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# **REVIEW**

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# Intramolecular singlet fission and triplet exciton harvesting in tetracene oligomers for solar energy conversion

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Singlet fission (SF), which can convert one singlet exciton into two triplet excitons, has the potential to enhance the solar energy conversion efficiency in devices such as photovoltaic cells and processes such as photocatalysis. Recently, intramolecular SF (iSF), which is observed in covalently linked molecular oligomers, has attracted significant research interest due to its unique advantages in revealing the SF mechanism and practical applications. In the case of exothermal chromophores (their singlet state energy  $(E_{S_i})$  is more than twice their triplet state energy  $(E_{T_i})$ , such as pentacene), efficient iSF is easily achieved. However, efficient iSF is limited in endothermic/isothermal chromophores ( $E_{S_{L}} \le 2E_{T,L}$ , such as tetracene). Compared to exothermal systems, the energy loss in the SF of endothermic/isothermal systems is much lower, which is more favorable for enhancing the solar energy conversion efficiency. Therefore, achieving efficient iSF in endothermic/isothermal systems is necessary. Unfortunately, the design strategy for efficient iSF based on endothermic/isothermal chromophores is not clear. Herein, we summarize the SF studies on covalently linked tetracene oligomers and analyze the key influencing factors on iSF in these systems. Additionally, we review the harvesting of the two formed triplets from the iSF of tetracene oligomers. It is hoped that the results of this review will provide some new insights into the design of highly efficient iSF materials based on endothermic/isothermic chromophores for practical applications.

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## Introduction

Singlet fission (SF) is a spin-allowed multiple-exciton generation (MEG) process, which can split one high-energy singlet exciton into two low-energy triplet excitons. When SF is incorporated in the solar energy conversion process, it is expected to successfully achieve single-photon excitation to produce two electrons and enhance the efficiency of the related process.1-5 For



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example, the Shockley–Queisser (SQ) limit of the photoelectric conversion efficiency of single-junction solar cells was improved from 33% to 45% upon the incorporation of SF.<sup>6,7</sup> Beside solar cells, SF has also been applied to enhance the quantum efficiencies of organic light-emitting diodes,<sup>8</sup> organic photodetectors,<sup>9,10</sup> photon multiplier devices<sup>11</sup> and nonlinear optical responses.<sup>12,13</sup> Recently, it has been reported that SF also shows potential application in quantum information and computing.<sup>14</sup>

SF was first observed in anthracene crystals in 1965,15 and subsequently confirmed in tetracene in the 1970s. 16-18 However, in the following decades, further advances in this field were negligible. It was only in 2006 that Hanna and Nozik found that SF has great potential for enhancing the efficiency of solar cells. Subsequently, SF has shown vigorous development, and great efforts have been devoted to developing new efficient SF materials and revealing the SF mechanism. Many chromophores have been found to meet the energy requirement of SF, where the singlet state energy  $(E_{S_s})$  should be larger or equal to twice the triplet state energy  $(E_{T_1})$   $(E_{S_1} \le 2E_{T_1})$ , such as acenes (anthracene,19 tetracene,20 pentacene,21-24 hexacene25-27 and their derivatives), polyene,28-34 1,3-diphenylisobenzofuran,35 diketopyrrolopyrrole,36-40 perylenediimide,41-46 boron dipyrromethene (BODIPY), 47-50 and other chromophores. 37,51-61 These molecules have been summarized in some comprehensive reviews on this field.7,53,62-65

Based on these chromophores, several intermolecular SF (xSF) materials, including crystal films, colloidal nanoparticles and concentrated solutions, have been developed. The study of the SF in these systems gave some insights into the SF mechanism and showed that the SF rate and efficiency are strongly dependent on the intermolecular packing, morphology and defects. However, the determination of the structures of xSF materials is challenging, and consequently it is difficult to establish their structure–property relationship. Therefore, intramolecular SF (iSF), which is observed in covalently linked molecular oligomers, has attracted significant research interest due to the well-defined structure and facile structural modification of these materials.<sup>24,66,67</sup> These advantages are more feasible for the establishment of the relationship between the molecular structure and SF properties.<sup>7,32,68-71</sup> More importantly,



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the inherent nature of iSF in these oligomers is not restricted by external conditions and iSF materials can be easily incorporated in solar energy conversion devices *via* solution processing techniques and still maintain their SF properties.<sup>72</sup>

To date, several iSF materials based on different chromophores have been prepared and their SF properties investigated.32,44,73-85 In the case of exothermic chromophores  $E_{\rm S_1} > 2E_{\rm T_2}$ , such as pentacene, regardless of their configuration, efficient iSF can be easily achieved.86,87 However, efficient iSF is difficult to achieve in endothermic or isothermic systems ( $E_{S_1} \le$  $2E_{\rm T}$ ) (such as tetracene, diketopyrrolopyrrole, and perylene diimide). Compared to exothermic systems, endothermic or isothermic systems exhibit lower energy loss in the SF process, which have greater potential to enhance the overall solar energy conversion efficiency. Thus, it is necessary to achieve efficient iSF in endothermic or isothermic systems. Presently, although there are some comprehensive reviews on the SF mechanism in the literature,<sup>7,26,62,88-93</sup> strategies for the design of efficient iSF materials based on endothermic or isothermic chromophores remain unclear. The revelation of the key factors influencing the iSF of endothermic or isothermic systems is crucial for designing novel efficient iSF materials based on these chromophores. In this review, firstly, we summarize SF studies on covalently linked tetracene oligomers, and then list several key factors on the iSF in these systems. Additionally, we review the study of harvesting two triplets from iSF in tetracene oligomers. Taking tetracene as an example, it is hoped that this work will provide some insights for the design of efficient iSF materials based on endothermic or isothermic chromophores and the application of iSF in solar energy conversion processes.

## 2 SF mechanism

Here, the generally accepted SF process is roughly depicted in Fig. 1. $^{7,65}$  Upon photoexcitation, the singlet state (S<sub>1</sub>) is

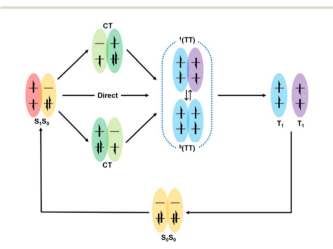


Fig. 1 Description of SF mechanism.  $S_0S_0$ : ground state;  $S_1S_0$ : singlet state; and CT: charge transfer state, where this intermediate CT state may be virtual (in the form of a super-exchange mechanism) or real.  $^1$ (TT): triplet pair state and  $^5$ (TT): quintet state (spin evolution can be performed between the multiexciton  $^1$ (TT) and multiexciton  $^5$ (TT) state).  $T_1$ : free triplet state.

generated, which collides with another adjacent molecule in the ground state  $(S_0)$  to produce the correlated triplet pair state (1(TT)) with an overall singlet spin character. This process can proceed via a direct or indirect mechanism. In the direct mechanism, the S<sub>1</sub> state converts into the <sup>1</sup>(TT) state without the involvement of any intermediate. Alternatively, the S<sub>1</sub> state can transform into the <sup>1</sup>(TT) state via an intermediate state that has a charge-transfer (CT) nature, which is the so-called indirect mechanism.7,63

Additionally, a coherent superposition state of the lowest lying absorbing state, CT state, and multiexciton state is formed upon photoexcitation, which can drive the SF process. This is the so-called quantum coherent mechanism. 88,89,94 Then, the two free triplets  $(T_1)$  can be formed *via* the decoherence of either the <sup>1</sup>(TT) state or the <sup>5</sup>(TT) state. Generally, the <sup>5</sup>(TT) state is formed via the spin evolution of the <sup>1</sup>(TT) state.<sup>7,89,92</sup> Initially, most of the research on SF only focused on the first step, namely, the formation of the <sup>1</sup>(TT) state. Subsequently, the dissociation of the <sup>1</sup>(TT) state into the T<sub>1</sub> state has also received attention. However, due to the similar transient absorption spectral feature between the <sup>1</sup>(TT) state and the T<sub>1</sub> state as well as the possible intermediate state (3(TT) or 5(TT)), this dissociation process is difficult to be clarified by transient absorption spectra alone.95 Fortunately, the electron spin change in these states allows the use of spin resonance techniques to resolve this dissociation process.96,97

The above-mentioned rough SF mechanism is suitable for xSF and iSF. Many iSF molecules demonstrate a direct SF process, 68,98-108 in which the iSF rate is dependent on the direct coupling strength between the S<sub>1</sub> state and the <sup>1</sup>(TT) state. The covalent linker plays an important role in iSF by controlling the coupling strength via through-bond or through-space interactions. Generally, conjugated linker-bridged oligomers exhibit much faster SF than bridged by nonconjugated linkers due to the stronger electronic coupling.106 However, a CT-assisted mechanism is also adopted in other iSF materials, as evident by the solvent-polarity-dependent rate yield. 24,42,45,74,77,82,109-118 The CT state couples strongly with the S<sub>1</sub> and <sup>1</sup>(TT) state via superexchange to serve as an intermediate state for the formation of <sup>1</sup>(TT) whether directly populated or only virtually involved. With the help of this CT-assisted mechanism, highly efficient iSF can still occur in weakly coupled systems. Besides the above-mentioned two mechanisms, several other new models have been established for iSF. For example, the 'vibronic coupling' or 'vibrationally coherent' mechanism has been proven in several systems, 73,119-121 corresponding to the case in which S<sub>1</sub> and <sup>1</sup>(TT) are nonadiabatically coupled through vibrational coordinates or form vibronically mixed states when their energy levels are resonant owing to excited vibrational quanta.73,122 This mechanism demonstrates an intimate link between the nuclear motions and SF, and can be approached from conical intersection, vibronically mixed  $S_1 - {}^{1}(TT)$ or symmetry-breaking vibrations.73,119-121,123,124 Recently, a quantum interferenceassisted mechanism was proposed for iSF.84 Constructive quantum interference can promote the formation of the <sup>1</sup>(TT) state; however, destructive quantum interference will slow

down the formation of <sup>1</sup>(TT). Besides the CT intermediate state, the excimer state is also observed to act as an intermediate for the formation of 1(TT).44 However, this mechanism is still controversial. In some covalent link oligomers, 23,100 the formation of the excimer state is considered to compete with SF. Thus, the real role of the excimer state in the SF process should be further investigated in the future. Additionally, a new spin exchange mechanism has been demonstrated to enable efficient SF via spin density localization,125 which is fundamentally different from the CT-mediated mechanism. In this mechanism, S<sub>1</sub> already has partial <sup>1</sup>(TT) characteristics and can quickly relax to <sup>1</sup>(TT).

# Covalently linked tetracene oligomers

Firstly, we discuss tetracene dimers, and then oligomers with more than two tetracene units. Dimers, as the smallest units capable of SF, can offer a better understanding of the SF mechanism. When the structural unit is increased, some extra factors including exciton delocalization and entropy increase should be considered.

#### 3.1 Tetracene dimers

In the past few years, several tetracene dimers with different conformations have been prepared. Depending on the connection position of tetracene, we classified these dimers into three groups, including 5,5'-linked dimers, 2,2'-linked dimers and other dimers linked via multiple-positions of tetracene (Fig. 2).

#### 3.1.1 5,5'-Linked tetracene dimers

3.1.1.1 Tetracene dimers bridged via 5-position of tetracene. In this part, initially we focus on tetracene dimers bridged through the 5-position of tetracene via a spacer (1-10, Fig. 2).

The pioneering work utilizing molecular dimers to study iSF was reported in 2006 by Bardeen's group, in which three tetracene dimers linked by para-phenylene (1 and 2) and meta-phenylene (3) groups were prepared. 68,126 The electronic coupling in these three dimers is dominated by through-bond interactions, which is very weak, as revealed by the small redshift in their absorption spectra compared to that of monomeric tetracene (Fig. 3a).

