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Introduction

Currently, singlet fission (SF), in which one singlet excited state is converted into two triplet excited states,^{1,2} serves as the primary means to overcome the Shockley–Queisser limit.^{3–6} A number of energetic and conformational requirements must be satisfied to realize SF. Among these requirements, the most important component is the energy of the singlet excited state, $E(S_1)$, relative to the energy of the triplet excited state, $E(T_1)$, namely $E(S_1) \ge 2(E(T_1))$.^{1,2,7,8} Early stage investigations of intermolecular SF focused on crystalline materials.^{5,9–15} With respect to SF efficiency, crystal packing, molecular orbital overlap, *etc.* of the chromophores are decisive.^{16–18}

SF has been documented for tetra-, penta-, and hexacene in the solid state,¹⁹⁻²⁴ as well as for tetra- and pentacene in

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Solution-based intramolecular singlet fission in cross-conjugated pentacene dimers†

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We show unambiguous and compelling evidence by means of pump-probe experiments, which are complemented by calculations using *ab initio* multireference perturbation theory, for intramolecular singlet fission (SF) within two synthetically tailored pentacene dimers with cross-conjugation, namely **XC1** and **XC2**. The two pentacene dimers differ in terms of electronic interactions as evidenced by perturbation of the ground state absorption spectra stemming from stronger through-bond contributions in **XC1** as confirmed by theory. Multiwavelength analysis, on one hand, and global analysis, on the other hand, confirm that the rapid singlet excited state decay and triplet excited state growth relate to SF. SF rate constants and quantum yields increase with solvent polarity. For example, **XC2** reveals triplet quantum yields and rate constants as high as $162 \pm 10\%$ and $(0.7 \pm 0.1) \times 10^{12} \text{ s}^{-1}$, respectively, in room temperature solutions.

solution.^{25–28} In the solid state, tetracene features nearly degenerate singlet excited states and correlated pairs of triplet excited states, *i.e.*, $E(S_1) \approx 2E(T_1)$, rendering the standard enthalpy for intermolecular SF negligible. In contrast, the singlet excited state energy in pristine pentacene in the solid state is more than twice the energy of a correlated pair of two triplet excited states, namely $E(S_1) \geq 2(E(T_1))$. Thus, SF is exoergic and unidirectional for pentacene. In hexacene, SF is linked to the formation of a correlated pair of triplet excited states and phonons.^{5,9,23,29–31}

In contrast to SF in the solid state, SF in solution has only recently been discovered for pentacenes, with yields reaching as high as 200%.^{26–28} Solution-state SF has also been demonstrated in tetracene dimers, but with a SF yield of only 3%.²⁵

It is logical to hypothesize that the relative proximity between neighboring acenes is critical to the process of SF, and the molecular geometry as well as the π - π overlap would be dominant factors for efficient SF.32-35 One can envision a plethora of acenes that might be synthesized to explore specific aspects of SF in dimeric structures.36-39 We have recently described the photophysical behavior of a set of three regioisomeric pentacene dimers, linked via an ortho-, meta-, and *para*-phenylene (Fig. 1).²⁷ The observed triplet quantum yield in these dimers is as high as $156 \pm 5\%$. It is interesting to note that the *meta*-linked dimer gives the best SF performance, a molecule in which the two pentacenes are not directly conjugated through the intervening phenylene ring, in contrast to either the ortho- or para-linked dimer. In light of the general structure of the *meta*-dimer, we speculate that moderating the electronic communication between two acene chromophores



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Fig. 1 Phenylene tethered pentacene dimers studied previously²⁷ and cross-conjugated pentacene dimers investigated in the present work.

might be an important design concept for the optimization of SF. In order to test this hypothesis, dimers based on diethynyland tetraethynylethenes (DEE and TEE, respectively) have been designed, in which the two pentacenes communicate in the ground state only *via* cross-conjugation.^{40,41} The constrained conjugation of DEEs and TEEs has been used, for example, to form molecular wires,^{42–44} switches,⁴⁵ and nonlinear optical materials.⁴⁶ Furthermore, the contrasting behavior of chromophores linked by *meta*-phenylene groups *versus* a cross-conjugated skeleton has been a topic of somewhat controversial discussion in organic electronics.^{47,48}

Considering the excited states that are involved, their nature, and their electronic coupling, two main mechanisms for SF have been proposed over the years, the direct and the mediated, and the predominance of one or another remains a subject of controversy.^{2,7,9,17,32,49-62} In the direct mechanism, the absorbing state transforms into a correlated pair of triplet excited states, also called multiexcitonic (ME) or doubly excited state, which later dissociates into two (non-interacting) T₁ states. In the mediated mechanism, the transformation of the absorbing state into the ME state, which then dissociates into two T₁ states, is facilitated by coupling to higher-lying CT states.2,55 Recent studies have refined these traditional viewpoints, providing more information about the detailed interplay of the different electronic states involved in the process. In particular, one of the studies postulates that the initial excitation of the molecular system produces a state that is not an eigenstate of the system but a coherent superposition of the (lowest-lying) absorbing state, CT states, and the ME state.^{9,17} The latter ultimately splits into two separated T₁ states after decoherence. Finally, a very recent model suggests that the non-interacting T1 states are directly populated from the absorbing state without the participation of any intermediate electronic state in the process.⁶²

