT) and time-dependent DFT (TD-DFT) methods in combination with the quantum mechanics/molecular mechanics (QM/MM) approach. In terms of the results, radiative and nonradiative rates and related TADF mechanism of these dinuclear Cu(I) TADF emitters are discussed in detail.

2. Computational methods

Initial models are built based on X-ray crystal structures in experiments.⁹ Full optimizations of crystal structures including cell parameters and molecular geometric parameters are carried out at the PBE level with periodic boundary condition (PBC).²³ The DNP basis set is used to expand electronic wavefunction and the default dispersion correction of Grimme is added to consider weak dispersion interaction.^{24,25} A Monkhorst-Pack *k*-point grids of $2 \times 2 \times 2$ are used. Default convergence criteria on gradients and displacements are used in geometry optimizations. These calculations are performed using the DMol³ module in Materials Studio 2016.^{26–28}

In order to simulate excited-state properties of these Cu(I) complexes in crystals, a combined quantum mechanics/

molecular mechanics (QM/MM) method is used.^{29,30} The central one molecule is chosen as the QM region; whereas, the surrounding twelve molecules are chosen as the MM region and frozen during geometry optimizations (see Fig. S1†). The universal force field (UFF) and QM methods are used for the MM and QM regions, respectively.³¹ This strategy has been demonstrated to work well in previous works.^{32–34} The minima are optimized by the DFT (S_0) and TD-DFT (S_1 and T_1) methods with the hybrid exchange-correlation functional (PBE1PBE).^{35–37} In all these calculations, the Stuttgart relativistic pseudopotential and its accompanying basis set (SDD, ECP60MWB) are applied for the Cu and I atoms; the 6-31G* [6-31+G*] basis set is used for C, H and N [P] atoms.^{38–41} All QM/MM geometry optimizations are carried out using Gaussian 09 (ref. 42) and all vibrational frequencies are calculated using Gaussian 16.⁴³ Oscillator strengths and spin-orbit coupling matrix elements that are used for rate constants of radiative fluorescence and phosphorescence emission and nonradiative ISC (k_{ISC}) and rISC (k_{rISC}) of compounds **1** and **2** are calculated using the Amsterdam density functional ADF program^{44–46} in which the TD-PBE1PBE method and the TZP basis set^{47,48} are used with the zero order regular approximation (ZORA).^{49–51}

3. Results and discussion

3.1 Molecular structure

Important geometric parameters of two complexes in crystal as well as the corresponding experimental values are summarized in Table S1† and optimized molecular structures in the S_0 , S_1 , and T_1 states are schematically shown in Fig. 2.

In the S_0 state, two complexes have similar molecular structures and the structural parameters in crystal are consistent with experimental values.⁹ It also confirms the reliability of the PBE1PBE functional and the QM/MM method used. As shown in Fig. 2, the Cu atoms in the two complexes are coordinated with the two P atoms from the diphosphine ligands that form five-membered ring chelates and exhibit highly distorted tetrahedral coordination. The Cu–I and Cu–P bond lengths are comparable in the two complexes in crystal. The Cu...Cu distance is well-known to exert an evident impact on photophysical properties of the polynuclear Cu(I) halide complexes and the short Cu...Cu distance, less than 2.7 Å, usually generates emissive cluster-centered excited states.⁵² In crystal, they are 2.783 Å and 2.805 Å for **1** and **2**, respectively. The distances are more or less equal to the sum of van der Waals radius of Cu(I) of 2.8 Å, indicating a small interaction between these two Cu atoms. The similar changes can be found in the I...I distances. These shortened distances should come from the influence of the surrounding molecules.

For **1**, both I–Cu–I and P–Cu–P bond angles deviate largely from the usual tetrahedral value of 109.5°. In particular, the P–Cu–P bond angles are only 88.3° in crystal because of the small angle of the diphosphine ligand with respect to the five-membered ring chelate. The value 90.0° in crystal for compound **2** is larger slightly than that of compound **1**. The chelate effect of diphosphine ligands may lead to stable complexes **1** and **2**. In addition, two Cu(I) centers are bridged by

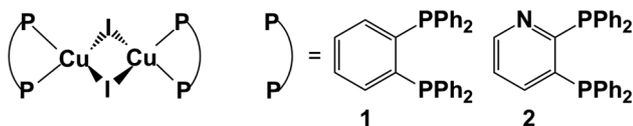


Fig. 1 Chemical structures of two Cu(I) complexes **1** and **2** studied in our present work. Also shown are two different substituent groups.