The presence of a delayed fluorescence signal in the fluorescence dynamics of dimers 1 and 2 indicates that these two dimers undergo SF (Fig. 3b). In contrast, SF is absent in dimer 3, which may be caused by its much smaller through-bond interactions than that in dimers 1 and 2. Notably, the SF efficiencies of dimers 1 and 2 are only  $\sim$ 3%, which is much lower than that of tetracene crystal films.127-131 This should be ascribed to their weak intramolecular electronic coupling in comparison with the strong coupling in the solid. Additionally, this study revealed that the fission process  $(S_1 \rightarrow {}^1(TT))$  is temperaturedependent in dimers 1 and 2, but the fusion process  $\binom{1}{TT} \rightarrow$  $S_1$ ) is independent of temperature (Fig. 3c). This is consistent with the fact that the fission process in tetracene derivatives is endothermic. Thus, this study provided some insights into the design of iSF molecules. Our group also synthesized a para-

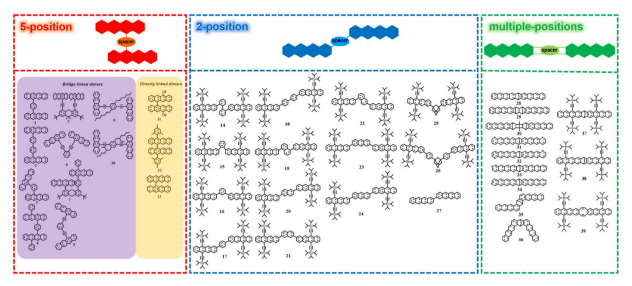


Fig. 2 Molecular structures of tetracene dimers linked via 5-position of tetracene, 2-position of tetracene and multiple-positions of tetracene.

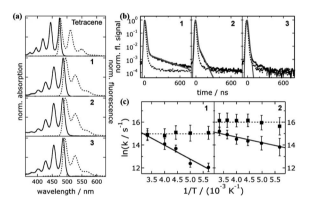


Fig. 3 (a) Absorption and fluorescence spectra of dimers 1-3 and tetracene. (b) Fluorescence dynamics of dimers 1-3. (c) Temperature dependence of fission (circle) and fusion (square) rates of dimers 1 and 2. Reproduced with permission. <sup>68</sup> Copyright 2007, the American Chemical Society.

phenylene-linked tetracene dimer (4) but with two extra phenyl groups on the two tetracene units, which improved the solubility of the dimer. However, its SF efficiency was still very low, which should also be caused by the weak intramolecular electronic coupling. These results suggest that phenylene-bridged tetracene dimers are not favorable for SF due to their weak through-bond electronic coupling.

To strength the intramolecular electronic coupling, a "face-to-face" stacked tetracene dimer linked by a xanthane bridge was prepared by our group (5).<sup>133</sup> Its absorption spectrum changed significantly compared to that of the corresponding monomer (Fig. 4a), suggesting the presence of strong electronic coupling between the two tetracene units. This strong coupling leads to the formation of an "excimer-like" state, as confirmed by its broad and structureless emission band.

However, the "excimer-like" state shows a short lifetime ( $\sim$ 0.23 ns), which is different from the traditional excimer-like state with a long lifetime. <sup>134–137</sup> This phenomenon is similar to

the observation in the excimer-mediated SF in a concentrated bis(triisopropylsilylethynyl)[TIPS]-tetracene solution, <sup>138</sup> suggesting the presence of SF in this tetracene dimer.

Unfortunately, we did not find direct evidence for SF in this dimer due to the use of single detection technology. Subsequently, Thompson's group investigated its excited-state dynamics in detail using femtosecond transient absorption (fs-TA) spectra and found that this dimer undergoes structural relaxation to form an excimer-like structure that decays into the  $^1$ (TT) state within 200 fs (Fig. 4b). $^{100}$  Nevertheless, the  $^1$ (TT) state cannot separate into the free triplet state due to its low energy and the strong coupling between the  $^1$ (TT) state and the  $S_0$  state, and decays directly into the  $S_0$  state quickly within 500 ps. It can be seen that the first step ( $S_1 \rightarrow ^1$ (TT)) of SF can be conducted fast in this class of strongly coupled tetracene dimers, but the second step ( $^1$ (TT)  $\rightarrow$  T<sub>1</sub> + T<sub>1</sub>) is impeded due to the strong exchange interaction between the two triplets of the  $^1$ (TT) state. $^{139}$  Specifically, to achieve the whole SF process efficiently,

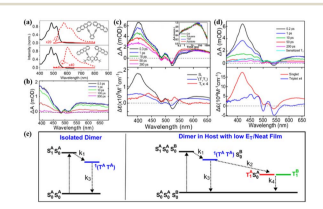


Fig. 4 (a) Absorption and fluorescence spectra of dimers 5 and 6. (b) fs-TA spectra of dimer 5 in solution. (c) and (d) fs-TA spectra of dimer 6 in solution (c) and solid film (d). (e) Kinetic model for dimer 6 in solution (left) and solid film (right). Reproduced with permission. Copyright 2016, the American Chemical Society.

the intramolecular electronic coupling should be decreased appropriately in molecular dimers.

Thus, a cofacial tetracene dimer (6) linked by ortho-phenylene with a twisted arrangement between tetracenes, exhibiting less overlap between the tetracene  $\pi$ -orbitals in comparison with dimer 5, was prepared by Thompson's group. 100 The electronic coupling between the two tetracene units in dimer 6 is still strong, but weaker than that in dimer 5, as can be seen from the smaller degree of the inversion of the 0-0 and 0-1 peaks (Fig. 4a). Similar to dimer 5, the fluorescence of dimer 6 was almost quenched completely in solution ( $\Phi_{\rm F} = 0.6\%$ ). However, the emission line shape retained the vibronic structure (Fig. 4a), suggesting that its emission does not originate from a deep-trap excimer state like that in dimer 5. The fs-TA experiment revealed that the <sup>1</sup>(TT) state is formed quickly in dimer 6 upon photoexcitation in solution with a time constant of  $\sim$ 2 ps (Fig. 4c). Similar to dimer 5, the <sup>1</sup>(TT) state could not further dissociate into free triplets in solution and decayed directly to the ground state with a time constant of  $\sim$ 500 ps (Fig. 4e). When dimer 6 was prepared as a solid film, the <sup>1</sup>(TT) state was also formed quickly ( $\tau = \sim 0.8$  ps) (Fig. 4d and e).

Notably, regardless of solution or solid film, the formation rate of the <sup>1</sup>(TT) state was much faster than that in crystalline tetracene, suggesting that the arrangement of the two tetracenes in dimer 6 is more favorable for SF than that in crystalline tetracene. More importantly, the <sup>1</sup>(TT) state can further separate into free triplets rapidly ( $\tau = \sim 0.2 \text{ ps}$ ) in the solid film *via* triplet energy transfer to the neighboring tetracene unit with a high triplet yield of 154%. This indicates that the triplet energy transfer is an essential second step required for the production of free triplets in endothermic SF systems. Additionally, to decrease the coupling strength, we synthesized a slip-stacked tetracene dimer composed of a tetracene unit and 5,12-diphenyltetracene (DPT) unit (7).140 In contrast to the "face-to-face" stacked structure of dimer 5, the slip-stacked structure of dimer 7 causes a large decrease in the degree of overlap between the two chromophores. Consequently, the electronic coupling between the two tetracene units in dimer 7 is almost negligible. The TA experiment revealed that ultrafast singlet energy transfer from the tetracene unit to the DPT unit proceeded with a time constant of  $\sim 0.32$  ps. This ultrafast singlet energy transfer successfully prevents the useless nonradioactive decays of the singlet state of tetracene and stores the excitation energy of tetracene as the singlet state of DPT. Unfortunately, no efficient SF was observed in this dimer, which may be caused by its weak intramolecular interaction.

Recently, Thompson's group prepared meta- and para-bis(ethynyl tetracenyl)benzene dimers (8–10), which are analogues to the above-mentioned tetracene dimer 6, to study the effect of through-bond versus through-space coupling on the overall SF process.99 For meta-bis(ethynyl tetracenyl)benzene dimer (8), the two tetracene units is cross-conjugated. The interchromophore interactions are dominated by through-bond coupling, but the interaction strength is very small, and thus it can be deduced from its nearly unchanged absorption spectrum compared to that of the corresponding monomer. Alternatively, ortho- and para-ethynylbenzene linkers enable

conjugation between the two tetracene units. Thus, the through-bond coupling between the two tetracene units in ortho- (6) and para-ethynylbenzene-linked dimers (9 and 10) is strong.

In the case of dimer 6, through-space coupling is also present besides through-bond coupling. Analogously to dimer 6, 100 dimers 9 and 10 are weakly emissive in solution ( $\Phi_{\rm F}$  < 1%), suggesting the presence of a nonradiative decay pathway in them. Instead, dimer 8 is highly emissive with a quantum yield of 62%. TA spectra revealed that ultrafast SF proceeded in dimers 9 and 10 ( $\tau_{SF} = 0.1$  and 0.4 ps for dimers 9 and 10, respectively) similar to that in dimer 6 ( $\tau_{SF} = 2$  ps) (Fig. 5b-d). However, no obvious triplet signal was observed in the TA spectra of dimer 8 (Fig. 5a), indicating the absence of efficient SF. This suggests that the conjugation linked mode is more favorable for SF than the cross-conjugation linked mode in endothermic SF systems. In contrast, a series of pentacene dimers linked by the same linker (ortho-, meta- and para-ethynylbenzene), where the energetics are favorable for SF, displayed ultrafast SF.24

Additionally, the faster SF rate of dimers 9 and 10 compared to that of dimer 6 indicates that the through-bond coupling operating in the para dimers is stronger than the combined through-space and through-bond coupling operating in the ortho dimer. Compared to the slow SF rate in tetracene dimer 1, in which the two tetracene units are directly attached to the para-phenyl group, dimers 9 and 10, where the two tetracene units are linked to the para-phenyl group via an ethynyl group, exhibited a much faster SF rate. This may be caused by the presence of two ethynyl groups, which can perturb the S<sub>1</sub> and T<sub>1</sub> state energies in a favorable way and ensure stronger electronic coupling between the two tetracene units.

Notably, the formed <sup>1</sup>(TT) state can further dissociate into two free triplets in the para dimers 9 and 10 instead of the ortho dimer 6, which is attributed to the rotational flexibility of the two tetracenes in dimers 9 and 10. This breaks the coupling between the two tetracenes and allows the triplets to separate from the <sup>1</sup>(TT) state. The rotation of the tetracene units in dimer 6 is hindered because they are confined to a cofacial orientation. This provides another method to facilitate the separation

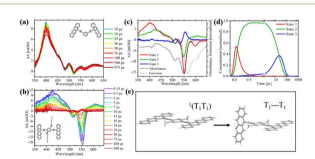


Fig. 5 (a) and (b) TA spectra of dimer 8 (a) and 9 (b) in solution. (c) and (d) Species associated difference spectra (c) and concentration curves (d) from the target analysis of the TA data of dimer 9. (e) Proposed decoupling of the <sup>1</sup>(TT) state in dimer 10 to give rise to two independent triplets. Reproduced with permission.99 Copyright 2018, the American Chemical Society.

of the  $^1$ (TT) state via the molecular rotation besides the above-mentioned triplet energy transfer. Although free triplets could be obtained in dimers **9** and **10** via the molecular rotation, the yield is relatively low ( $\Phi_{\rm T}=70\%$  and 94% for dimers **9** and **10**, respectively), which is ascribed to the fast recombination rate of the triplets (20 and 76 ps for dimer **9** and **10**) compared to the rate of their formation (19 and 48 ps for dimers **9** and **10**), respectively. This may be a disadvantage of molecular dimers, where the formed two triplets are confined in one molecule, and thus recombine rapidly.

Moreover, dimer 9 linked by hexyloxy-bis(ethynyl)benzene ( $\tau_{SF}=0.1$  ps) showed a faster SF rate compared to dimer 10 linked by 2-ethylhexylbis(ethynyl)benzene ( $\tau_{SF}=0.4$  ps), which is because the hexyloxy-bis(ethynyl)benzene linker has a much closer molecular orbital energy to that of tetracene, inducing better mixing of the linker orbital with tetracene, and thus stronger coupling in dimer 9. This highlights the important role of the linker energetics in SF for endothermic SF systems. Nevertheless, it is important to note that different types of substituents may increase different degrees of steric hindrance when linking substituents on phenylene spacer, thus limiting the conformational flexibility and blocking SF.  $^{101}$ 

3.1.1.2 Directly linked tetracene dimers via 5-position of tetracene. Here, we discuss the directly linked dimer via the 5-position of tetracene (11–13, Fig. 2), 118,141,142 in which the two tetracene units are linked directly through a C–C bond and interact via through-bond coupling.