In the present contribution, we have designed and examined two novel pentacene dimers **XC1** and **XC2**, that feature a cross-conjugated framework to mediate the electronic coupling between the two pentacene moieties. The cross-conjugated structural modification is expected to be particularly helpful as a means to either enhance or attenuate the through bond electronic couplings between the pentacene chromophores.⁴¹ Furthermore, **XC1** and **XC2** feature rigid structures, with geometries that are similar to that found in *ortho*-dimer (*vide infra*).

Results and discussion

Synthesis

Cross-conjugated dimers **XC1** and **XC2** were synthesized from the reaction of masked pentacene **1**, which has proven to be a valuable precursor for the stepwise assembly of pentacene derivatives *via* transition metal catalyzed reactions.^{27,37,39} Thus, the reaction of **1** with either dibromoolefin 2⁶³ or 3⁶⁴ was carried out using a Sonogashira cross-coupling protocol, and the TEE and DEE cores of **4** and **5** were formed in acceptable yields of 63% and 60%, respectively (Scheme 1).

Subsequently, Sn(n)-mediated reductive aromatization afforded XC1 and XC2 in 82% and 95% yield, respectively. Dimers XC1 and XC2 are stable to air and moisture under



Scheme 1 Synthesis of XC1 and XC2.

normal laboratory conditions, and they have good solubility in typical organic solvents.

A single crystal of **XC2** suitable for X-ray crystallographic analysis was grown from a THF solution layered with MeOH and allowed to evaporate at 4 °C. The solid-state structure emphasizes the proximity of the two pentacenes. Fig. 2 documents that through space interactions are accessible in both the solid and solution states *via* carbon–carbon contacts between the two pentacene units as short as 3.34 Å (in the crystal), a result that agrees well with that found theoretically (3.41 Å, see ESI[†]).

Singlet and triplet excited state energetics

We turned to absorption and emission spectroscopies combined with electrochemical analyses to establish the singlet and triplet excited state energetics. In terms of ground state features, we note short wavelength absorptions in the region of 350-500 nm and long wavelength absorptions in the 500-775 nm region (Fig. 3). As a result of the rigid structure of XC1 and XC2, both sets of absorptions show distinct vibrational fine structures. The spectroscopically relevant long wavelength absorptions are at 681 nm (XC1) and 683 nm (XC2). In concentration dependent experiments, which obey the Lambert-Beer law for dilute solutions, the following extinction coefficients were determined in benzonitrile: 28 800 M⁻¹ cm^{-1} (XC1) and 48 900 M^{-1} cm^{-1} (XC2). It is notable that the values for XC1 and XC2 are less or exactly twice that of TIPS pentacene 7 (Fig. 4), respectively. A likely rationale infers stronger electronic communications between the two pentacenes in XC1 relative to XC2.

In terms of excited state features, exciting into the long wavelength absorptions leads to fluorescence maxima at 691 and 688 nm and fluorescence quantum yields of $0.03 \pm 0.01\%$ and $0.12 \pm 0.04\%$ for **XC1** and **XC2**, respectively, in benzonitrile. Notably, these features are all linked to the singlet excited state with energies of 1.81 eV. Finally, no appreciable triplet excited state phosphorescence is observed for **XC1** or **XC2** in either solutions at room temperature or in frozen media experiments upon excitation in the long wavelength absorption region.

To complement absorption and fluorescence studies, squarewave voltammetric analyses of **XC1** and **XC2** have been carried out in benzonitrile solutions containing 0.1 M TBAPF₆



Fig. 2 X-ray crystallographic structure of **XC2** illustrating the spatial arrangement and proximity of the two pentacene units in the dimer. THF solvent molecules and hydrogen atoms have been omitted for clarity.



Fig. 3 Top: Room temperature absorption spectra of XC1 (black line) and XC2 (red line) in benzonitrile. Bottom: Normalized fluorescence spectra of XC1 (black dots) and XC2 (red dots) in benzonitrile solutions following photoexcitation at 625 nm with an optical density of 0.05. The corresponding fluorescence quantum yields in benzonitrile are 0.03 \pm 0.01% for XC1 and 0.12 \pm 0.04% for XC2 (measured relative to a zinc phthalocyanine used as a reference compound⁶⁵).