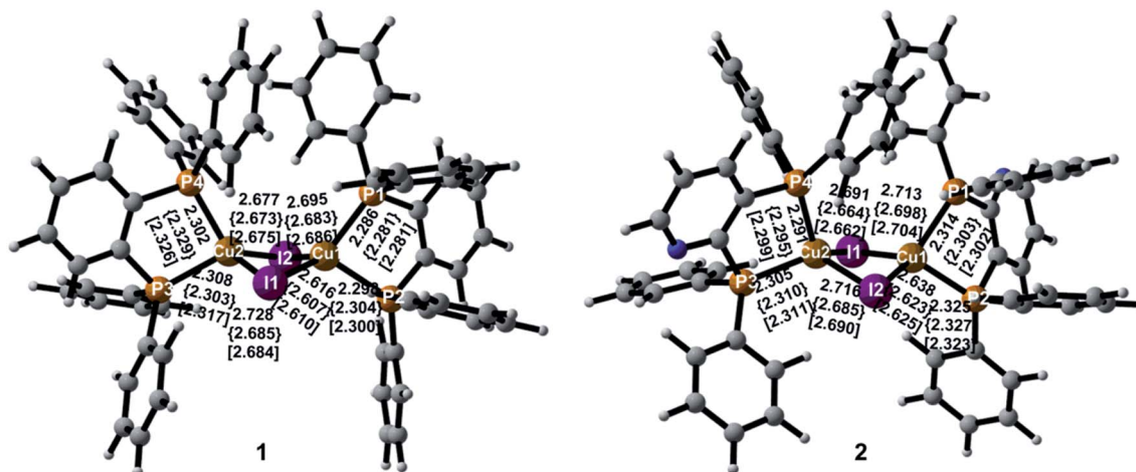


Fig. 2 Optimized minimum-energy structures with atomic numbering and selected geometric parameters in the S_0 , S_1 (curly brackets), and T_1 [square brackets] states.

two I atoms to form a dinuclear structure with a bent rhombic $\{Cu_2I_2\}$ core. The four-membered butterfly Cu_2I_2 rings of the complexes are significantly distorted by bending along the $I\cdots I$ axis, which is different from most of halide-bridged $Cu(I)$ complexes having a planar rhombic Cu_2X_2 geometry.^{53,54} The dihedral angles between the two CuI_2 triangles in the $\{Cu_2I_2\}$ core are 141.5° and 144.1° for **1** and **2** in crystal, respectively, implying a large steric hindrance of the diphenyl groups attached on the phosphines.

Different geometric changes of complexes **1** and **2** in the excited states are related to their luminescent properties. Comparing geometric parameters of S_0 , S_1 , and T_1 minima in crystal, we can see that the Cu–I bond lengths and the $I\cdots I$ distances are shortened for **1** and **2**. The changes of the $Cu\cdots Cu$ distances are different from those of the Cu–I bond lengths. For **1**, they are computed to 2.918 \AA and 2.934 \AA in S_1 and T_1 , respectively, which are longer than 2.783 \AA in S_0 . There are negligible changes for the Cu–P bond lengths in S_1 and T_1 . The same trend is also observed for compound **2**. In addition, Fig. 3 displays the difference of geometric parameters for **1** and **2** between S_0 and S_1 (T_1) states. It is worth mentioning that the S_1 and T_1 minima are structurally similar to each other, which may be caused by the dppb and dppy ligands constructing a sterically bulky coordination environment around the Cu atoms. As a result, the structural distortion of the MLCT excited state may be suppressed,⁵⁵ which will benefit the $Cu(I)$ complex to achieve high emission quantum yield, 0.99 for **1** and 0.48 for **2** in crystal.⁹ This corroborates experimental studies focused on the design of TADF for heteroleptic $Cu(I)$ complexes.^{54,55}

3.2 Frontier molecular orbitals

Frontier orbital analysis for compound **1** and its related orbital energies at the S_0 and S_1 minima are shown in Fig. 4. The frontier orbital characters of **1** and **2** at the T_1 minima as well as those of **2** at the S_0 and S_1 minima are displayed in Fig. S2 and S3,[†] respectively.

As shown in the left panel of Fig. 4, the highest occupied molecular orbital (HOMO) of compound **1** at the S_0 minimum is mainly derived from the d orbitals of the Cu centers and the p orbitals of the bridging I atoms. In contrast, the lowest unoccupied molecular orbital (LUMO) is localized at one dppb moiety of the bridging diposphine ligands. The nature of molecular orbitals of compound **1** at the S_1 minimum are almost constant except a little different weights from the different fragment groups (see the right panel of Fig. 4). The minimum-energy structures at both S_1 and T_1 states of **1** are similar to each other, thus subtly changes are found in the orbital analysis for the T_1 minimum compared with that for the S_1 one (see Fig. S2[†]). In addition, the calculations reveal that compositions for compound **2** at the S_0 and S_1 minima are almost identical with those of compound **1** and are thus placed in Fig. S3.[†]

Since the photoexcited electron and the remaining hole are spatially separated in the π^* orbital of the organic ligand and the 3d orbital of the Cu atom, the energy gap between HOMO and LUMO tends to be small enough to permit an inverse intersystem crossing process from the T_1 to S_1 states enabling TADF. At the S_0 minima, the HOMO–LUMO energy gaps are 4.07 and 3.97 eV for compounds **1** and **2**, respectively. However, they decrease to 3.30 and 3.14 eV at the S_1 minima and 3.26 and 3.15 eV at the T_1 minima for compounds **1** and **2**. Obviously, this remarkable reduction of the HOMO–LUMO energy gaps from the S_0 to S_1 minima of compounds **1** and **2** is responsible for the large Stokes shift experimentally observed in the emission spectra (see below, Fig. 5). On the other hand, the energy gap becomes smaller in compound **2** than that of **1**, which is caused by the introduction of the pyridyl N atom with large electronegativity to the aromatic ring that stabilizes the π^* orbitals. The similar variation of the HOMO–LUMO energy gaps is also found in recent theoretical works on dinuclear TADF $Cu(I)$ molecules.⁵⁷