Notably, the two tetracene units can rotate around this single bond, inducing the formation of the symmetry breaking charge transfer (SBCT) state, which shows a significant solvation effect, similar to that in single-bond-linked anthracene and pentacene dimers. <sup>114,115,144–146</sup> The presence of the charge transfer (CT) state in this type of dimer enables the role of the CT state in the SF process to be determined.

In 2016, a stable directly linked tetracene dimer (11) was synthesized by our group by introducing an electronwithdrawing cyano group in the tetracene skeleton.141 Its singlet state is delocalized on the two tetracene units and mixed strongly with the CT state. Unfortunately, we failed to observe the occurrence of SF. This gives the impression that the orthogonal conformation linked by a single bond is not sufficient to support the occurrence of SF. However, in 2019, this vague understanding was broken by Musser's group. 118 They found that activated iSF could occur in a single-bond-linked orthogonal tetracene dimer with two phenyl groups in the tetracene skeleton (12). As documented in anthracene and pentacene dimers with an orthogonal conformation, 114,115,144-146 its photophysical properties are strongly dependent on the torsional angle  $(\Phi)$  between the two tetracene units. The displacement along  $\Phi$  leads to mixing between the localized excited state (LE) and the CT state (Fig. 6a), reaching the maximum mixing at the local minima of the LE state at 70°. This introduces a strong solvation effect on its fluorescence (Fig. 6b). In low-polar solvents, the emission is dominated by a well-defined vibronic progression, which is attributed to the relaxed LE state. With an increase in the solvent polarity, the emission becomes featureless and gradually shows a redshift,

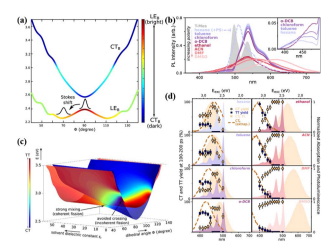


Fig. 6 (a) Potential energy surfaces of the LE<sub>B</sub> and CT<sub>B</sub> states along the torsional coordinate ( $\Phi$ ) for dimer 12. (b) Steady-state photoluminescence of dimer 12 in a range of solvents (solid lines). (c) Adiabatic potential energy surfaces along the dihedral angle  $\Phi$  and the solvent dielectric constant, showing their CT (blue) and TT (red) character. (d) Yield of long-lived CT (circles) and TT pairs (squares), extracted from excitation-dependent transient absorption measurements for solvents of increasing polarity (left to right). Reproduced with permission. <sup>143</sup> Copyright 2019, the American Chemical Society.

which indicates the participation of the CT state. 115,144 The transient absorption spectra showed that activated iSF could proceed with the excitation energy above the threshold energy (2.8 eV). Notably, iSF is also strongly dependent on the solvent polarity (Fig. 6d). In nonpolar solvents, the <sup>1</sup>(TT) yield is nearly constant above the threshold energy, but the <sup>1</sup>(TT) yield shows an obvious gradual onset in polar solvents. Moreover, in intermediate-polarity solvents iSF has the largest <sup>1</sup>(TT) yield and the fastest rate. This iSF behaviour was attributed to the large mixing of the CT state and the 1(TT) state in intermediatepolarity solvents, which makes SF coherent and accelerates the SF rate (Fig. 6c). 143 In low-polarity and high-polarity solvents, the mixing is minimal and makes SF incoherent. This result deeply reveals the role of the CT state in the SF process, in which the hot iSF can be mediated by the CT state via the coherent or incoherent mechanism and these two mechanisms can be switched by the solvent polarity by controlling the mixing degree between the CT state and the <sup>1</sup>(TT) state. Additionally, Shizu et al. predicted that the <sup>1</sup>(TT) state was formed efficiently in a directly linked tetracene dimer (13) due to the small <sup>1</sup>(S<sub>0</sub>S<sub>1</sub>) - <sup>1</sup>(TT) energy difference and the large vibronic coupling at the  $^{1}(S_{0}S_{1})$  geometry.  $^{142}$ 

Recently, Mattos *et al.* further studied the role of the SBCT state in SF through a computational study of dimer 13.<sup>82</sup> They quantified the SF mechanism proposed by Alvertis *et al.*,<sup>118</sup> in which solvent-induced symmetry breaking leads to a high-energy CT state that interacts with the <sup>1</sup>(TT) state, resulting in SF. An approximate assessment of the nonadiabatic interactions between the different electronic states highlighted that the CT state is essential in facilitating the transition from the bright excitonic state to the <sup>1</sup>(TT) state, leading to SF. Additionally, several types of symmetry-breaking inter- and intra-

fragment vibrations play a crucial role in a concerted mechanism with the solvent environment and with the symmetric interfragment torsion, which tune the admixture of excitonic and CT states. This work offers a new perspective on how solvent-induced SBCT can be understood and how it cooperates with the intramolecular mechanisms in SF. The result of the three latter works suggests that the orthogonal tetracene dimeric structure linked directly by the single-bond is advantageous for achieving efficient iSF, and the CT state here is generally considered to be an intermediate state for the occurrence of SF.

3.1.2 2,2'-Linked tetracene dimers. The 2-position is another active site of tetracene. Tetracene dimers linked via the 2-position will show different electronic couplings in comparison with that linked via the 5-position. This will influence the SF dynamics. 103 Thus, several tetracene dimers linked via the 2various linkers position with have been reported (Fig. 2).75,98,147,148

Although efficient iSF has been achieved in the abovementioned tetracene dimers linked via the 5-position of tetracene, the lifetime of the obtained triplet is very short. Thus, the short-lived triplets are difficult to be harvested in solar energy conversion. Consequently, realizing the long-lived triplet state is necessary in tetracene derivatives for the application of SF. The 2,2'-biphenyl linker has been employed to construct pentacene dimers for achieving a high yield of long-lived triplets.149 Inspired by this, Nakamura et al. prepared a 2,2'-biphenyllinked tetracene dimer (14, Fig. 2) with a bent-shaped structure.75 The steady-state absorption and fluorescence spectra demonstrated that the electronic coupling between the two tetracene units is relatively weak in this dimer.

The fs-TA spectra showed that the triplet signal appears rapidly with a time constant of  $\sim$ 14 ps (Fig. 7A). Then, the triplet state decays with a biexponential function (Fig. 7B). The faster component originates from the mixture of the singlet and <sup>1</sup>(TT) states because of the competitive reaction between iSF and triplet-triplet annihilation (TTA). The other much slower one is attributed to the free triplet state, 150 which could be confirmed by its ultra-long lifetime (0.29 ms) (Fig. 7C). The quantum yield of the free triplet state is up to  $\sim$ 175% in this dimer, which should be attributed to the weak interchromophore electronic

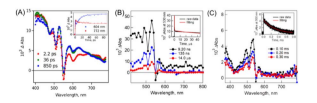


Fig. 7 (A) fs-TA spectra of dimer 14 in PhCN (excited at 350 nm). The inset shows the time profiles at 604 nm (blue) and 772 nm (red). (B) ns-TA spectra of dimer 14 in PhCN (excited at 550 nm). The inset shows the time profile at 530 nm (black) and fitting curve at 530 nm (red). (C) μs-TA spectra of dimer 14 in PhCN (excited at 532 nm). The inset shows the time profile at 500 nm (black) and fitting curve at 500 nm (red). Reproduced with permission.<sup>75</sup> Copyright 2019, the American Chemical Society.

coupling. However, the reason for the fast SF rate in this dimer with very weak electronic coupling is still unclear and should be further investigated in the future.

This is useful for the design of novel iSF materials with a high yield of long-lived triplet states. Besides electronic coupling, the conformational flexibility of the dimer structure may play an important role in SF.99 Thus, Nakamura et al. studied the synergistic role of conformational flexibility and electronic coupling in SF utilizing four tetracene dimers (15-18) linked by different phenylene-based linkers.98 The steadystate spectra and electrochemical measurements revealed that the intramolecular electronic coupling of dimer 16 linked by the meta-phenyl group is relatively weaker than the ortho-(dimer 15) and para- (dimer 17) phenyl-linked dimers. The electronic coupling of dimer 18 bridged by the 4,4'-biphenyl group should be somehow weaker than that of the other three dimers. The energies of the rotational barriers of these four dimers were calculated using DFT to discuss the conformational flexibility between the two tetracene units. The rotational barriers of dimers 18, 16 and 17 are 0.075, 0.13, and 0.081 eV, respectively. In contrast, the barrier of dimer 15 could not be determined because of the bulkiness of the TIPS acetylene unit. This suggests that dimers 17 and 18 possess larger conformational flexibilities than dimer 15 and 16. Thus, these four dimers are categorized into four different characters, i.e., dimer 15 with strong electronic coupling and low conformational flexibility, dimer 16 with weak electronic coupling and low conformational flexibility, dimer 17 with strong electronic coupling and high conformational flexibility, and dimer 18 with weak electronic coupling and low conformational flexibility. The TA experiment revealed that SF could proceed in all four dimers (Fig. 8a-h). However, the formation and separation rate of the <sup>1</sup>(TT) state and the corresponding inverse process (TTA) are significantly dependent on the electronic coupling strength and the conformational flexibility degree. Dimers 15 and 17 with strong electronic coupling have a faster <sup>1</sup>(TT) formation and TTA rate than dimer 16 and 18 with weak electronic coupling, suggesting that the larger electronic coupling will induce faster SF and TTA rate. Although the strong electronic coupling could enable a faster SF rate, it also accelerated the TTA process, which will decrease the overall triplet yield. Thus, the electronic coupling strength should be balanced to ensure a fast SF rate and slow TTA rate for achieving a high triplet yield. Notably, dimer 18 with the largest conformational flexibility has the fastest <sup>1</sup>(TT) separation rate among the four dimers, suggesting that high conformational flexibility will accelerate the dissociation of the <sup>1</sup>(TT) state. Additionally, the quintet (<sup>5</sup>(TT)) intermediate state for the dissociation of the <sup>1</sup>(TT) state into the free triplet state was observed in the TREPR spectra of dimers 15, 17 and 18 (Fig. 8i). Different from dimers 15 and 17, the weakly coupled triplet state (T + T) also appeared in the TREPR spectra of dimer 18. This is caused by its flexible conformation, as confirmed by the calculated conformation of the  ${}^{5}(TT)$  and (T + T) states (Fig. 8j). The  ${}^{5}(TT)$ state possesses a nearly planer conformation with a strong spin-spin exchange interaction (J) induced by the large direct

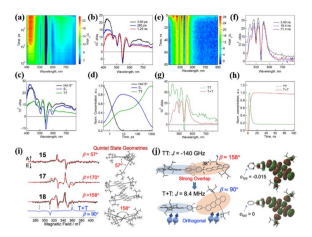


Fig. 8 (a) and (b) fs-TA spectra of dimer 18 in THF ( $\lambda_{ex} = 350$  nm). (c) and (d) Species-associated TA spectra and corresponding decay profile of dimer 18. (e) and (f) ps-TA spectra of dimer 18 in THF ( $\lambda_{ex}$  = 350 nm). (g) and (h) Species-associated TA spectra and corresponding decay profile of dimer 18. (i) TREPR spectra at 0.4 µs after the laser irradiation of dimers 15, 17 and 18 in MTHF at 77 K. Simulated spectra of the strongly coupled quintet multiexcitons are shown by the red lines with the dihedral angles  $(\beta)$  between the triplet excitons, as indicated by the right geometries. The spin-correlated triplet pair (SCTP) spectrum of the weakly coupled T + T state for dimer 18 was also computed by applying the electron spin polarization transfer (ESPT) model with J = 8.4 MHz, as shown by the blue line, and added to the quintet spectrum to obtain the red line. (j) Geometries of the <sup>5</sup>(TT) and T + T multiexcitons in dimer 18 from the simulations of the TREPR spectra (left) and the corresponding electron density distribution (right). Dihedral angle change between the adjacent aromatic planes is indicated by the arrows at the centers, denoting that an almost orthogonal conformation leads to the decoupling in J. Reproduced with permission.98 Copyright 2021, the American Chemical Society.

orbital overlap. However, an orthogonal conformation is adopted in the T<sub>1</sub> state, and consequently the two triplets are no longer correlated.