Fig. 4 Molecular structures of C₆₀ derivative 6 and TIPS pentacene 7.

as supporting electrolyte. From the one electron oxidations at +0.44 and +0.35 V, as well as the one electron reductions at -1.27 and -1.39 V for **XC1** and **XC2** (*versus* Fc/Fc⁺), respectively, highest occupied molecular orbital/lowest unoccupied molecular orbital gaps of 1.71 and 1.74 eV are calculated, which are in sound agreement with the absorption/fluorescence assays.

TIPS pentacene

First, we examined 7 in 2-methyltetrahydrofuran (Me-THF, 3.0×10^{-4} M) with excitation at 656 nm to stimulate population of only vibrational states of the first singlet excited state manifold (Fig. 5). Additionally, the photon flux has been adjusted to 6.6×10^9 photons per pulse to excite, on average, 10% or less of 7 in its ground state in order to rule out multiple excitations.



Fig. 5 Upper left: Chemical structure of TIPS pentacene 7. Upper right: Differential absorption spectra (visible and near-infrared) of the spectra shown in the lower right with time delays of 2.6 ns (black) and 132 ns (gray). Lower left: Time absorption profiles of the spectra shown in the lower right at 447 nm (black), and 498 nm (red) illustrating the dynamics of the singlet excited state formation followed by the singlet to triplet transformation in the form of intersystem crossing, and triplet decay. Lower right: Differential absorption changes (visible and near-infrared) obtained upon femtosecond pump probe experiments (656 nm) of 7 (3.0 \times 10⁻⁴ M) in argon-saturated Me-THF at room temperature with several time delays between 0 and 50 μ s.

For photoexcited 7, the major deactivation pathway includes a 12.3 ± 0.1 ns intersystem crossing, through which the singlet excited state transforms into the corresponding triplet manifold. Once formed, the triplet excited state deactivates with $24.0 \pm 0.5 \,\mu$ s to reinstate the singlet ground state of 7. As far as the singlet excited state characteristics are concerned, maxima at 447, 508, 538, 572, 845, and 1400 nm are noted, as well as minima at 595 and 641 nm. An additional feature at 710 nm is observed, which corresponds to stimulated emission. Triplet excited state characteristics include maxima at 465 and 498 nm and minima at 546, 587, and 641 nm. Owing to a moderate triplet quantum yield of only 16% for 7 the resulting characteristics turn out to be extremely weak.

Triplet-triplet sensitization

To investigate the triplet excited states characteristics *via* an inefficient intersystem crossing – *vide supra* – we turned to intermolecular, that is, diffusion or activation controlled, triplet-triplet sensitization experiments using the well-studied C_{60} derivative **6** as a photosensitizer to examine the triplet excited state characteristics of **7** as well as of **XC1** and **XC2** (Fig. 4, 6 and S9–S17†). Incentives for the use of **6** are an excitation wavelength independent triplet quantum yield of unity, a triplet excited state energy 1.5 eV higher than that of **7**, **XC1**, and **XC2**, a triplet excited state molar extinction coefficient of 16 000 M^{-1} cm⁻¹ at its transient maximum at 695 nm, and a lack of chemical reactions with pentacene derivatives in the



Fig. 6 Upper left: Chemical structure of XC1. Upper right: Differential absorption spectrum (visible and near-infrared) of the spectra shown in the lower right with a time delay of 5.1 μ s representing the triplet excited state of XC1. Lower left: Time absorption profiles of the spectra shown in the lower right at 510 nm (black), 584 nm (red), 678 nm (orange), and 740 nm (gray) illustrating the dynamics of the 6 singlet excited state formation followed by the intersystem crossing to the corresponding 6 triplet excited state and transduction of triplet excited state energy to XC1. Lower right: Differential absorption changes (visible and near-infrared) obtained upon femtosecond pump probe experiments (480 nm) of 6 (8.0×10^{-5} M) and XC1 (1.0×10^{-4} M) in argon-saturated toluene at room temperature with several time delays between 0 and 400 μ s.

applied concentration range.⁶⁶ Excitation of 6 at 480 nm $(8.0 \times 10^{-5} \text{ M})$ resulted in differential absorption changes on the nanosecond/microsecond time scale featuring a 695 nm maximum and a 840 nm shoulder. These originate from a 1.5 ns intersystem crossing between the corresponding singlet and triplet excited manifolds. In the absence of 7, XC1, XC2, and molecular oxygen, these features decay within 33.0 \pm 1.0 µs due to triplet-triplet and triplet-ground state annihilations. Addition of 7, XC1, or XC2 in the concentration range from 10⁻⁵ to 10⁻⁴ M affords nearly diffusion-controlled deactivations of the triplet excited state of 6 with rate constants on the order of 2.5×10^9 M⁻¹ s⁻¹.^{67,68} Simultaneously with such a faster decay newly developing transient features are noted including a maximum at 510 nm and a minimum at 678 nm for XC1 as well as a maximum at 508 nm and minima at 622 and 679 nm for XC2. Here, the excited state minima are precise mirror images of the ground state absorption maxima. Notable is also the close resemblance with the features seen for 7 and several other pentacence dimers,²⁷ from which we conclude a transduction of triplet excited state energy to XC1 and XC2. The lifetimes of the triplet excited states of XC1 and XC2 are bimolecular in nature. From the corresponding rate constants, namely $(1.1 \pm 0.1) \times 10^8$ and $(0.7 \pm 0.1) \times 10^8$ M⁻¹ s⁻¹ for XC1 and XC2, respectively, we deduce activation limited triplet excited state decays including triplet-triplet annihilation as a major component and collision induced triplet-ground state annihilation as a minor component.