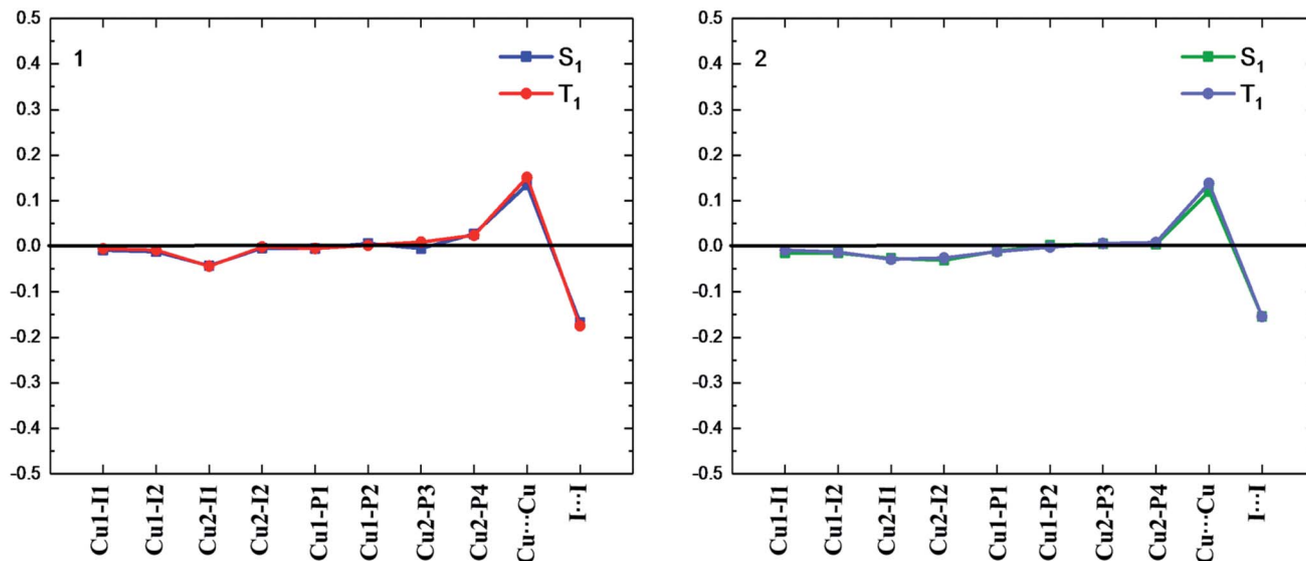


Fig. 3 Bond length differences (in Å) of the S₁ and T₁ minima for complexes 1 and 2 relative to the counterparts of the S₀ minimum (black line).

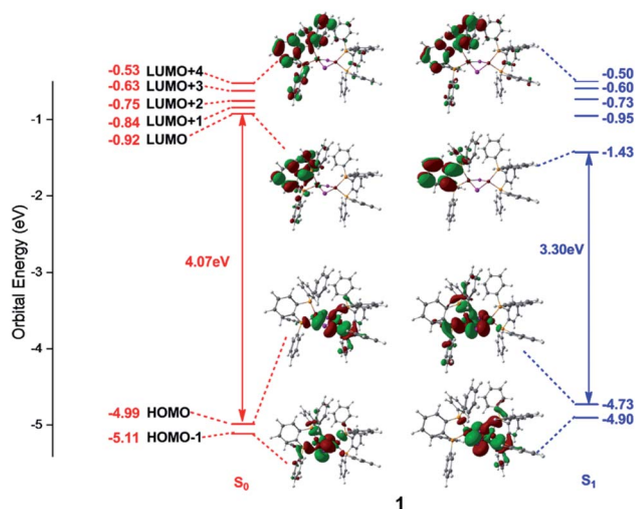


Fig. 4 Frontier molecular orbitals and related orbital energies (in eV) at the S₀ and S₁ minima of complex 1.

3.3 Absorption spectra

On the basis of the S₀ minima of compounds 1 and 2, we have simulated the corresponding absorption spectra. As shown in Fig. 5, there is a broad shoulder at *ca.* 400 nm in compound 1. Electronic structure analysis shows that the absorption at 374 nm mainly stems from the S₀ → S₁ electronic excitation, for which the HOMO–LUMO electronic configuration contributes the most. The HOMO is mainly distributed over the Cu and I atoms, while the LUMO is localized on the dppb ligands. Thus, this absorption peak is of the mixed metal-to-ligand and halide-to-ligand charge transfer (M + X)LCT character. The origin of the lowest absorption of compound 2 is similar to that of 1 and the calculated absorption of 2 at 392 nm has a small redshift compared to that of 1 at 374 nm, which could be due to the

stabilization of the π* orbitals of the bridging diphosphine ligands introduced by the pyridyl N atom.

For 1, the computed 331 nm peak is very close to the experimentally measured 330 nm peak in crystal. The 332 nm peak of 2 is almost equal to that in 1 and also reproduces the experimental value of 328 nm.⁹ These peaks are mainly caused by the electronic excitation transitions of S₀ → S₈ for 1 and S₀ → S₁₀ for 2. Moreover, these two complexes have strong absorption peaks at 291 nm (see Table S3[†]), which are also assigned to the electronic transitions H → L+1 for 1 and H – 1 → L+7 for 2, respectively. They are of the same electronic transition character *i.e.* (M + L)CT (see Fig. S3[†]).

3.4 Emission properties

Vertical emission energies and related electronic configurations of compounds 1 and 2 in crystal from their S₁ and T₁ excited states are calculated and listed in Table 1, which also includes wavelengths and oscillator strengths. The corresponding fluorescence emission spectra based on the S₁ minima are simulated in Fig. 5.