Specifically, intramolecular decoupling occurs in the <sup>1</sup>(TT) state of dimer 18 by the conformation change, and thus leads to the highest free triplet yield (196%) among the dimers (67%, 125% and 117% for dimers 15, 16 and 17, respectively). This also highlights the important role of molecular conformation change for the separation of the <sup>1</sup>(TT) state, similar to the report on dimers 9 and 10 by Thompson's group.99 This work provides a new perspective for the development of novel iSF materials with long-lived triplets.

Campos' group reported the preparation of a series of tetracene dimers (16, 17, and 19-24, Fig. 2) linked by a bridge with constructive (CQI) or destructive (DQI) quantum interference effect to reveal the effect of quantum interference (QI) on the formation dynamics of the <sup>1</sup>(TT) state (Fig. 9a).<sup>84</sup> The CQI bridges include 14Ph (17), 15N (19), 26N (20) and 26A (24), while 13Ph (16), 27N (21), 16N (22) and 27A (23) show the DQI effect (Fig. 9a). As revealed by the TA experiment, the formation rate of the <sup>1</sup>(TT) state in the dimers with the DQI effect is always slower than that with the CQI effect across different bridge lengths and geometries (Fig. 9b), emphasizing the important role of CQI in SF.

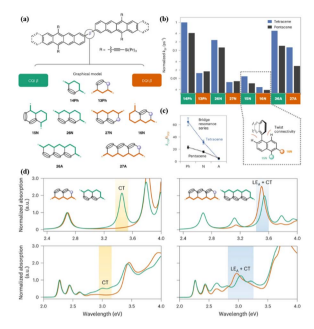


Fig. 9 (a) Bridged tetracene and pentacene dimers (top), where  $\beta=$ 14Ph (17), 13Ph (16), 15N (19), 26N (20), 27N (21), 16N (22), 26A (23) and 27A (24). (b) Comparison of SF rates (normalized to the largest  $k_{SF}$ ) across all Ph, N and A bridges in this work. Inset: chromophores attached to the 1 and/or 5 positions and the 2, 6 or 7 positions of naphthalene. (c) Changes in the CQI and DQI SF rate constants as the bridge length increases. (d) Calculated (top) and experimental (bottom) normalized absorption spectra for naphthalene-bridged tetracene dimer (left) and anthracene-bridged tetracene dimer (right). Green traces correspond to a CQI bridge (26N or 26A) and orange traces correspond to a DQI bridge (27N or 27A). The calculations assume a planar conformation. The inclusion of intermonomer rotation decreases the intensity of the CT absorption, but the absorption energy is largely unaffected. Reproduced with permission.84 Copyright 2022, Nature.

This phenomenon may be due to the more pronounced CT character in the dimers with the CQI effect compared to that with the DQI effect, as shown in the 26N- and 27N-bridged dimers (20 and 21), respectively (Fig. 9d). Notably, although the 26A- and 27A-bridged dimers show a similar CT strength, the <sup>1</sup>(TT) formation rate of the 27A dimer is much slower than that of the 26A dimer (Fig. 9d), which is caused by the near-zero transition dipole moment between the ground state and the CT state in the 27A dimer. Thus, strong dipole coupling should also be considered for SF. As the bridge length increased, the difference between the <sup>1</sup>(TT) formation rate constants of the CQI and DQI dimers became smaller (Fig. 9c). This is mainly due to the larger resonance interactions between tetracene and the longer bridge caused by the much closer molecular orbital energy of the bridge to tetracene, which can accelerate SF. This effect will compensate the DQI effect to some extent. Additionally, the chromophore attached to the 1 and/or 5 positions of naphthalene (dimer 19) has a larger chromophore distortion than the chromophore attached to the 2, 6 or 7 positions (dimer 22), which will reduce the orbital overlap of the chromophores and slow down the SF. However, the dimer attached to the 1 and/or 5 positions of naphthalene has a larger SF rate compared

to that attached to the 2, 6 or 7 positions. This suggests that QI remains a substantial effect in the limit of both strong and weak interchromophore coupling. This work reveals critical considerations of the bridge topology and frontier molecular orbital energies in applying QI conductance principles to predict the rates of multiexciton generation.

The above-mentioned tetracene dimers are almost covalently bonded via conjugated or cross-conjugated linkages. Generally, these linkers result in a decrease in the S<sub>1</sub> and T<sub>1</sub> energies, which makes SF unfavorable thermodynamically. 151 Thus, nonconjugated linkages including sp<sup>3</sup>-hybridized atoms may be a good choice for maintaining the intrinsic high energies of S<sub>1</sub> and T<sub>1</sub> and enable the moderate interactions needed to trigger iSF. Thus, Matsui et al. prepared two tetracene dimers (25 and 26) linked by an adamantane moiety. 147 The two tetracene units in dimer 25 are bonded to the adamantane moiety directly, but in dimer 26, the two tetracene units are bonded to the adamantane moiety via the phenyl group. According to their steadystate spectra (Fig. 10a), their S<sub>1</sub> and T<sub>1</sub> state energies were estimated to be 2.27 and 1.02 eV, respectively, suggesting that the SF in these two dimers is exergonic. Notably, the T<sub>1</sub> energy is stabilized a lot compared to the pure tetracene (1.2 eV), which is due to the presence of silylethynyl groups. The decreased T<sub>1</sub> energy is contradictory to the value of the T<sub>1</sub> energy (1.2 eV) for the same tetracene monomer with two substituted silylethynyl groups reported by Anthony et al. 138 In anthracene derivatives, the introduction of two silylethynyl groups could indeed disturb the S<sub>1</sub> and T<sub>1</sub> energies and make SF more feasible. 19,152 The role of the silylethynyl group in the decrease of the T1 energy should further investigated. The presence of the delayed

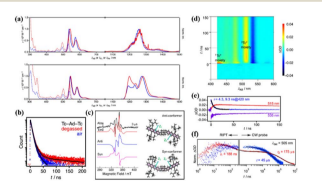


Fig. 10 (a) UV-vis absorption (dotted), fluorescence, and phosphorescence (solid) spectra of dimers 25 and 26 (red) and the corresponding monomer (blue) in MCH. Phosphorescence spectra were measured using methylcyclohexane (MCH) matrices at 77 K. (b) Fluorescence decay profiles of dimer 25 in degassed (red) and aerated (blue) MCH detected at the respective  $\lambda_{FL}$  maxima. Black curves are fitted lines. (c) Time-resolved EPR spectra of dimer 25 in degassed MTHF at 110 K (left). (d) TA spectra of dimer 25 in degassed MCH. (e) Single-wavelength dynamics probed at 420, 515 and 535 nm. (f) Nanoto-millisecond decay profiles for (red) dimer 25 and (blue) corresponding monomer traced at 505 nm using RIPT and CW probe methods for the region shorter and longer than 200 ns, respectively.  $\Delta$ OD values were normalized at 200 ns. Two components of the decay profile possess time constants of 188 ns and 175 µs for dimer 25. Reproduced with permission.<sup>147</sup> Copyright 2019, the American Chemical Society.

fluorescence (Fig. 10b) and the quintet state signal in timeresolved EPR spectra (Fig. 10c) of dimer 25 suggest that iSF occurs in dimer 25. In contrast, no SF signal was observed in dimer 26 due to the extremely weak electronic coupling induced by the large inter-chromophore distance.

The presence of iSF in dimer 25 was also confirmed by the TA experiment (Fig. 10d-f). The triplet state decayed with a twoexponential function, i.e., 188 ns and 175 µs. The fast component is ascribed to the TTA process, and the long-lived component originated from the free triplet state. Notably, two different conformers were presented in the quintet state of dimer 25 (Fig. 10c), suggesting that the quintet state is populated by the spin relaxation of the <sup>1</sup>(TT) state induced by the conformation change.153 The triplet yield of dimer 25 is still relatively high (92%), resulting from the slow TTA and the fast separation of the <sup>1</sup>(TT) state, which is a consequence of the weak electronic coupling. As can be seen, the incorporation of rigid aliphatic linkages in dimers provides a way to control the 3D orientations of  $\pi$ -conjugated systems, while leaving the intrinsic S<sub>1</sub> and T<sub>1</sub> levels of the original monomers, causing exergonic iSF to take place and preventing the undesirable TTA.

Additionally, a directly linked tetracene dimer (27) via the 2position of tetracene was reported by Müller et al. and the photophysical property of its solid state was investigated in detail.148 However, no SF signal was observed in its solid film, which is because the large decrease in the S<sub>1</sub> state energy caused the energy requirement of SF to no longer be fulfilled.

3.1.3 Dimers linked *via* multiple-positions of tetracene. All the above-mentioned tetracene dimers have in common the property that they are linked by a single bond, and thus the tetracene unit can be rotated around the linkage, making their conformation more flexible. Recently, a series of tetracene dimers connected by multiple-positions of tetracene with rigid linkages was reported, which exhibited completely different photophysical properties from flexible dimers. This enabled other factors influencing SF besides the conformation change to be evaluated. Damrauer's group prepared a wide range of norbornyl-bridged tetracene dimers and studied experimentally and theoretically their photophysical in detail (28-38, Fig. 2).  $^{123,154-160}$  This class of compounds is characterized by  $\mathrm{C}_{2\mathrm{V}}$ point-group. Due to the orbital symmetry, the adiabatic coupling from S<sub>1</sub> to <sup>1</sup>(TT) is general zero, resulting in the blocking of SF.

This can be broken by vibration, and an effective conversion from S<sub>1</sub> to <sup>1</sup>(TT) can be achieved as long as there is moderate nonadiabatic coupling. 123,156,157 Thus, the bridge geometry, including the length of the bridge and the orientation of the bridge, was used to control the SF thermodynamics and the electronic coupling between the chromophores in these dimers.154

Firstly, the effect of bridge length on the SF driving force and electronic coupling was investigated theoretically using three dimers (28-30).154 With the elongation of the bridge length from dimer 28 to 30, the interchromophore electronic coupling becomes weak gradually. However, the SF driving force does not change with the length of the bridge. Despite the sizeable electronic coupling and reasonable CT energy in these three dimers, the SF rate is expected to be nonoptimal due to the nonhorizontal CT matrix elements being zero as a result of the  $C_{2V}$  symmetry. This was also confirmed by the experimental results, where dimer 28 showed slow iSF with a low triplet yield  $(\sim 6\%)$ .<sup>158</sup> Thus, vibrational motions that break the long-axis reflection plane will be required.

Based on their vibronic coupling predictions, 159 the norbornyl bridge was attached at the 1,2- instead of the 2,3-positions of tetracene to construct two isomers (trans-dimer (35) and cis-dimer (36)) and break the  $C_{2V}$  symmetry. <sup>156</sup> A more favorable SF driving force was achieved for dimers 35 and 36 compared to dimers 28-30, which can be understood as manifesting from the enhanced wave function delocalization inclusive of the bridge in dimers 35 and 36, serving to lower both the T<sub>1</sub> and S<sub>1</sub> energies. Notably, in these two dimers, it appears possible to increase the interchromophore electronic coupling (manifested by the larger Davydov splitting), while simultaneously increasing the SF driving force. Furthermore, the coupling strength of cis-dimer 36 is considered to be greater than that of the *trans*-one due to three reasons. Firstly, the transition dipole moments of the two tetracene units in the cis-dimer are very close, which enhances the Coulomb interaction. Secondly, the two cis-unit edges provide a larger steric contribution to the exchange interaction. Thirdly, given that the direct throughspace orbital interaction occurs not only in the region of the bridge but also inside each tetracene arm, the cis-dimer through electron transfer and hole transfer electron coupling is stronger. 154 Besides, the CT energy in the cis-dimer is significantly lower than that in the trans-dimer, and all these factors are favorable to improve the SF efficiency in the cis-dimer. 156 In addition, heteroatom-substituted tetracene dimers (33 and 34) were also used to break the  $C_{2V}$  symmetry. <sup>156</sup> As expected, the electronic coupling is asymmetric. In these dimers, the "horizontal" coupling shows a common sign given that both chromophores in each dimer are juxtaposed directly across from each other via the norbornyl bridge, resulting in a fixed value. This means that SF will occur via the constructive interference of electron transfer and hole transfer pathways in the C2symmetric species. This constructive interference increases the diabatic coupling significantly and leads to faster SF.