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With the triplet excited state molar extinction coefficients of 6 at hand, we calculated the triplet excited state extinction coefficients at the 508 nm maximum, the 622 nm minimum, and the 679 nm minimum at each concentration of XC2, as well as at the 510 nm maximum and the 678 nm minimum of **XC1.** By plotting the extinction coefficients as a function of concentration, we noticed a saturation behavior towards the high concentration regime. At an extrapolation to 5×10^{-4} M the triplet-triplet sensitization is quantitative relative to the initially photoexcited 6. Of importance is the fact that the molar extinction coefficients of 21 200 and 52 800 M⁻¹ cm⁻¹ for XC2 at the 622 nm minimum and the 679 nm minimum, respectively, are in sound agreement with the ground state molar extinction coefficients of 30 800 and 57 900 M⁻¹ cm⁻¹ at 620 and 678 nm, respectively. Likewise the value of 28 700 M⁻¹ cm⁻¹ for XC1 at its 678 nm minimum matches that seen at 675 nm in the ground state absorption. From the aforementioned we infer quantitative bleaching of the ground state for XC1 or XC2 and, in turn, triplet excited states, in which both pentacenes are excited. Considering pseudo first order conditions, namely a laser intensity that is equivalent to approximately 10^{-7} M triplet excited state of 6 and 5.0×10^{-5} M of XC1 or XC2, we postulate that the triplet excited state molar extinction coefficients are only half of the extrapolated values, that is, 45 200, 10 600 and 26 400 M^{-1} cm⁻¹ at the 508 nm maximum, the 622 nm minimum, and the 679 nm minimum, respectively. In fact, with this assumption they are identical to what has been seen for 7, which features molar extinction coefficients about half of that of XC2.

To substantiate our postulate we performed actinometry using pristine C₆₀.⁶⁹ Here, we used solutions of pristine C₆₀ with four different concentrations and solutions of either 7, XC1, or XC2, each of them with exactly the same absorptions at the 387 nm excitation wavelength, and pumped them with the same laser power. In Fig. S18,^{\dagger} the intensities of the C₆₀ singlet excited state markers, on one hand, and the 7, XC1, and XC2 singlet as well as triplet excited state markers, on the other hand, are plotted as a function of absorption. This dependence documents the linear dependence of each excited marker within the range of absorptions. Correlating the respective slopes for pristine C₆₀ with those for 7, XC1, or XC2 enable the determination of the molar extinction coefficients for 7, XC1, and XC2 at, for example, time delays of 1.5 ps. To this end, we determined singlet excited state molar extinction coefficients for XC2 of 14400 $M^{-1}\ cm^{-1}$ at the 622 nm minimum and of 24 100 M^{-1} cm⁻¹ at the 677 nm minimum. For XC1, the corresponding singlet excited state molar extinction coefficient is 20 200 $M^{-1}\ \text{cm}^{-1}$ at 679 nm. All of these values relate to half of the ground state extinction coefficients/ ground state bleaching and suggest singlet excited states, in which only a single pentacene is photoexcited. The latter assumption is in sound agreement with the low laser flux of 6.6×10^9 photons per pulse. Additional support was provided by the fact that the 647 nm value of 15 000 M^{-1} cm⁻¹ for 7 correlates quantitatively with the ground state extinction coefficient/ground state bleaching. In case that 7, XC1, and XC2

show quantitative intersystem crossing, that is, affording triplet quantum yields of 100%, the triplet excited state molar extinction coefficients at 647, 679, and 677 nm, respectively, should match those of the singlet excited states.

Singlet fission in XC1 and XC2

For XC2 in benzonitrile, the singlet excited state features are discernable as a 455 nm maximum and a broad band in the NIR as well as minima at 627 and 687 nm directly upon photoexcitation at 656 nm (Fig. 7). In contrast to that observed for 7, where a slow intersystem crossing dominates the initial 20 ns, the singlet excited state features of XC2 are replaced within 15 ps by characteristics that match those seen in the triplettriplet sensitization experiments (Fig. S17[†]). In particular, these are maxima at 486, 517, 853, and 976 nm and minima at 628 and 685 nm. From multiwavelength analyses, which afforded strictly unimolecular, intramolecular dynamics in the range from 1×10^{-5} to 1×10^{-4} M of XC2, we conclude that these features correspond to the rapidly decaying singlet excited state $((0.7 \pm 0.1) \times 10^{12} \text{ s}^{-1})$ and the simultaneously grow of the triplet excited state markers. Please note that there is a slight blue shift, from 687 to 685 nm, as a consequence of this transformation. A closer look at delay times up to 2000 ps, however, reveals the metastability of the newly formed triplet excited state. Considering, on one hand, that the experimental conditions preclude multiple excitations of XC2 and, on the other hand, that triplet-triplet annihilation contributes most strongly to the triplet excited state decay we tested the