In experiments, the emission spectrum of the dinuclear complex 1 in crystal is very similar to that of the mononuclear complex [CuI(dppb)PPh₃] (λ_{em} = 492 nm).¹⁶ This can be understood very well taking into account that the Cu...Cu distance of 1 is close to the sum of the van der Waals radii of two Cu atoms (see above). This results in very weak interaction between two Cu atoms in both ground- and excited-states; thus, the emission from 1 resembles that of the mononuclear complex.¹⁶

From Table 1, we can find that the HOMO–LUMO electronic configuration is predominantly responsible for the S₁ → S₀ and T₁ → S₀ electronic de-excitation transitions, which are also of mixed metal-to-ligand and halide-to-ligand charge-transfer character (M + X)LCT. By contrast, the introduction of the N atoms to the cross-linking phenyl groups of the diphosphine



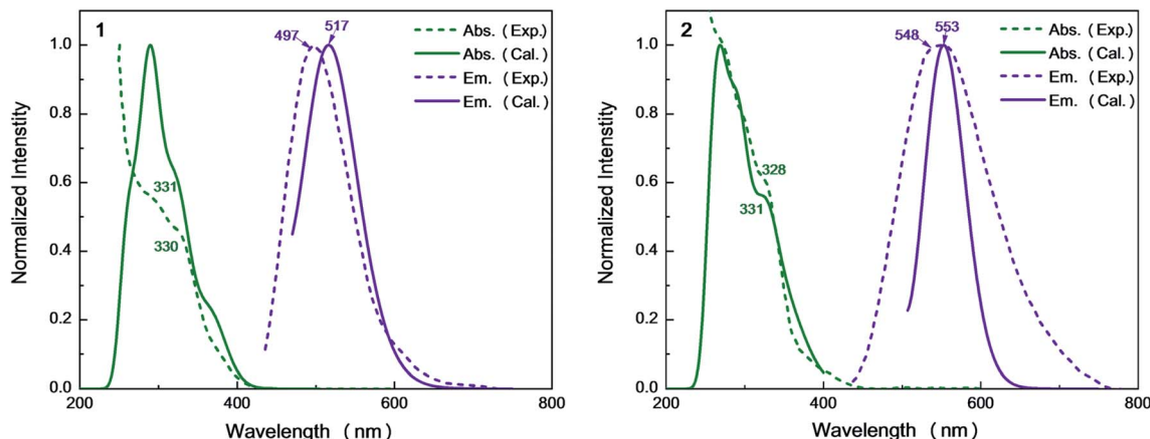


Fig. 5 Simulated absorption and emission spectra of complexes 1 and 2 with experimental results.

ligands in 2 greatly shifts the emission from bright-greenish-blue ($\lambda_{\text{em}} = 497$ nm) to yellow ($\lambda_{\text{em}} = 548$ nm) as observed in experiments.⁹ The calculated λ_{max} values of the dinuclear complexes 1 and 2 are in the order 1 (517 nm) < 2 (553 nm), which are consistent with the experimental values in crystal.⁹

3.5 Excited state properties

Excited-state properties play an important role in determining luminescence mechanism. In order to determine quantitatively related radiative and nonradiative processes of compounds 1 and 2, we have calculated rate constants of forward and inverse intersystem crossing processes between S_1 and T_1 [ISC (k_{ISC}) and rISC (k_{rISC})] and those of fluorescence (k_{r}^{F}) and phosphorescence (k_{r}^{P}) from S_1 and T_1 in crystal.

3.5.1 ISC rate constants. The radiationless rate constants for the intersystem crossing processes are obtained by using the classical Marcus theory,^{58–60} which has been extensively used recently.^{61,62} Its basic formulae are briefly presented below to form a self-contained work. The rate constant from the initial *i* to final *f* electronic states is expressed by the classical Fermi golden rule:

$$k_{i \rightarrow f} = \frac{2\pi}{\hbar} |H_{if}|^2 \rho_{\text{FC}} \quad (1)$$

where H_{if} is the effective spin-orbit coupling between the initial and final states. In the classical regime, this formula could be expressed by the famous Marcus equation where the Franck-Condon weighted density of states, ρ_{FC} , is written as

$$\rho_{\text{FC}} = \frac{1}{\sqrt{4\pi\lambda_{\text{reorg}}k_{\text{B}}T}} \exp\left[-\frac{(\Delta E + \lambda_{\text{reorg}})^2}{4\pi\lambda_{\text{reorg}}k_{\text{B}}T}\right] \quad (2)$$

in which k_{B} is the Boltzmann constant; ΔE is defined as the difference between adiabatic energies of the final and initial states; T is the temperature; λ_{reorg} is the Marcus reorganization energy. In terms of these formula, one can find that ISC (k_{ISC}) and rISC (k_{rISC}) rates are closely related to spin-orbit coupling (SOC) values, singlet-triplet energy differences (ΔE_{ST}), and reorganization energies (λ_{reorg}). In the next sections, we will discuss these factors separately to gain in-depth insights. The computed rate constants as well as spin-orbit coupling matrix elements (SOCMEs), and reorganization energies (λ_{reorg}) are collected in Table 2.