Additionally, the effect of the substituent position on iSF was investigated utilizing two isomers (37 and 38), where the acetylene substitution pattern on each chromophore in dimer 38 is moved outward by a ring relative to the bridge (Fig. 11a). 123,155,157 The steady-state absorption and fluorescence spectra showed that there is nearly no difference between these two dimers (Fig. 11b), suggesting that their S<sub>1</sub> state energy is almost identical. The TA experiment revealed that ultrafast iSF occurs in these two dimers ( $\tau_{\rm SF}=10$  ps and 5 ps for dimer 37 and 38) (Fig. 11c and e), respectively. The fast iSF rate is initially surprising given the absence of a reaction driving force and the unfavorable structural symmetry in these two dimers, which is expected to limit the diabatic coupling between the reactant and product.154,159 However, their analysis suggests that the fewparameter rate constant expression of the Marcus theory explains both individual and comparative findings in the set of systems, thus establishing benchmarks for the diabatic

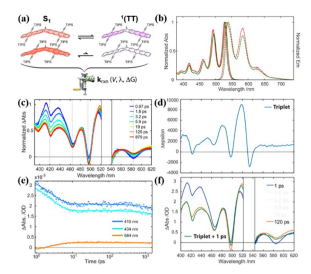


Fig. 11 (a) Illustration of electronic coupling strength in the  $S_1$  and  $^1(TT)$  states for dimers 37 and 38. (b) Normalized steady-state electronic absorption (solid) and emission (dashed) spectra for dimers 37 (red) and 38 (green) in toluene. (c) TA spectra of dimer 38 in toluene following ultrafast excitation at 530 nm. The spectral region around the excitation wavelength is removed due to pump scatter. (d) Triplet  $\Delta_{\mathcal{E}}$  spectrum for dimer 38 from sensitization experiment in toluene. (e) Selected single wavelength kinetic traces (data points) taken from the full-spectrum data with applied model fits (lines) retrieved from global analysis. (f) Selected spectral slices for dimer 38 at 1 ps (blue) and 120 ps (red) along with a reconstructed TA spectrum (green), which is comprised of a superposition between the 1 ps TA spectrum and the sensitized triplet  $\Delta_{\mathcal{E}}$  spectrum from (d). Reproduced with permission. Copyright 2019, the American Chemical Society.

coupling and reorganization energy needed for efficient <sup>1</sup>(TT) formation. 159,161,162 This highlights the importance of diabatic coupling and reorganization energy for SF with the Marcus theory for the first time. Due to the close energy alignment of the  $S_1$  and  $^1(TT)$  state, fast equilibrium between them is established in these two dimers, as confirmed by the perfect match between the TA spectrum at 120 ps and TA spectrum obtained via the superposition of the  $S_1$  and  $T_1$  states (Fig. 11d and f). This is also consistent with their relatively high fluorescence quantum yields ( $\sim$ 72%). The equilibrium constant (K) was calculated to be 0.1 and 1 for dimers 38 and 38, respectively. According to the K value, the  $^{1}(TT)$  yields of dimers 37 and 38 were estimated to be  $\sim$ 10% and  $\sim$ 50%, respectively. These markedly different <sup>1</sup>(TT) yields are caused by the different exchange interactions induced by the different placement of the acetylene side groups in these two dimers.97 In dimer 38, the relative placement of the acetylene side groups draws the triplet excitons further away from each another, thereby lowering the overall energy of the 1(TT) state and enabling its substantial participation ( $K \sim 1$ ) in equilibrium with the S<sub>1</sub> singlet exciton state. This suggests that the magnitude of the exchange interaction in the correlated triplet manifold plays a critical role in dictating the <sup>1</sup>(TT) yield in tetracene systems. However, no free triplet state has been found in these dimers.

Yamakado *et al.* also reported a dimer (39) linked *via* multiple-positions of tetracene in which the two tetracene units

are connected by an  $8\pi$  cyclooctatetraene (COT) ring. This linkage is different from the freedom rotation of a single bond73,103 and the fixed effect of the rigid spacer,106,157,158,160 and it is characterized by flapping motion (Fig. 12a). The electronic coupling between the two tetracene units of this dimer is relatively weak. This dimer exhibits  $C_{2V}$  symmetry at its equilibrium geometries (Fig. 12d), whereas the electronic coupling between  $S_1$  and  ${}^1(TT)$  is zero.  ${}^{143,154,158,159}$  Thus, there should be no SF in this dimer. However, efficient iSF is observed in it, as revealed by TA spectra (Fig. 12c), which is attributed to the flexible COT structure. This structure can promote SF by inducing conformational flexibility to cause symmetry-breaking vibrational modes (Fig. 12d). Different from the COT-linked anthracene dimer, the conformational planar dynamics is absent in this dimer because the energy potential no longer gives a minimum in its planar conformation as a consequence of the decreased electronic contribution of the  $8\pi$  COT ring with the extension of the acene unit. 163 This work shows a new platform for the design of iSF materials utilizing the fused COT linkage with conformation flexibility, and again highlights the importance to break the molecular symmetry for activating SF.

Next, we summarize the SF parameters of these dimers in Table 1. Among the 5,5'-linked dimers, dimers 1-4 bridged by para-phenylene or meta-phenylene have weak electronic coupling between the two tetracene units and show a low SF efficiency. When a triple bond is inserted between the tetracene unit and para-phenylene or meta-phenylene (dimers 6, 9 and 10), the electronic coupling is enhanced significantly due to the free rotation of the two tetracene units along the triple bond, which accelerates the SF dramatically because the SF rate is proportional to the square of the effective electronic coupling.<sup>164</sup> This indicates that relatively strong electronic coupling is

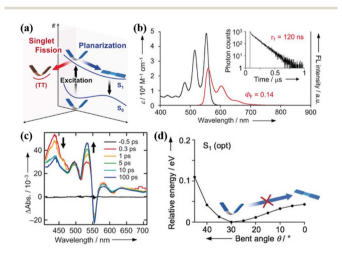


Fig. 12 (a) Energy diagram of the COT-fused dimers. Two possible dynamics, conformational planarization in  $S_1$  and SF to a correlated triplet pair <sup>1</sup>(TT), are indicated. (b) UV/vis absorption (black) and FL (red) spectra of dimer **39**. Fluorescence quantum yield ( $\Phi_{\rm F}$ ) is considered in the area of the fluorescence spectra. The fluorescence decay curve of dimer 39 is shown in the inset of (b). (c) TA spectra of dimer 39 in CH<sub>2</sub>Cl<sub>2</sub>. (d) S<sub>1</sub> energy profiles with respect to conformational planarization of dimer 39. Reproduced with permission. 163 Copyright 2018, Wiley-VCH.

necessary for efficient SF. However, relatively strong electronic coupling will hinder the dissociation of the <sup>1</sup>(TT) state and result in fast decay of the 1(TT) state. Furthermore, too strong coupling will lead to the formation of an excimer state, which competes strongly with the occurrence of SF, similar to dimer 5. Compared to 5,5'-linked dimers, 2,2'-linked dimers are more conducive for the occurrence of SF and achieving a high yield of long-lived free triplet states, which can be attributed to the proper electronic coupling and the larger conformational flexibility in 2,2'-linked dimers. 75,98,147 In these dimers, the electronic coupling is not only sufficient to ensure fast 1(TT) formation, but also weak enough to guarantee the 1(TT) dissociation into the free triplet state. More importantly, the strong conformational flexibility can also decrease the intertriplet spin-spin exchange interaction (*J*), and then decelerate the TTA process and accelerate the dissociation of the <sup>1</sup>(TT) state into the free triplet state, resulting in a high yield of free triplet states. In comparison with 5,5'-linked and 2,2'-linked dimers, most of the reported multiple-positions linked dimers show no SF, which is due to their orbital symmetry, making the adiabatic coupling from  $S_1$  to  ${}^{1}(TT)$  is zero, and then blocking SF. Only when the symmetry is broken, can SF occur. In this case, the symmetry can be broken by vibration. For example, in dimer 39, the conformationally flexible cyclooctatetraene linkage could break the molecular symmetry to activate SF. This highlights the important role of the vibration on SF. Through a systematic comparison of these dimers, we conclude that 2,2'-linked dimers are more favourable for the occurrence of SF in the tetracene chromophore.

#### 3.2 Tetracene oligomers with more than two tetracene units

In molecular dimers, the two triplets formed via SF are confined to the two subunits and easily annihilated by each other (TTA) to the ground state rapidly due to the strong electronic coupling. 103,106,166 This, together with the strong spin-exchange interaction between the two correlated triplets, may slow down the spin dephasing of the correlated triplet state. 167,168 Thus, the separation of the two correlated triplets into free triplets is difficult in molecular dimers. In endothermic SF systems, such as tetracene, the recombination of the two correlated triplets into the S<sub>1</sub> state can also proceed. However, the free triplets with long lifetime are more convenient to be harvested in solar energy conversion process. Consequently, achieving the separation of the two correlated triplets to yield free triplets is urgent for the application of SF, especially in endothermic SF systems. In crystalline tetracene and pentacene films, 92,169-171 the two correlated triplets can diffuse apart from each other via triplet energy transfer to the adjacent molecule at the  $S_0$  state, resulting in the generation of free triplets. Thus, applying this concept to iSF materials may also promote the dissociation of the two correlated triplets into two free triplets.

To verify this hypothesis, our group synthesized a series of tetracene oligomers including trimer (40, Fig. 13) and tetramer (41, Fig. 13), with dimer 4 as a comparison. 132,172,173 The electronic coupling in all these oligomers is relatively weak, as revealed by the small redshift in their absorption peak

Table 1 SF parameters of the discussed dimers

Connection position	Dimer	$ au_{\mathrm{SF/ps}}$	$\Phi_{ m TT}/\%$	$\tau_{\rm TT}/{\rm ns}$	$\Phi_{\mathrm{T}\cdots\mathrm{T}/\mathrm{T_{1}}}$ /%	$\tau_{T\cdots T/T_1}/\mu s$	Solvent	Ref.
5-Position	1	$3.6 \times 10^5$	2-3	_	_	_	Toluene	68
	2	$2.5  imes 10^5$	2-3	_	_	_	Toluene	68
	3	_	_	_	_	_	Toluene	68
	4	$1.23\times10^4$	_	_	21	10.9	Toluene	132
	5	_	0	_	_	_	_	100 and 133
	6	2	_	0.5	_	_	THF	165
		1	_	$5  imes 10^{-3}$	154	$4\times 10^{-4}$	Film	100 and 139
	7	_	_	_	_		_	140
	8	$1.5\times 10^4$	1	_	_	_	$CH_2Cl_2$	99
	9	0.1	_	$1.9\times10^{-2}$	94%	$2\times 10^{-5}$	$CH_2Cl_2$	99
	10	0.4	_	$4.8\times10^{-2}$	70%	$7.6 \times 10^{-5}$	$CH_2Cl_2$	99
	11	_	_	_	_	_	_	141
	12	3-40	_	_	_	_	_	118
	13	_	$\sim 10$	_	_	_	_	142
2-Position	14	14.1	_	_	$175\pm5$	294	PhCN	75
	15	14.3	_	33.3	$67 \pm 4$	222	THF	98
	16	90.9	_	12.5	$125\pm 8$	277.8	THF	98
		453	192	129	_	145	Toluene	84
	17	23.3	_	7.7	$117\pm 8$	294	THF	98
		7	199	121	_	141	Toluene	84
	18	400	_	2.9	$196\pm12$	333.3	THF	98
	19	30	199	140	_	136	Toluene	84
	20	952	185	129	_	145	Toluene	84
	21	591	190	141	_	170	Toluene	84
	22	1437	178	23	_	158	Toluene	84
	23	55	199	126	_	155	Toluene	84
	24	14	199	157	_	133	Toluene	84
	25	$6.6 \times 10^{3}$	126	13.5	92	175.4	Methylcyclohexane	147
	26	_	0	_	_	_	Methylcyclohexane	147
	27	_	_	_	_	_	_	148
Multiple-positions	28	$7.1 \times 10^4$	6.3	_	_	_	_	158
	29	_	_	_	_	_	_	154
	30	_	_	_	_	_	_	154
	31	_	_	_	_	_	_	156
	32	_	_	_	_	_	_	156
	33	_	_	_	_	_	_	156
	34	_	_	_	_	_	_	156
	35	_		_	_	_	_	154 and 156
	36	_	_	_	_		_	154 and 156
	37	9.1	<0.1	_	_	_	Toluene	157
	38	$5\pm0.5$	$50 \pm 8$	_	_		Toluene	123
	39	3	180	_	_	_	$CH_2Cl_2$	163