Fig. 7 Upper left: Chemical structure of **XC2**. Upper right: Differential absorption spectra (visible and near-infrared) of the spectra shown in the lower right with time delays of 1.8 ps (black) and 15.0 ps (gray). Lower left: Time absorption profiles of the spectra shown in the lower right at 486 nm (black), 517 nm (red), 628 nm (orange), and 687 nm (gray) illustrating the dynamics of the singlet excited state formation followed by the singlet to triplet transformation in the form of singlet fission and triplet-triplet annihilation. Lower right: Differential absorption changes (visible and near-infrared) obtained upon femtosecond pump probe experiments (656 nm) of **XC2** (5.0×10^{-5} M) in argon-saturated benzonitrile at room temperature with several time delays between 0 and 7500 ps.

impact of different concentrations in the range from 1×10^{-5} to 1×10^{-4} M. Regardless of the concentration, we note only unimolecular and invariant dynamics with a rate constant of $(2.54 \pm 0.08) \times 10^9$ s⁻¹. As such, we hypothesize unimolecular triplet-triplet annihilations, which requires that two triplets are localized within XC2 and this is only feasible by SF - vide infra. Spectroscopic corroboration for two photoexcited pentacenes came from a 2-3 nm blue-shift not only relative to the singlet excited state bleaching but also relative to the triplet excited state bleaching seen in direct excitation and triplettriplet sensitization experiments, respectively. The product of the triplet-triplet annihilations is the singlet ground state of XC2 via radiationless relaxation,⁷⁰ and independent on concentration or laser power. But, a spectroscopically invisible excited state cannot be ruled out. As a complement to multiwavelength analyses, we performed global analyses with the Glotaran software package (Fig. 9).⁷¹ The results, as they were obtained from the global fitting, are a qualitative and quantitative match to those gathered in the multiwavelength approach for XC2 and XC1. For example, the singlet excited state in XC2 transforms sequentially into the triplet excited state and ground state within 0.7 \pm 0.1 and 445 \pm 20 ps, respectively.

Very similar observations are made for **XC1** in benzonitrile (Fig. 8 and 9), in terms of differential absorption changes and kinetics *versus* **XC2**. In particular, the singlet excited state for **XC1** converts with similar decay and growth kinetics of $(0.65 \pm$



Fig. 8 Upper left: Chemical structure of **XC1**. Upper right: Differential absorption spectra (visible and near-infrared) of the spectra shown in the lower right with time delays of 1.8 ps (black) and 13.8 ps (gray). Lower left: Time absorption profiles of the spectra shown in the lower right at 484 nm (black), 513 nm (red), 646 nm (orange), and 685 nm (gray) illustrating the dynamics of the singlet excited state formation followed by the singlet to triplet transformation in the form of singlet fission and triplet-triplet annihilation. Lower right: Differential absorption changes (visible and near-infrared) obtained upon femtosecond pump probe experiments (656 nm) of **XC1** (5.0×10^{-5} M) in argon-saturated benzonitrile at room temperature with several time delays between 0 and 7500 ps.

 $(0.1) \times 10^{12} \text{ s}^{-1}$ via SF into a metastable triplet excited state. In THF and toluene the corresponding kinetics tend to be as slow as $(0.63 \pm 0.1) \times 10^{12}$ and $(0.50 \pm 0.1) \times 10^{12}$ s⁻¹. Analyses of the long wavelength minima reveals a blue-shift of 2-3 nm when comparing the singlet and triplet excited state spectra. Similar to XC2, the metastability observed for XC1 is seen in the form of unimolecular, intramolecular triplet-triplet annihilations $((2.08 \pm 0.07) \times 10^9 \text{ s}^{-1})$. The latter results in fast and quantitative ground state recovery on a time scale of less than 3000 ps. Resembling XC2, the two steps, namely SF and the decay of the correspondingly formed triplet excited state, are clearly deconvoluted. If at all, the different cross conjugation in XC2 results in slightly faster triplet excited state formation and slightly faster triplet excited state decay relative to XC1, but the differences are small. The absorption spectra - vide supra point possibly to stronger electronic coupling in XC1, which is likely derived from through-bond interactions. As such, we hypothesize that solution SF seems more favorable in the latter scenario.