Table 2 Spin-orbit coupling matrix elements (SOCMEs/cm⁻¹), reorganization energies (λ_{reorg} /eV), intersystem crossing rates (k_{ISC} /s⁻¹), reverse intersystem crossing rates (k_{rISC} /s⁻¹), fluorescence rate constants ($k_{\text{r}}^{\text{F}} \times 10^6$ /s⁻¹), and phosphorescence rate constants ($k_{\text{r}}^{\text{P}} \times 10^3$ /s⁻¹) of compounds 1 and 2 in crystal

	At S_1		At T_1	
	1	2	1	2
SOCMEs (cm ⁻¹)	13.14	18.57	8.34	20.72
λ_{reorg} (eV)	0.0256	0.0108	0.0241	0.0096
$k_{\text{ISC}}/k_{\text{rISC}}$ (s ⁻¹)	8.25×10^8	2.34×10^9	6.27×10^5	3.36×10^7
$k_{\text{r}}^{\text{F}} \times 10^6$ (s ⁻¹)	3.91	0.33	—	—
$k_{\text{r}}^{\text{P}} \times 10^3$ (s ⁻¹)	—	—	1.11	2.70

Table 1 Emission energies (E /eV), wavelength (λ /nm), oscillator strengths (f), electronic configurations, and characters involved in the $S_1 \rightarrow S_0$ and $T_1 \rightarrow S_0$ electronic transitions of complexes 1 and 2 in crystal

	State	E (eV)	λ (nm)	f	Electronic configuration	Assignment
1	S_1	2.40	517	0.0157	H \leftarrow L (94.4%)	(M + X)LCT
	T_1	2.19	567	0.0000	H \leftarrow L (93.1%)	(M + X)LCT
2	S_1	2.24	553	0.0015	H \leftarrow L (96.8%)	(M + X)LCT
	T_1	2.14	579	0.0000	H \leftarrow L (93.3%)	(M + X)LCT



First, we focus on the SOCMEs values of compounds **1** and **2** calculated at the S_1 and T_1 minima in crystal (see Table 2). After comparing these values, one can find that the SOC value 13.14 cm^{-1} of compound **1** at the S_1 minimum is a little larger than that 8.34 cm^{-1} at the T_1 minimum; but, the SOC value 18.57 cm^{-1} of compound **2** at the S_1 minimum is tiny smaller than 20.72 cm^{-1} at the T_1 minimum. These SOCMEs values are comparable to other complexes with TADF and are large enough to ensure effective forward and reverse intersystem crossing processes.

Second, it is generally accepted that singlet-triplet energy difference (ΔE_{ST}) is an important parameter to regulate reverse ISC processes. Table 3 lists the energies of the S_1 and T_1 minima of compounds **1** and **2** in crystal. One can see the S_1 and T_1 states' energies of compound **2** are a little decreased compared to those of compound **1**. For example, the values 2.75 eV in the S_1 state and 2.66 eV in the T_1 state of compound **2** are smaller than 2.89 eV and 2.74 eV in S_1 and T_1 for compound **1**, respectively. Nevertheless, the ΔE_{ST} values are still small in both compounds, *i.e.* 0.15 eV in **1** and 0.09 eV in **2**. Considering that both **1** and **2** have small ΔE_{ST} values, which make their rISC processes from the T_1 to S_1 states possible in the view of energy. Meanwhile, we have also calculated higher-lying triplet excited states to judge whether they are also involved in TADF. From Table S2,[†] it is clear that there is only a triplet state *i.e.* T_1 lower than S_1 in energy for compounds **1** and **2**. In other words, no higher-lying triplet excited states are involved in the rISC processes.

Finally, we analyze the changes of reorganization energies (λ_{reorg}), which can help us understand the radiationless processes either from S_1 to T_1 or from T_1 to S_1 . The reorganization energy is calculated through the total Huang–Rhys factor S , which is collected as a sum of the Huang–Rhys factors of all vibrational frequencies S_j via the following equation:

$$S = \sum_j^n S_j = \sum_j^n \frac{\omega_j}{2\hbar} \Delta Q_j^2 \quad (3)$$

where ω_j is the normal mode frequency of the j th mode of the initial state and ΔQ_j is the normal mode displacement between the initial and the final states. The ω_j can be obtained *via* diagonalizing the mass-weighted Hessian matrix of the optimized initial state: $L^T H L = \omega^2$; in the same time, the transformation matrix L is obtained.

In order to calculate ΔQ_j , the following relationship for Cartesian coordinates is defined:

$$\mathbf{q}_{\text{init}} = \mathbf{q}_{\text{final}} + \Delta \mathbf{q} \quad (4)$$

in which \mathbf{q}_{init} and $\mathbf{q}_{\text{final}}$ represent mass-weighted Cartesian coordinates of initial and final states; then, the normal mode coordinates \mathbf{Q} can be transformed from Cartesian coordinates \mathbf{q} using the relation $\mathbf{Q} = L^T \mathbf{q}$. Finally, we arrive at the following equation:

$$\mathbf{Q}_{\text{init}} = L_{\text{init}}^T L_{\text{final}} \mathbf{Q}_{\text{final}} + L_{\text{init}}^T \Delta \mathbf{q} \quad (5)$$

where L_{final} and L_{init} correspond to the transform matrices of the final and initial states that diagonalize the corresponding mass-weighted Hessian matrices. $L_{\text{init}} L_{\text{final}}^T (=D)$ is called Duschinsky rotation matrix in the literature, so $\Delta \mathbf{Q}$ could be calculated by $\Delta \mathbf{Q} = L_{\text{init}}^T \Delta \mathbf{q}$. Once S and S_j are calculated, the reorganization energy

$$\lambda_{\text{reorg}} = \sum_j^n \lambda_{\text{reorg}_j} = \sum_j^n S_j \omega_j \hbar \quad (6)$$

can be calculated in a straightforward way.