compared to that of the corresponding monomer. However, the degree of redshift from the dimer to tetramer gradually became large (Fig. 14a), suggesting that the corresponding electronic coupling gradually becomes stronger. This may be attributed to the increased planarity between/among the tetracene subunits in this series of molecules. 132,174 Their fluorescence spectra showed a similar vibrational structure (Fig. 14b), but a gradually decreased fluorescence quantum yield and a gradually accelerated fluorescence decay rate were observed on going from the dimer to tetramer, indicating the presence of an accelerated nonradiative decay pathway in these oligomers (Fig. 14c). 175 In the fs-TA spectra, the triplet state signal emerged at ~490 nm with a decrease in the S<sub>1</sub> absorption intensity in these oligomers (Fig. 14e), which originated from iSF. The triplet formation rate gradually became faster from dimer to tetramer, suggesting accelerated iSF (Fig. 14d). More importantly, the triplet state

showed a long lifetime (>100  $\mu$ s) (Fig. 14f), which suggests that the free triplet state is formed in these oligomers. A gradually enhanced free triplet yield was obtained in these oligomers (21%, 96% and 128% for dimer, trimer and tetramer, respectively) (Fig. 14d). This was the highest iSF yield with long-lived triplet state achieved in a tetracene compound in solution at that time, and it is also the only case with a triplet quantum yield exceeding 100%. The high long-lived triplet yield of tetramer 41 makes it a good candidate for the application of SF in solar energy conversion processes.

Subsequently, in cooperation with Zhang's group, the iSF mechanism in these oligomers was illustrated clearly using magnetic-field-dependent time-resolved fluorescence and TA spectroscopic technology.<sup>173</sup> In these oligomers, the <sup>1</sup>(TT) state is formed first, and it then dissociates directly into free triplets in the dimer (Fig. 15a) or free triplets in the trimer and tetramer

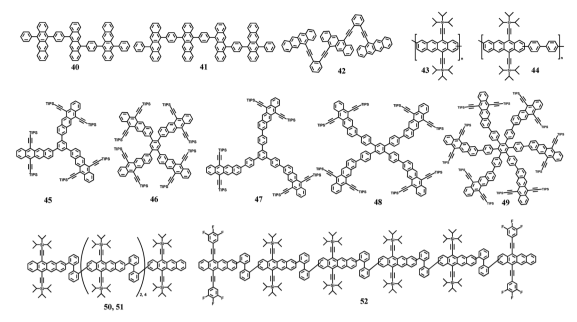


Fig. 13 Molecular structures of tetracene oligomers.

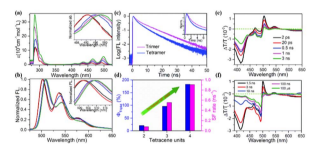


Fig. 14 (a) and (b) Absorption (a) and fluorescence (b) spectra of dimer 4, trimer 40 and tetramer 41 and their corresponding monomers. (c) Fluorescence dynamics of trimer 40 and tetramer 41. (d) Triplet yield and SF rate of dimer 4, trimer 40 and tetramer 41. (e) and (f) fs- (e) and ns- (f) TA of tetramer 41. Reproduced with permission. <sup>172</sup> Copyright 2018, The Royal Society of Chemistry.

via a spatially separated <sup>1</sup>(TT) state (denoted as <sup>1</sup>(T···T)) (Fig. 15b). However, the dissociation of the <sup>1</sup>(TT) state is very difficult in the dimer due to the limited diffusion space. The presence of the 1(TT) and 1(T···T) state in the trimer and tetramer was confirmed by the two-rise stage ( $\sim$ 10 ps and  $\sim$ 400 ps) in their triplet dynamics (Fig. 15c-h). The fast and slow components are ascribed to the formation of the  $^{1}(TT)$  and  $^{1}(T\cdots$ T) state, respectively. The assignment of the slow rise component to the  ${}^{1}(T \cdots T)$  state in the trimer and tetramer was further proven by its suppression with an external magnetic field. That is because the dynamics of the 1(T···T) state with a weak exchange interaction is much more sensitive to the Zeeman interaction induced by the external magnetic (Fig. 16b). 168,176-178

Due to the suppressed generation of the  ${}^{1}(T \cdots T)$  state, the free triplet yields of the trimer and tetramer also decreased when applying an external magnetic field (Fig. 16c), confirming

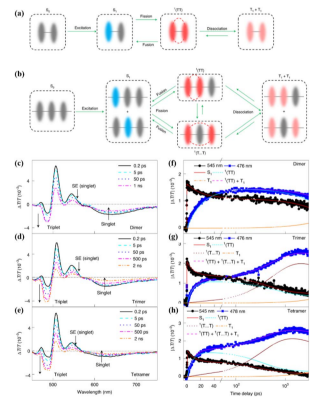


Fig. 15 (a) and (b) Scheme of the iSF process in the dimer (a), trimer and larger array systems (b). (c)–(e) fs-TA spectra of dimer (c), trimer (d) and tetramer (e). (f)-(h) Kinetic curves probed at the characteristic wavelengths showing the dynamics of the singlet and triplet states in dimer (f), trimer (g) and tetramer (h). The lines represent the dynamics of the different species (S1,  ${}^{1}(TT)$ ,  ${}^{1}(T\cdots T)$ , T1 and the sum of the triplet populations) derived with the kinetic model (a) and (b) by a global fitting analysis. Reproduced with permission.<sup>173</sup> Copyright 2021, Springer Nature.

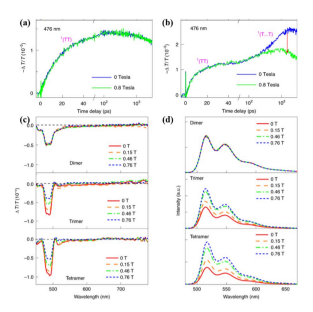


Fig. 16 (a) and (b) Dynamics of the singlet and triplet states of dimer (a) and trimer (b) with and without application of an external magnetic field of 0.8 T. (c) TA spectra of these oligomers recorded at a time delay of 100 ns, while applying a magnetic field of different magnitudes. (d) Fluorescence spectra of these oligomers with an applied magnetic field of different magnitudes. Reproduced with permission. Copyright 2021, Springer Nature.

the important contribution of the  $^1(T\cdots T)$  state to free triplet generation. Additionally, the fluorescence intensity of the trimer and tetramer was enhanced when an external magnetic field was applied (Fig. 16d), which is because the suppressed generation of the  $^1(T\cdots T)$  state makes a certain portion of the singlet population that is originally converted into free triplets through the  $^1(T\cdots T)$  state recombine through the radiative emission pathway with a longer fluorescence lifetime.

However, the fast component, representing the formation of the <sup>1</sup>(TT) state, is nearly not independent of the magnetic field due to its strong intertriplet coupling (Fig. 16b).<sup>178</sup> Thus, the triplet dynamics, triplet yield and fluorescence intensity of the dimer are not dependent on the magnetic field (Fig. 16a and d). Different from the report by Korovina *et al.*,<sup>99</sup> rotational motions of the chromophores are important for the dissociation of the <sup>1</sup>(TT) state. This result highlights the important role of the <sup>1</sup>(T···T) state in the generation of the free triplet and provides an alternative route for highly efficient <sup>1</sup>(TT) dissociation, representing an important step towards the practical applications of iSF materials because the free triplets can be harvested much more efficiently than the <sup>1</sup>(TT) state through interfacial charge transfer.<sup>179</sup>

In addition, we also prepared a tetracene trimer with a face-to-face stacked structure (42), which showed strong inter-chromophore electronic coupling. Different from the corresponding dimer (6), the TA spectrum of the triplet obtained from iSF for trimer 42 more closely resembled that of the free triplet obtained from the sensitization experiment. Furthermore, the lifetime of the triplet of trimer 42 is much longer than that of the  $^1(TT)$  state obtained in dimer 6, but shorter than that of the free triplet. These results suggest that the  $^1(T\cdots T)$  state

may be formed in trimer **42**. However, no long-lived free triplet is formed in this trimer, similar to the above-mentioned trimer **40** and tetramer **41**, which may be due to the strong electronic coupling between the two adjacent tetracene units.

Campos' group reported two tetracene polymers linked *via* the 2-positon of tetracene, where one was bridged directly (43), while the other bridged *via* a diphenyl group (44). The TA experiment revealed that ultrafast iSF proceeded in these two polymers, which had a short lifetime (~8 ns). No long-lived free triplet was generated in these two polymers. However, in tetracene dimer 18 reported by Hasobe's group, with a similar structure to polymer 44, a high yield of long-lived free triplet was achieved. This difference may because the conformational flexibility that can promote the <sup>1</sup>(TT) separation is hampered in polymer 44 with long chain character.

Recently, Kim *et al.* reported a series of tetracene dendrimers (45–49) to mimic complicated SF dynamics in amorphous solids. <sup>107</sup> These dendrimers can be classified into two categories, *i.e.*, one type containing 45 and 46, showing relatively fast iSF, and the other including 47–49, which exhibit slow iSF. Notably, iSF proceeded with two time constants for dimer 46, which is attributed to the formation of two <sup>1</sup>(TT) states of different boundness. The less-bound <sup>1</sup>(TT) state at *para*-linked tetracene is formed *via* through-bond interactions with a fast rate, whereas the close bound <sup>1</sup>(TT) state at *ortho*-linked tetracene is formed *via* through-space interactions with a slow rate. This work demonstrates the effect of structural heterogeneity on SF and highlights the importance of microscopic ordering between SF chromophores to maximize the efficiency of the exciton multiplication process for application in organic photovoltaic devices.