To corroborate SF as the mechanism for singlet to triplet conversion we have used three different approaches, that is, the sensitization method, the bleaching method, and the actinometry method, to determine the triplet quantum yields. In the first approach, 6 is used as an external reference to derive the transient extinction coefficients. Firstly, we have revisited the triplet-triplet sensitization experiments, whose differential absorption spectra in the high concentration regime, where the triplet energy transfer efficiency is close to unity, enable derivation of triplet molar extinction coefficients for XC2 of 45 200, 10 600 and 26 400 M⁻¹ cm⁻¹ at the 508 nm maximum, the 622 nm minimum, and the 679 nm minimum, respectively. Next, we have determined the singlet molar extinction coefficients with the help of conventional femtosecond transient absorption measurements. XC2, for example, features only very weak ground state absorption at 456 nm in benzonitrile, and, as such, we take the amplitude of the differential absorptions at 455 nm after 1 ps and relate it entirely to the singlet excited state concentration formed upon excitation. Finally, the differential absorption changes of 0.0242 a.u. at 455 nm and 0.0679 a.u. at 518 nm are converted into concentrations of the singlet excited state of 4.64×10^{-6} M and of the triplet excited state of 7.51×10^{-6} M. This ultimately converts to a triplet quantum yield of $162 \pm 10\%$ for XC2.

Independent confirmation of the triplet quantum yield comes from the second approach (bleaching method), in which no external reference is used to calculate the transient extinction coefficients of the singlet and triplet excited state. In this case, the amplitudes of the ground state bleaching in benzonitrile at 687 nm (-0.0245 a.u) and 685 nm (-0.0314 a.u.) are used to calculate the transient concentrations as 4.64×10^{-6} and 5.96×10^{-6} M for the singlet and the triplet excited state, respectively. This affords a triplet quantum yield of $128 \pm 5\%$ for **XC2**. The high triplet quantum yield along with its ultrafast formation provide strong evidence that the triplet excited state for **XC2** forms by intramolecular SF between the two pentacenes. Moreover, higher quantum yields in more polarizable solvents –



Fig. 9 Deconvoluted transient absorption spectra of the singlet (black) and the triplet excited state (red) of XC1 and XC2 as obtained by global analysis (visible and near-infrared). Inset: Evolution of the population of the singlet (black) and the triplet excited state (red).

mean values of $128 \pm 10\%$ in toluene *versus* $162 \pm 10\%$ in benzonitrile – corroborate the charge transfer nature of the mediating step. Analogously, the triplet quantum yields for **XC1** are derived as $127 \pm 10\%$ with an external reference and $124 \pm 5\%$ without an external reference in benzonitrile. In toluene, the corresponding yields for **XC1** are lower with values of $119 \pm$ 10% and $114 \pm 5\%$, respectively (Table 1).⁷² In a third approach we used the results from the actinometry experiments to calculate the triplet quantum yields. Correlating the slopes for the respective singlet and triplet excited state features enable the determination of triplet quantum yields of $111 \pm 5\%$ for **XC1** and $107 \pm 5\%$ for **XC2** in toluene, in good agreement with the results obtained by means of the first and second approach.

Theory

To further solidify the experimental results, we have investigated the electronic states of relevance for SF using multireference perturbation theory methods (see ESI† for details). The calculated vertical excitation energies, dipole moments, oscillator strengths, and character of the relevant adiabatic states are summarized in Table 2. For **XC1** and **XC2**, the ME state is the lowest-lying singlet excited state (see Table 2 and ESI†). Furthermore, the vertical excitation energies of the lowest-lying absorbing S₂ state of **XC1** (1.88 eV) and **XC2** (1.90 eV) agree well with the experimental data. Both **XC1** and **XC2** exhibit CT states (S₄ and S₅) with moderate dipole moments that are relatively high in energy (~0.5–0.6 eV) compared to the S₂ state.

Table 1 Triplet excited state quantum yields (Φ_{Triplet}) for XC1 and XC2 calculated *via* the sensitization method, the bleaching method, and the actinometry

		Sensitization method	Bleaching method	Actinometry	
		Triplet quantum yield [%]			
XC1	Toluene	119	114	111	
	THF	131	121	_	
	Benzonitrile	127	124		
XC2	Toluene	128	109	107	
	THF	144	113	_	
	Benzonitrile	162	128	_	