From Table 2, the reorganization energies λ_{reorg} for compound **1** either from S_1 to T_1 (0.0256 eV) or from T_1 to S_1 (0.0241 eV) are much close to each other. For compound **2**, the λ_{reorg} values from S_1 to T_1 or from T_1 to S_1 are similar in crystal, 0.0108 eV from S_1 to T_1 vs. 0.0096 eV from T_1 to S_1 . The reorganization energies λ_{reorg} of compound **1** either from S_1 to T_1 or from T_1 to S_1 are slightly larger than those of compound **2** in crystal phase. In addition, we have examined the distribution of the Huang–Rhys factors. Fig. S5[†] depicts the Huang–Rhys factor for each vibrational mode involved in the ISC and rISC processes of compounds **1** and **2** in crystal. The low-frequency vibrational modes related to the torsional motion of the dppb and dpppy groups are mainly responsible for these intersystem crossing processes between S_1 and T_1 (either ISC or rISC). Comparing these Huang–Rhys factors, one can find that the vibrational modes with large Huang–Rhys factors in the low-frequency region are suppressed to certain extent in compound **2**. This is consistent with the results of λ_{reorg} discussed above.

On the basis of SOC, ΔE_{ST} , and λ_{reorg} , we have calculated the related intersystem crossing rates k_{ISC} and k_{rISC} of these two Cu(I) dimers in crystal. The k_{ISC} and k_{rISC} values of compound **2** are larger than those of compound **1** at room temperature. For example, the k_{ISC} and k_{rISC} values of compound **2** (2.34×10^9 and $3.36 \times 10^7 \text{ s}^{-1}$) are larger than those of compound **1** (8.25×10^8 and $6.27 \times 10^5 \text{ s}^{-1}$), respectively. This change trend is in good agreement with that of involved SOC values (see Table 2). Moreover, the rates k_{rISC} of either compounds **1** or **2** are smaller than their forward intersystem crossing rates k_{ISC} in crystal. For example, the k_{rISC} and k_{ISC} values are predicted to be 6.27×10^5 [3.36×10^7] vs. 8.25×10^8 [$2.34 \times 10^9 \text{ s}^{-1}$] for **1** [**2**]. This is because the forward ISC process is energetically favorable whereas the reverse one is merely active when the thermal energy provided by the temperature can overcome the related reverse barriers. Finally, we have checked the temperature effects on these intersystem crossing rates k_{ISC} and k_{rISC} (see Fig. 6). Obviously, both k_{ISC} and k_{rISC} rates of compounds **1** and **2** increase with higher temperature, which is caused by the fact that more vibrationally excited states are involved in the

Table 3 Relative energy (eV) of PBE1PBE optimized structures of complexes **1** and **2** in crystal and their energy differences

	1	2
S_1	2.89	2.75
T_1	2.74	2.66
$\Delta E_{S_1-T_1}$	0.15	0.09



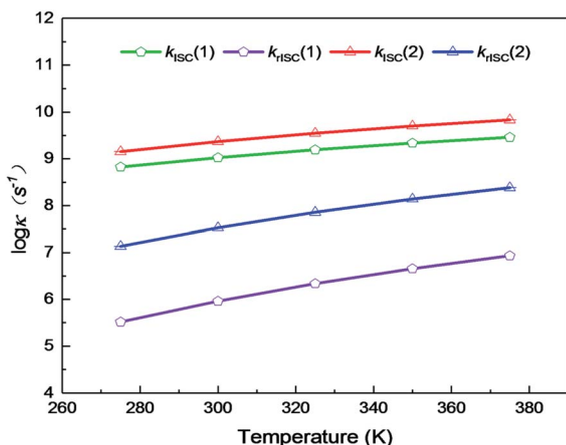


Fig. 6 Temperature dependence of ISC and rISC rates of complexes 1 and 2 from 275 to 375 K.

intersystem crossing processes when the temperature increases.⁵⁸ Furthermore, one can see that the reverse intersystem crossing rate constants k_{rISC} are more sensitive to the temperature than those of the forward ones k_{ISC} in the temperature range of 275 to 375 K.

From above eqn (1) and (2), it is clear that k_{ISC} and k_{rISC} rate constants are dependent on not only SOC values but also $(\Delta E + \lambda_{reorg})^2$. Fig. 7 shows the relationship of the forward and reverse ISC rate constants and ΔE for compounds 1 and 2 in crystal. In the equations, ΔE is defined as the difference between adiabatic energies of the final and initial states. For the reverse ISC processes from T_1 to S_1 , $\Delta E_{ST} = (E_{S_1} - E_{T_1})$ is positive for our studied systems, so k_{rISC} decreases upon increasing ΔE because λ_{reorg} is a fixed positive value. By contrast, k_{ISC} is affected by both ΔE and λ_{reorg} through $(\Delta E + \lambda_{reorg})^2$ because $\Delta E_{TS} = (E_{T_1} - E_{S_1})$ is negative for our studied systems. Specifically, when ΔE_{ST} is larger than the reorganization energy λ_{reorg} , k_{ISC} decreases upon increasing ΔE_{TS} . Because $\Delta E_{TS} = -\Delta E_{ST}$, k_{ISC} is found to

decrease with decreasing ΔE_{ST} . More interestingly, the curvatures of parabolic functions for the k_{ISC} and k_{rISC} rate constants of compound 1 as a function of ΔE_{ST} are slightly smaller than those of compound 2 (see Fig. 7). This can be understood based on eqn (2) and (3). The logarithmic function of the rate constant k can be rewritten as follows

$$\ln k = \text{constant} - \frac{(\Delta E + \lambda_{reorg})^2}{4\lambda_{reorg}k_B T} \quad (7)$$

in which

$$\text{constant} = \ln \left(\frac{1}{h} |H_{if}|^2 \sqrt{\frac{\pi}{\lambda_{reorg} k_B T}} \right) \quad (8)$$

From this equation, it is clear that the function curvature is proportional to $|1/\lambda|$ when the temperature is fixed. Because the λ values for compound 1 are larger than those of compound 2, as shown in Table 2, the former curvatures are smaller than the latter ones. In addition, one can find that the parabolic peaks are always at the points of $\Delta E = -\lambda_{reorg}$ where the logarithmic function in eqn (7) achieves a maximum value.