Under the guidance of dimer 14, Hasobe's group further synthesized a series of linear homo-oligomers (tetramer (Tc)<sub>4</sub> (50), hexamer (Tc)<sub>6</sub> (51)) and hetero-hexamer (TcF<sub>3</sub>-(Tc)<sub>4</sub>-TcF<sub>3</sub>, 52) with a 3,4,5-trifluoropheny substituent on the terminal position of the two tetracenes (Fig. 13). 181 TcF3 acts as an energy acceptor, exciton marker and vibronic promotor (Fig. 17a). The systematic comparison of these homo-oligomers and heterooligomer enables the effect of oligomerization on the <sup>1</sup>(TT) dissociation to be clarified. The TA spectra showed that iSF proceeded in all these compounds (Fig. 17c and d). The free triplet yield of homo-oligomers was enhanced gradually with an increase in the number of number of tetracene units (165%, 173% and 182% for (Tc)<sub>2</sub>, (Tc)<sub>4</sub>, and (Tc)<sub>6</sub>, respectively), which is attributed to the increased entropy change  $(\Delta S)$  of <sup>1</sup>(TT) dissociation with the increase in the number of tetracene units caused by the larger change  $(\Delta \Delta W)$  in the number of  $T_1 + T_1$  with respect to <sup>1</sup>(TT), as revealed by the calculation of the thermodynamic parameters (Fig. 17e and f). More importantly, the trapped triplet excitons were observed ( $\Phi_{TrT} = 176\%$ ,  $\tau_{TrT} = 0.30$ ms) after the exciton migration in TcF<sub>3</sub>-(Tc)<sub>4</sub>-TcF<sub>3</sub> due to the introduction of the energy acceptor (Fig. 17d). Furthermore, the  $\Delta S$  value increased in TcF<sub>3</sub>-(Tc)<sub>4</sub>-TcF<sub>3</sub> compared to (Tc)<sub>6</sub>, which is ascribed to the enhanced  $\Delta \Delta W$  caused by the vibronic levels by the rotation of the trifluorophenyl groups at the terminal TcF<sub>3</sub>. This effect also contributed to the <sup>1</sup>(TT) dissociation in TcF<sub>3</sub>-(Tc)<sub>4</sub>-TcF<sub>3</sub> besides the increase in the number of tetracene units. This work presents a new design strategy for achieving

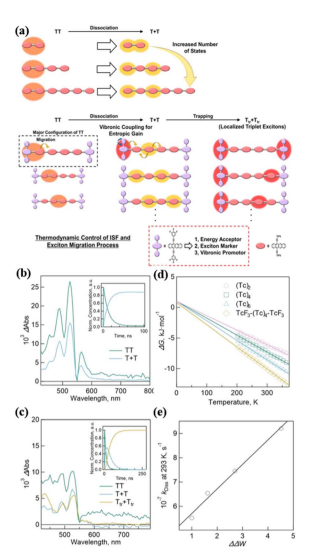


Fig. 17 (a) Conceptual schemes for the triplet exciton migration including <sup>1</sup>(TT) dissociation and exciton trapping processes in heterohexamer. (b) Species-associated spectra (SAS) of (Tc)<sub>6</sub>. The inset shows decay profiles of  $^{1}(TT)$  (green) and T+T (blue). (c) SAS of  $TcF_{3}-(Tc)_{4} TcF_3$ . The inset shows decay profiles of  $^1(TT)$  (green), T + T (blue) and  $T_{tr}$ +  $T_{tr}$  (orange). (d) The plot of  $\Delta G$  vs. temperatures. (e) Plot of  $k_{Diss}$  vs.  $\Delta\Delta W$ . Reproduced with permission.<sup>181</sup> Copyright 2023, Wiley.

efficient exciton transport processes in iSF oligomers by promoting 1(TT) dissociation and controlling the sequential exciton trapping process.

# Triplet harvesting in tetracene oligomers

Harvesting the two triplet states obtained from SF efficiently is a precondition for achieving its practical application. To achieve highly efficient extraction of the two triplets, the triplet state should have a long lifetime. Interestingly, dimer 14 shows a high yield ( $\sim$ 175%) of long-lived triplet (0.29 ms). With this dimer in hand, Nakamura et al. investigated the triplet electron transfer from dimer 14 to the electron acceptor chloranil.

The TA experiment using a mixture of dimer 14 and chloranil demonstrated that the ESA signal of the tetracene cation and the chloranil anion emerges as the triplet signal vanishes (Fig. 18b and c), suggesting the presence of electron transfer from the triplet state of the dimer to chloranil. With an increase in the chloranil concentration, the electron transfer rate gradually became fast (Fig. 18d), confirming that the electron transfer is a diffusion-controlled process. 182,183 More importantly, the quantum yield of the electron transfer reached  $\sim$ 173%, which is comparable to the triplet yield ( $\sim$ 175%). This suggests that quantitative two-electron transfer proceeded from the two triplets of dimer 14 to chloranil. This provides a new perspective for the construction of future solar energy conversion systems with SF.

To harvest two triplet excitons of singlet fission (SF) via twoelectron transfer efficiently, revelation of the key factors that influence the two-electron transfer process is necessary. Thus, our group investigated the effect of the intertriplet distance on the two-electron transfer process utilizing our previously prepared tetracene oligomers (dimer 4, trimer 40 and tetramer 41) (electron donor) and TCNQ (electron acceptor) for the first time (Fig. 19).184 In these compounds, the lifetimes of the triplet state are almost identical, but the separation distances between the two triplet states are different. This allowed us to reveal the effect of the separation distance between the two triplet excitons on the electron-transfer process independently. The TA spectra of the mixture of oligomers and TCNQ showed that the ESA signal of the TCNQ anion appears with the decay of the triplet state (Fig. 19c and d), suggesting that the electron transfer

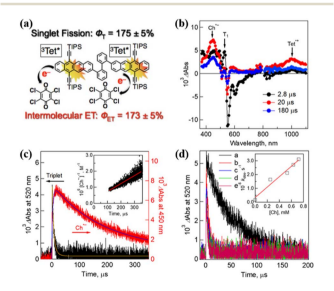


Fig. 18 (a) Schematic diagram of intermolecular electron transfer from iSF of tetracene dimer 14 to chloranil. (b) µs-TA spectra of dimer 14 (50  $\mu$ M) in the presence of 250  $\mu$ M chloranil in Ar-saturated PhCN after excitation at 532 nm. (b) Corresponding time profiles at 450 and 520 nm. The inset shows the second-order plots. (c) Time profiles of absorption at 520 nm in the presence of different concentrations of chloranil: (a) 0 mM, (b) 0.25 mM, (c) 0.53 mM, (d) 0.63 mM, and (e) 0.72 mM. The inset shows the plot of the pseudo-first-order rate constant  $(k_{obs})$  vs. the concentration of chloranil. Reproduced with permission.75 Copyright 2019, the American Chemical Society.

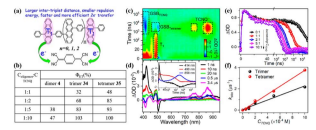


Fig. 19 (a) Schematic diagram of intermolecular electron transfer from iSF of dimer 4, trimer 40 and tetramer 41 to TCNQ. (b) Electron transfer efficiency ( $\Phi$ ) from dimer 4, trimer 40 and tetramer 41 to TCNQ. (c) ns-TA absorption spectra of tetramer 41 in the presence of TCNQ in degassed PhCN. (d) Single-wavelength dynamics for a mixture of the tetramer 41 and TCNQ monitored at different wavelengths. (e) Triplet dynamics of the tetramer probed at 490 nm in the presence of different concentrations of TCNQ in degassed PhCN. (f) Plots of the pseudo-first-order rate constant ( $k_{\rm obs}$ ) vs. concentration of TCNQ. Reproduced with permission. (184 Copyright 2022, the American Chemical Society.

occurs from the triplet state of the oligomers to TCNQ. With an increase in the concentration of TCNQ, the electron transfer rate gradually became faster (Fig. 19e), indicating that the electron transfer is a diffusion-controlled process. More importantly, the electron transfer efficiency reached 100% at high TCNQ concentrations, suggesting that a two-electron transfer process occurred. Notably, at the same TCNQ concentration, the electron transfer efficiency from dimer 4 to trimer 40, and then tetramer 41 increased gradually, and the corresponding electron transfer rate increased. This suggests that a larger intertriplet distance in iSF molecules is more favorable for the occurrence of two-electron transfer. This is attributed to the more negative free energy change induced by the smaller Coulomb repulsion energy between the two positive charges generated after the two-electron transfer. 185,186 A small intertriplet distance leads to a large Coulomb repulsion energy, which can cause a more positive free energy change and make the second-electron transfer less efficient or even be hindered. This work demonstrates the important role of the Coulomb repulsion energy in the two-electron transfer from the two triplets of iSF molecules for the first time and provides useful guidelines to design iSF molecules that can exhibit efficient twotriplet-electron transfer.

Additionally, Compos; group fabricated active layers composed of polymer 43 and a series of acceptor molecules (PC60BM, C60 and PDIF-CN2, Fig. 20a) into organic photovoltaic (OPV) devices. <sup>180</sup> OPV devices that primarily harvest triplets from an SF material should show an increased photocurrent at low magnetic fields, followed by a decrease and eventual saturation with an increase in the magnetic field. <sup>187,188</sup> The reason for this feature is that the number of triplet–triplet pairs with singlet character increases from three at a low magnetic field. Instead, the number of triplet–triplet pairs with singlet character decreases to two at a high magnetic field. This change increases, and then decreases the SF rate, leading to a positive, and then negative change in photocurrent if the device operates *via* majority triplet transport as a result of SF.

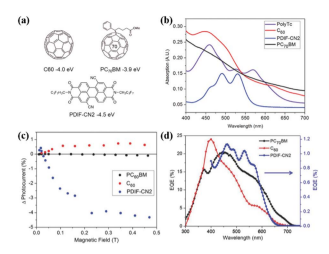


Fig. 20 (a) Structures of the acceptors used in this work, with their LUMO levels relative to vacuum. (b) Steady-state absorption profiles of the individual layers of acceptors and polymer 43. (c) Change in photocurrent with an applied magnetic field on polymer OPVs incorporating PC60BM (black) C60 (red) or PDIF-CN2 (blue) as the electron acceptor. (d) EQE characteristics of polymer OPVs incorporating PC70BM (black) C60 (red) or PDIF-CN2 (blue) as the electron acceptor. EQE values for the PDIF-CN2-containing device were scaled by 20 for clarity. Reproduced with permission. Copyright 2017, Wiley-VCH.

This characteristic effect of photocurrent enhancement at low magnetic field, and reduction at high magnetic field is observed for the acceptor PDIF-CN2, but not for C60 or PC60BM. Furthermore, the external quantum efficiency (EQE) is enhanced, benefitting from the contribution from polymer 43 (Fig. 20d). This, together with the characteristic behaviour of the photocurrent as function of magnetic field, indicate the extraction of multiple charge carriers generated from iSF in polymer 43 when PDIF-CN2 is used as the acceptor. This is because the much lower LUMO of PDIF-CN2 in comparison to that of fullerenes provides a suitable driving force for the dissociation of the triplets generated. This work represents the first example of iSF-generated triplet extraction in devices, exhibiting the potential of iSF materials for use in next-generation devices.

Recently, Guldi's group prepared a photoelectrode composed of carboxylate-anchored pentacene dimer and ZnO/ indium-zinc oxide (IZO), and successfully injected the triplet excitons of the pentacene dimer generated via iSF into ZnO/ IZO. <sup>190</sup> This photoelectrode achieved an electron injection rate of  $4.4 \times 10^{11} \ {\rm s}^{-1}$  and carrier multiplication rate of 130%. Unfortunately, the overall device efficiencies of the abovementioned two examples are rather moderate. Thus, further optimization of iSF molecules and the device structure is needed for achieving high device efficiencies.

# 5 Influence factors on endothermic SF systems

A high yield of long-lived triplet states from SF is more favorable for its application in solar energy conversion. Next, we summarize the factors influencing the overall SF dynamics (including the formation and separation of the <sup>1</sup>(TT) state) of tetracene oligomers. This will give significant insights for achieving high yield of long-lived free triplet states in endothermic SF systems.

#### 5.1 Electronic coupling

Electronic coupling plays a crucial role in the overall SF dynamics of tetracene oligomers. A relatively strong interchromophore electronic coupling is needed for the fast and efficient formation of the <sup>1</sup>(TT) state in tetracene derivatives. Weakly coupled tetracene oligomers show slow and inefficient <sup>1</sup>(TT) formation. This is consistent with the fact that the SF rate is proportion to the strength of the electronic coupling predicted by the Fermi Golden Rule.191 However, in exothermic SF systems, such as pentacene, efficient formation of the <sup>1</sup>(TT) state can be still realized although weak coupling is adopted.86 These studies indicate that this effect is particularly pronounced in endothermic and isothermal systems compared to that in exothermal systems. However, the strong electronic coupling will accelerate the direct deactivation process of the  $^{1}(TT)$  state to the ground state  $(^{1}(TT) \rightarrow S_0 + S_0)$  and the reverse reaction of SF (TTA), and thus prevent the dissociation of the <sup>1</sup>(TT) state into the free triplet state with long lifetime. 81,139 Furthermore, too strong electronic coupling will decrease the S<sub>1</sub> energy seriously. This will make SF more endothermic, and thus slow down the formation rate of the 1(TT) state. Thus, a balanced electronic coupling strength is necessary for achieving a high yield of the overall SF process. As is known, electronic coupling is dependent on the intermolecular orientation and distance, which can be controlled by various linkers. Notably, it is difficult to achieve efficient overall SF in tetracene oligomers via simple electronic coupling regulation. Thus, other factors should be considered to achieve high efficiency of long-lived free triplet states.