To shed light on the SF mechanism, we have determined the diabatic electronic states and their couplings employing Truhlar's four-fold-way diabatization procedure (see ESI[†]).^{73,74} We report in the following the results obtained for XC1 and **XC2**. Following previous works,^{2,18,60} the diabatic states are classified as the singlet ground state $|S_0S_0\rangle$, the correlated triplet pair (or ME) state $|T_1T_1\rangle$, the locally excited (in one pentacene chromophore) singlet states $|S_1S_0\rangle$ and $|S_0S_1\rangle$ and the charge transfer states $|CA\rangle$ and $|AC\rangle$. In the latter, one of the two pentacenes is formally a radical cation (C), while the other is a radical anion (A). For XC1 and XC2, the diabatic coupling matrices are presented in Table 3. As Table 3 documents, XC1 and XC2 each exhibit a moderate electronic coupling between the locally excited states $(|S_1S_0\rangle$ and $|S_0S_1\rangle$) and $|T_1T_1\rangle$. Specifically, $|S_1S_0\rangle$ and $|S_0S_1\rangle$ couple to $|T_1T_1\rangle$ with a strength of 9 and -12 meV, respectively, in XC1. In XC2, the corresponding values are -1 and 2 meV. In contrast, the electronic coupling between $|S_1S_0\rangle$ or $|S_0S_1\rangle$, and the charge transfer states |CA> and |AC> range from 30 to 140 meV. As such, this is nearly an order of magnitude larger than that between the locally excited and the correlated triplet pair state $|T_1T_1\rangle$ (Table 3). $|T_1T_1\rangle$ also couples to the $|CA\rangle$ and $|AC\rangle$ states with 10-60 meV in XC1 and 23-27 meV in XC2.

To dissect contributions from either the direct or the mediated SF mechanism and the role of CT states, we estimated the magnitude of the effective coupling, V_{eff} , of the lowest lying absorbing singlet state $|S_1S_0\rangle$ to $|T_1T_1\rangle$ (similar for $|S_0S_1\rangle$ to $|T_1T_1\rangle$) as:^{55,75}

$$V_{\rm eff} \approx V_{\rm S_1S_0,T_1T_1} - 2 \frac{V_{\rm S_1S_0,CA}V_{\rm CA,T_1T_1} + V_{\rm S_1S_0,AC}V_{\rm AC,T_1T_1}}{[E(\rm CT) - E(\rm T_1T_1)] + [E(\rm CT) - E(\rm S_1S_0)]} \ (1)$$

where E(i) are the energies of the corresponding diabatic states, E(CT) is the mean of E(CA) and E(AC), and $V_{i,j}$ are the couplings between the states *i* and *j*. The first term in eqn (1) denotes the direct coupling and the second term the indirect coupling mediated by the CT states. The results in Table 4 indicate that in both **XC1** and **XC2** the direct coupling of $|S_1S_0\rangle$ to $|T_1T_1\rangle$ is significantly smaller than the indirect coupling *via* CT states. The differences between both types of coupling are smaller in the case of **XC2** relative to **XC1**. For the second bright state $|S_0S_1\rangle$, which is quasi-degenerate to $|S_1S_0\rangle$,

Table 2 Vertical excitation energy (ΔE in eV),^a modulus of the dipole moment (μ , D),^b oscillator strength (f),^b and character for the lowest lying singlet excited states (char.)^c of **XC1** and **XC2**

	XC1				XC2			
State	ΔE	μ	f	Char.	ΔE	μ	f	Char.
S_1	1.82	2.31	< 0.001	ME	1.84	0.06	< 0.001	ME
S ₂	1.88	4.43	0.880	LE	1.90	1.74	0.770	LE
$\tilde{S_3}$	1.94	3.39	0.610	LE	1.95	1.74	0.640	LE
S_4	2.31	8.23	0.004	CT	2.36	8.31	0.005	CT
S ₅	2.40	8.79	0.002	CT	2.40	8.33	0.001	CT

^{*a*} Calculated at the XMCQDPT/DZV level of theory. ^{*b*} Calculated at the CASSCF level of theory. ^{*c*} Character of excited state: ME = multiexcitonic state, LE = excited states that correlate with the plus and minus combinations of locally excited states of both pentacene monomers, CT = charge transfer states.

Table 3 Energies and coupling matrix elements (meV) of the low-lying diabatic electronic states of XC1 (top) and XC2 (bottom) calculated at the XMCQDPT/DZV level of theory

XC1	$ S_0S_0\rangle$	$ T_1T_1\rangle$	$ S_1S_0 angle$	$ S_0S_1\rangle$	$ CA\rangle$	$ AC\rangle$
$ S_0S_0\rangle$	0	10	261	105	81	18
$ T_1T_1\rangle$	10	1779	9	-12	61	11
$ S_1S_0\rangle$	261	9	1863	-29	141	71
$ S_0S_1\rangle$	105	-12	-29	1885	-31	34
CA	81	61	141	-31	2217	21
$ AC\rangle$	18	11	71	34	21	2323
XC2	$ S_0S_0\rangle$	$\left T_{1}T_{1} \right\rangle$	$ S_1S_0\rangle$	$ S_0S_1 angle$	$ CA\rangle$	$ AC\rangle$
S _o S _o >	0	-13	-119	139	-140	-118
$ T_1T_1\rangle$	-13	1806	-1	2	-27	23
$ S_1S_0\rangle$	-119	-1	2009	-94	-136	-72
$ S_0S_1\rangle$	139	2	-94	2025	138	87
$ CA\rangle$	-140	-27	-136	138	2172	-130
AC	-118	23	-72	87	-130	2282