3.5.2 Radiative rates. As mentioned in previous works,⁶³ highly luminescent TADF materials should have an efficient radiative decay rate (more than 10^6 s^{-1}) to overcome competitive nonradiative pathway from S_1 to S_0 . Accordingly, we have calculated the rate constants of fluorescence (k_r^F) and phosphorescence (k_r^P) according to the following formula using the ADF software:

$$k_r^{F/P} = \frac{2\pi\nu^2 e^2}{\epsilon_0 m c^3} f_{S_1 \text{ or } T_1 \rightarrow S_0} \quad (9)$$

in which ν is the emission energy; e is the elementary electric charge; ϵ_0 is the vacuum permittivity; m is the mass of electrons; c is the speed of light; and $f_{S_1 \text{ or } T_1 \rightarrow S_0}$ is the oscillator strength for $S_1 \rightarrow S_0$ or $T_1 \rightarrow S_0$ radiative transition.

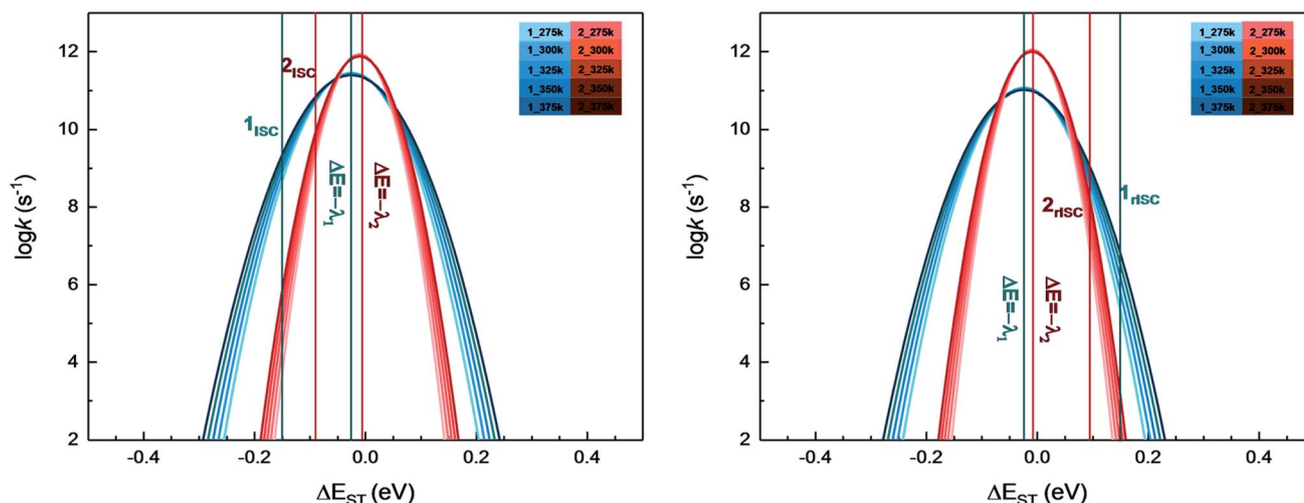


Fig. 7 Forward (left) and reverse (right) intersystem crossing rate constants ISC and rISC as a function of ΔE_{ST} at different temperatures of complexes 1 (blue) and 2 (red).



From Table 2, it can be found that the fluorescence and phosphorescence emission rates of compound **2** are smaller than those of compound **1** in crystal. The k_r^F rate of compound **2** is $0.33 \times 10^6 \text{ s}^{-1}$, which is smaller than $3.91 \times 10^6 \text{ s}^{-1}$ of compound **1**. Moreover, one can see that the phosphorescence emission rates are much smaller than the fluorescence emission rates (*ca.* 10^3 vs. 10^6 s^{-1}). Our presently calculated fluorescence radiative rates k_r^F are in good agreement with experiments, in which they are estimated to be 0.25×10^6 and $0.12 \times 10^6 \text{ s}^{-1}$ for compounds **1** and **2**, respectively. These efficient radiative decay rates may overcome competitive non-radiative pathway from S_1 to S_0 to enable TADF.

3.6 TADF mechanism

The existence of TADF indicates not only that the S_1/T_1 SOC should be significant for efficient reverse ISC but also that the S_1-T_1 energy gap has to be small enough so that thermal energy can overcome this gap efficiently. In the following, we will discuss the TADF mechanism of compounds **1** and **2**.

A three-state model of the S_0 , S_1 , and T_1 states is enough and used to study the TADF mechanism in these two Cu(I) dimer compounds because the T_2 state is energetically higher than the S_1 and T_1 states at either S_1 or T_1 minima as discussed above. Fig. 8 summarizes a reasonable speculation about the TADF working mechanism of compound **1**. This compound can effectively convert its S_1 to T_1 states with an ISC rate constant of $8.25 \times 10^8 \text{ s}^{-1}$; importantly, the rISC process back to the S_1 state is also very faster at the room temperature ($6.27 \times 10^5 \text{ s}^{-1}$). These ISC and rISC rates are two orders of magnitude larger than the fluorescence and phosphorescence emission rates from the S_1 and T_1 states [$k_{\text{ISC}} (10^8 \text{ s}^{-1}) > k_r^F (10^6 \text{ s}^{-1})$, $k_{\text{rISC}} (10^5 \text{ s}^{-1}) > k_r^F (10^3 \text{ s}^{-1})$; see Table 2]. In addition, the phosphorescence emission rates are overall about 3 orders of magnitude smaller than the fluorescence emission rates. Thereby, compound **1** mainly fluoresces from the S_1 state because the T_1 population can be quickly transferred back to the S_1 state through the fast rISC process. The internal conversion (IC) process from S_1 to S_0 and the ISC process from T_1 to S_0 are much slower due to huge energy gaps between S_1 and S_0 and between T_1 and S_0 , so these nonradiative decay rates are neglected in our model.⁶⁴ Even though the k_r^F rate of compound **2** is smaller than that of compound **1**, the TADF mechanism of **2** is similar to that

of **1**. The present analysis gives in-depth understanding on the TADF properties of the two dinuclear Cu(I) iodide complexes.⁹