#### 5.2 Conformational flexibility

As discussed by Hasobe's and Thompson's group,98,99 conformational flexibility plays an important role in the dissociation of the <sup>1</sup>(TT) state to generate free triplet states in tetracene derivatives. The conformational change at the <sup>1</sup>(TT) state can decrease the intertriplet spin-spin exchange interaction (J) by reducing the orbital overlap, and then decelerate the TTA process and accelerate the dissociation of the <sup>1</sup>(TT) state to the free triplet state, resulting in a high yield of free triplet states. For example, the two tetracene units of compound 18 at the <sup>1</sup>(TT) state are nearly planer, which shows strong orbital overlap. However, the conformation at the T1 state is changed to orthogonal with a negligible orbital overlap.48 The reduced orbital overlap induces electronic decoupling between the two triplets, and then leads to the generation of free triplets. This pivotal role has also been highlighted in some other oligomers composed of pentacene and perylene. 58,101,105,111,192,193 In perylene oligomers, the torsional disorder results in the formation of free triplet states from the separation of the  $^1(TT)$  state.  $^{58}$  Thus, a too rigid structure of tetracene oligomer is not a preferred

conformation to achieve a highly efficient overall SF process. Based on the premise of the appropriate intermolecular electronic coupling strength, a more flexible conformation that is determined by the linker may result in a high free triplet yield. Besides, for oligomers with a highly symmetrical structure, their conformation change may break the molecular symmetry to activate SF.<sup>43,163</sup>

#### 5.3 Covalent linker energy-level alignment

The effect of energy-level alignment of the covalent linker on SF dynamics has been illustrated by several groups. 68,84,87,99 The closer energy of the frontier molecular orbital of the linker with that of the chromophore, the better mixing of the linker orbitals with that of the chromophores, and thus the stronger resonance coupling between the chromophores. Stronger resonance coupling will result in the faster formation of the <sup>1</sup>(TT) state. To promote the dissociation of the <sup>1</sup>(TT) state into the T<sub>1</sub> state, a larger interchromophore distance will be more favorable. However, a decreased SF rate will be induced simultaneously. Fortunately, the resonance effect can overcome this negative effect. For example, compound 23 showed faster SF than of 16, despite having a longer linker.84 Similarly, this phenomenon was also observed in a series of pentacene dimers.84 Specifically, the selection of a covalent linker with an appropriate energy level and length will be more favorable for achieving a high yield of long-lived triplet states. Thus, the energy-level alignment, length and flexibility of the covalent linker should be considered when choosing the linker to connect the chromophores for achieving efficient SF.

#### 5.4 Oligomer size

As discovered by us and Zhang's group, 132,172,173 with an increase in the number of repeating unit, the formation rate and yield of the free triplet state increase significantly. This is because the increased number of structural units can provide sufficient statistical space to promote the formation of the spatially separated triplet pair state (1(T···T)), which mediates the generation of the free triplet state. This effect is more important for endothermic SF systems because the formation of the <sup>1</sup>(T··· T) state can effectively avoid the recombination of the <sup>1</sup>(TT) state to the S<sub>1</sub> state, which is a competing process with its separation into free triplet states. This provides an alternative route for the highly efficient dissociation of the <sup>1</sup>(TT) state into free triplet states. This effect also contributes to SF in a series of perylene oligomers.58 Additionally, an increase in the number of repeat units in oligomers can result in an increase in entropy, which is beneficial for the formation and separation of the <sup>1</sup>(TT) state.58,191,194

#### 5.5 Aggregation state behaviour

The ultimate goal for SF is its integration in photovoltaic devices. Considering that the active layer of solar cells is in the solid state, the effect of the aggregation behaviour of SF molecules in the solid state on SF dynamics should be addressed. Many crystal films and nanoparticles of tetracene derivatives with different arrangements show SF. However, the SF rate and

efficiency are strongly dependent on the molecular packing. In this case, herringbone packing may be the preferred conformation for SF in the solid state. For example, unsubstituted tetracene, rubrene and 5,12-bis(phenylethynyl) tetracene films, with herringbone accumulation, generally show efficient SF. 195-198 Alternatively, a 5,12-diphenyl tetracene film with a coplanar arrangement demonstrated relatively inefficient SF.20 This is consistent with the computation result that SF can occur efficiently in acene when the molecule adopts a herringbone arrangement. 199,200 However, very little work has been reported to reveal the effect of the aggregating behavior of iSF molecules on SF in the solid state, especially for tetracene. 100,180,200-202 When an iSF molecule is prepared in the solid state, its intermolecular interactions are dominated by through-space coupling besides intramolecular interactions. Thus, iSF and xSF will be both accessed possibly, resulting in their competition. This depends on the relative strength of the intramolecular and intermolecular interactions. For example, Korovina et al. observed ultrafast iSF in amorphous films of dimer 6 with strong intramolecular coupling.100 More importantly, <sup>1</sup>(TT) dissociation was also observed due to the entropic gains, which is often an issue in covalent dimers in solution. However, when the intramolecular coupling was weak, the through-space intermolecular coupling led to xSF instead of iSF. This was proven by the study of the terrylene-3,4:11,12-bis(dicarboximide) dimer in the solid state, which exhibited strong xSF.201 This provides an approach to achieve high yield of longlived free triplet states by combining strong intramolecular coupling and weak intermolecular coupling. Xia's group demonstrated that the behaviour of changing the aggregation state of the iSF material from the solution phase to the film could effectively cut off the transportation channel of the triplet excitons and obtain long-lived triplet excitons without affecting the rapid generation of triplet states.200

As can be seen, there are many factors that affect the overall SF process including the formation and separation of the <sup>1</sup>(TT) state. It is easy to understand that among the factors listed above, electronic coupling plays a decisive role in the SF process, which directly determines whether the material can undergo SF, especially for endothermic and isothermal systems. Furthermore, the electronic coupling also influences the dissociation of the <sup>1</sup>(TT) state. A moderate electronic coupling strength can ensure that both the formation and separation of the <sup>1</sup>(TT) state proceed simultaneously. The aggregation behaviour of iSF molecules in the solid state also plays an important role in the overall SF process by controlling the intramolecular and intermolecular interactions. In the case of covalent linker energy-level alignment, it will accelerate the formation of the <sup>1</sup>(TT) state by increasing the resonance interaction with the chromophore. Instead, the conformational flexibility and oligomer size mainly influence the separation process of the <sup>1</sup>(TT) state. Increasing the conformational flexibility and oligomer size can promote the dissociation of the <sup>1</sup>(TT) state by suppressing the TTA process and the reverse process of SF. Sometimes, different factors show the opposite effect. Consequently, it is difficult to make an absolute definition on how to get a high yield of long-lived free triplet excitons.

However, through a systematic comparison of the abovementioned tetracene oligomers, we can get some insights into the design of efficient SF materials based on endothermic chromophores. A structure with an appropriate electronic coupling strength and large conformational flexibility is more favorable for the generation of free triplet excitons. Additionally, an increase in the oligomer size based on the premise of proper electronic coupling can also promote the dissociation of the <sup>1</sup>(TT) state into free triplet states.

## 6 Conclusions and outlook

In this review, initially, we described the progress in iSF of tetracene oligomers, especially the relationship between their molecular structures and SF properties. Then, the factors that affect the overall SF process in tetracene oligomers were summarized, including electronic coupling, conformational flexibility, covalent linker energy, oligomer size and aggregation behaviour. The results showed that a molecular structure with appropriate electronic coupling strength and large conformational flexibility may be a good choice for achieving a high yield of long-lived free triplet states to construct iSF materials based on endothermic SF chromophores. Additionally, based on the premise of an appropriate electronic coupling strength, molecular oligomers with three or more structural units will promote the dissociation of the <sup>1</sup>(TT) state into free triplet states.

Many breakthroughs have been made in SF field in recent years. Several new series of SF materials have been developed, and their related SF mechanism has been established. However, several challenges still exist, which should be addressed in the future application of SF in solar conversion processes. Firstly, the number of SF materials, with fast rate, high triplet state energy (1.2-1.3 eV), high yield (>100%) of long-lived free triplet states and good light/heat stability, is still small. This will limit the application of SF to some extent. A fast SF rate can avoid extra competing processes, such as energy and charge transfer occurring from the singlet state at molecule/semiconductor interfaces in the device. The high triplet state energy is more feasible for tandem with solar cells (such as Si). The long-lived triplet state guarantees the efficient harvesting of two triplet states in solar conversion processes. Notably, a high yield of long-lived triplet states is difficult to be achieved. This is because it needs highly efficient <sup>1</sup>(TT) formation and separation simultaneously. Generally, efficient <sup>1</sup>(TT) formation needs strong electronic coupling, especially for endothermic and isothermic systems, but strong electronic coupling suppresses the dissociation of the <sup>1</sup>(TT) state into long-lived free triplet states. Therefore, achieving a high yield of long-lived triplet states by controlling the electronic coupling strength and other factors listed above is the main difficulty before the application of SF. Secondly, how to efficiently harvest the two triplet states of SF is another difficulty. Triplet electron transfer and triplet energy transfer have been demonstrated to be useful methods to harvest the two triplet states with quantum dots and metal oxides as well as organic molecules as energy or electron acceptors, 4,75,189,190,203-212 but

there are few examples with high efficiency. Furthermore, the real extraction mechanism of the electron transfer or the energy transfer is controversial. Additionally, some photovoltaic devices incorporating SF showed that several competitive routes exist in the extraction of the two triplets, such as energy/ electron transfer from the singlet state, TTA and the direct decay of the <sup>1</sup>(TT) state into the ground state. <sup>93,187,213-215</sup> More importantly, the exact working mechanism of SF-based devices remains ambiguous. Thus, the resolution of all the abovementioned difficulties is urgent to achieve the practical application of SF.

Endothermic iSF systems have some advantages for application in solar energy conversion process due to the lower energy losses in these SF systems. Although highly efficient iSF has been achieved in tetracene oligomers, tetracene derivatives are less stable toward to light and oxygen, which limits their practical application. Therefore, it is urgent to achieve highly efficient iSF in some molecular oligomers constructed by stable endothermic chromophores. Of course, improving the stability of tetracene is also a good choice. In this case, several methods can be used to improve the stability of the tetracene. Firstly, the introduction of some specific functional groups in tetracene has a positive effect on its oxidative stability. For example, the ethynyl group can improve the oxidation stability of tetracene.216 It also avoids the thermal decomposition of the resulting endoperoxides (EPOs). Moreover, the free radical stabilization of the intermediate produced by the photooxidation of tetracene is conducive to C-O cleavage, and thus it can promote the re-conversion of EPOs to the parent tetracene. Secondly, the introduction of heteroatoms is another method to enhance the stability of tetracene by affecting its electronic and properties. 217-221 For example, the introduction of nitrogen and fluorine (or chlorine) atoms as electron-withdrawing groups can reduce the energy of the frontier molecular orbital, which is expected to improve the stability of tetracene. With an increase in the number of heteroatoms, this effect is more obvious. Thirdly, the introduction of heterocycles (such as thiophene and pyridine) in the tetracene framework can also increase its stability.218,222 The stability of the oligomers prepared with the modified tetracene will be enhanced significantly. Additionally, to maximize the efficiency of solar cells, it is necessary to broaden the absorption of tetracene in the red or NIR region. However, although the polymerization of tetracenes into polymers is a choice, their solubility and processability will decrease remarkably due to the increase in their molecular rigidity. In this case, the introduction of solubilizing groups (such as long branched aliphatic chains) in the tetracene skeleton can improve the polymer solubility efficiently. This method has been widely applied in the design of polymer photovoltaic materials.223,224 Furthermore, the introduction of conjugated groups (such as alkynyl group and cyano group) in the tetracene parent can redshift its absorption significantly. More importantly, these groups can also increase the solubility of tetracene to some extent. Thus, polymerizing tetracenes with redshifted absorption into polymers may be another effective method.

## **Author contributions**

T. Wang and H. Liu wrote the manuscript. X. Wang, L. Tang, J. Zhou, X. Song and L. Lv, helped search related literature. W. Chen helped revise the manuscript. H. Liu and X. Li conceived the idea, designed the structure of the review, revised the draft of the manuscript and supervised the work. X. Li and Y. Chen provided funding support.

### Conflicts of interest

The authors declare no competing financial interests.

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