Table 4 Direct, indirect, and total coupling (meV) of the lowest lying absorbing singlet states $|S_1S_0\rangle$ and $|S_0S_1\rangle$ to $|T_1T_1\rangle$ for XC1 and XC2, calculated at the XMQCDPT/DZV level of theory using Truhlar's four-foldway diabatization method

System	State	Direct	Indirect	Total
XC1	$ S_1S_0\rangle$	9.39	-23.69	-14.30
	$ S_0S_1\rangle$	-12.09	3.94	-8.15
XC2	$ S_1S_0\rangle$	-0.83	-7.62	-8.45
	$ S_0S_1\rangle$	1.78	6.72	8.50

XC1 exhibits an effective coupling smaller than that found for $|S_1S_0\rangle$ and where the major contribution stems from the direct coupling of $|S_0S_1\rangle$ to $|T_1T_1\rangle$. On the other hand, **XC2** shows an effective coupling between $|S_1S_0\rangle$ and $|T_1T_1\rangle$ where the major contribution corresponds to the indirect coupling *via* CT states. Considering that the CT states are higher in energy than the absorbing states we postulate that SF is likely to proceed predominantly *via* a superexchange-like mechanism in both molecular systems, although in the case of **XC1** we expect a more significant contribution of the direct mechanism to the overall process than in **XC2** (see Table 4). This is further corroborated by the experimentally found dependence

of the triplet quantum yield on the polarizability of the solvents employed. The results in Table 4 also indicate that the coupling is somewhat larger in **XC1** compared to **XC2**, which may explain the stronger broadening of the absorption spectrum of **XC1** (see Fig. 3).

Conclusions

We have designed cross-conjugated pentacene dimers, **XC1** and **XC2**, to fine-tune the electronic coupling element between the two pentacenes and to gain control over intramolecular SF in room temperature solutions. To this end, **XC1** features slightly stronger electronic coupling, governed by throughbond interactions, than **XC2**. Kinetic and spectroscopic analyses were based on applications of multiwavelength analysis, on one hand, and global analysis using the program Glotaran, on the other hand. Both led to identical results and corroborate that the cross-conjugated tethers maintain the electronic couplings lending, however, to slightly higher triplet quantum yields for **XC2** than for **XC1**. As a matter of fact, the highest triplet quantum yield observed is $162 \pm 10\%$.

Our synthetic, spectroscopic, and computational efforts highlight the importance of establishing the right balance between triplet excited state formation/singlet excited state deactivation and triplet excited state decay towards the optimization of SF. An outstanding question remains the loss mechanism(s) from the singlet state in our dimers, and ongoing work in our laboratories focuses on realizing SF quantum yields close to unity through further controlling intramolecular interactions in synthetically designed dimers.

Experimental section

The synthesis of **XC1** and **XC2**, materials, general methods, and computational details are described in the ESI[†].

For the photophysical characterization (see ESI[†]), the samples were placed in fluorometric cuvettes with different pathways and, when necessary, purged with argon. Steady-state UV-vis absorption spectra were acquired at room temperature (RT) using a Perkin Elmer Lambda 2 spectrometer. Steady-

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state fluorescence spectra were carried out at a FluoroMax3 spectrometer from Horiba in the visible detection range (RT) and at a FluoroLog3 spectrometer from Horiba in the near infrared. Femtosecond (fs) transient absorption experiments were carried out with an amplified Ti:Sapphire CPA-2110 fs laser system (ClarkMXR: output 775 nm, 1 kHz, 150 fs pulse width) using transient absorption pump/probe detection systems (Helios and Eos, Ultrafast Systems).

The theoretical characterization was based on the following methods. The ground state equilibrium geometries of XC1 and **XC2** were optimized using density functional theory (DFT),⁷⁶ employing the B3LYP correlation-exchange functional and the def2-TZVP basis set.77,78 Dispersion interactions were taken into account using Grimme's empirical dispersion correction.⁷⁸ Vertical excitation energies were calculated using the extended multi-configurational quasi-degenerate perturbation theory (XMCQDPT)⁷⁹ method and a double- ζ basis set,⁸⁰ employing state average (SA) complete active space self-consistent field (CASSCF)^{81,82} wave-functions as reference and an intruder state avoidance shift of 0.02 au.83 The active space employed in the SA-CASSCF calculations comprised four electrons in four orbitals (HOMO and LUMO per pentacene unit) and six roots with equal weights were used. Dipole moments and oscillator strengths were calculated at the SA-CASSCF level. Diabatic states were built using Truhlar's four-fold-way of diabatization method implemented in GAMESS.73,74,84 All calculations were carried out using the electronic structure packages TURBOMOLE⁸⁵ and GAMESS.⁸⁶

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