The present results for the TADF of dinuclear Cu(I) iodide complexes also show that structural rigidity around the central Cu(I) atom is important. First, it benefits rapid intersystem crossing processes between singlets and triplets. Second, efficient deactivation pathways due to structural distortion are effectively suppressed. For example, the emission of dinuclear Cu(I) complexes with less-bulky ligands, *i.e.* $[\text{Cu}(\text{PPh}_3)_2(\text{H}_2\text{O})]\{\mu\text{-MeOpyz}\}\{\text{Cu}(\text{PPh}_3)_2(\text{H}_2\text{O})\}(\text{BF}_4)_2$ with $\text{PPh}_3 =$ triphenylphosphine and $\text{MeOpyz} = 2\text{-methoxyppyrazine}$, is observed at 580 nm with a remarkably lower quantum yield of 0.05, which has been proved due to efficient excited-state deactivation through structural distortion around the Cu atom from tetrahedral to quasi-square-planar coordination structures.⁵⁶

Finally, TADF is very sensitive to subtle structural changes of emitting materials. Our studied two compounds are not similar because they have distinctly different ligands: one is related to pyridine and the other is related to benzene. Both ligands indeed brings us significant differences. First, the energy gap becomes smaller in compound **2** than that of compound **1**, which is caused by the introduction of the pyridyl N atom with large electronegativity to the aromatic ring that stabilizes the π^* orbitals. Second, the introduction of the N atom to the cross-linking phenyl groups of the diphosphine ligands in compound **2** greatly shifts the emission from bright-greenish-blue (497 nm) to yellow (548 nm). Third, the k_{ISC} and k_{rISC} values of compound **2** are larger than those of compound **1** at room temperature. For example, the k_{ISC} and k_{rISC} values of compound **2** are larger than those of compound **1**, respectively (see Table 2). Fourth, the fluorescence and phosphorescence emission rates of compound **2** are smaller than those of compound **1** in crystal.

It should be noted that the TADF mechanism of copper-based organometallic compounds is rather different from those of pure organic TADF emitters. Recently, Penfold *et al.* have summarized many efficient organic TADF emitters and related TADF mechanisms.⁶⁵ For example, different D-A-D molecules with very similar energy gaps exhibit large variations in k_{rISC} . They found that luminescence could be switched from TADF to phosphorescence by sterically hindering the motion of D and A group. Interested readers are referred to this literature.⁶⁵

4. Conclusions

The structures, excited properties, and luminescence of two dinuclear Cu(I) complexes with dppb and dppy ligands $[\text{Cu}_2(\mu\text{-I})_2(\text{dppb})_2]$ (**1**) and $[\text{Cu}_2(\mu\text{-I})_2(\text{dppy})_2]$ (**2**) as potential TADF emitters have been comprehensively studied by means of combined QM/MM calculations. On the basis of ground and excited-state geometries we have found that the S_1 and T_1 states have very small energy gaps ΔE_{ST} , less than 0.15 eV, which makes the forward and reverse intersystem crossing ISC and rISC processes are much faster than the corresponding fluorescence and phosphorescence radiative

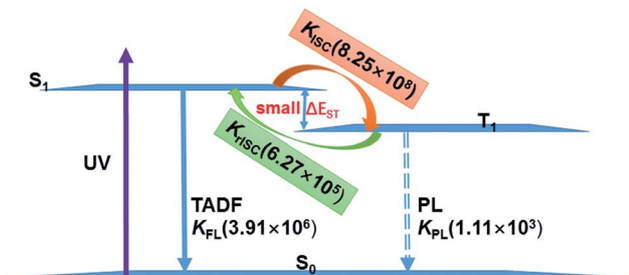


Fig. 8 TADF working mechanism of complex **1** including related radiative and radiationless rates calculated by our present theoretical studies.



processes [$k_{\text{ISC}} (10^8 \text{ s}^{-1}) > k_{\text{r}}^{\text{F}} (10^6 \text{ s}^{-1})$, $k_{\text{rISC}} (10^5 \text{ s}^{-1}) > k_{\text{r}}^{\text{P}} (10^3 \text{ s}^{-1})$]. Since the reverse rISC rates are much faster than the phosphorescence emission rates, the T_1 population can be quickly transferred to the S_1 state leading to a delay fluorescence emission. Through analyzing the Huang–Rhys factors we have found that the low-frequency vibrational modes related to the torsional motion of the dppb and dpppy groups are mainly in charge of these intersystem crossing processes ISC or rISC. Our present calculations not only elucidate recent experimental phenomena but also help understand, and design similar TADF emitters, e.g. with rigid ligands to inhibit structural distortions of emitting states and nonradiative processes, etc.

Conflicts of interest

The authors declared that they have no conflicts of interest to this work.

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