Distinct properties of the triplet pair state from singlet fission

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Singlet fission, the conversion of a singlet exciton (S1) to two triplets (2 × T1), may increase the solar energy conversion efficiency beyond the Shockley-Queisser limit. This process is believed to involve the correlated triplet pair state (TT). Despite extensive research, the nature of the (TT) state and its spectroscopic signature remain actively debated. We use an end-connected pentacene dimer (BP0) as a model system and show evidence for a tightly bound (TT) state. It is characterized in the near-infrared (IR) region (~1.0 eV) by a distinct excited-state absorption (ESA) spectral feature, which closely resembles that of the S1 state; both show vibronic progressions of the aromatic ring breathing mode. We assign these near-IR spectra to (TT) → S1 and S1 → S2 transitions; S1 and S2, likely come from the antisymmetric and symmetric linear combinations, respectively, of the S2 state localized on each pentacene unit in the dimer molecule. The (TT) → S1 transition is an indicator of the intertriplet electronic coupling strength, because inserting a phenylene spacer or twisting the dihedral angle between the two pentacene chromophores decreases the intertriplet electronic coupling and diminishes this ESA peak. In addition to spectroscopic signature, the tightly bound (TT) state also shows chemical reactivity that is distinctively different from that of an individual T1 state. Using an electron-accepting iron oxide molecular cluster [Fe8O4] linked to the pentacene or pentacene dimer (BP0), we show that electron transfer to the cluster occurs efficiently from an individual T1 in pentacene but not from the tightly bound (TT) state. Thus, reducing intertriplet electronic coupling in (TT) via molecular design might be necessary for the efficient harvesting of triplets from intramolecular singlet fission.

INTRODUCTION

Singlet fission is a many-body photophysical process in molecules where the photoexcited singlet (S1) splits into two triplets (2 × T1) with spin conservation (1, 2). Since its discovery, efficient singlet fission has been reported mostly for solids and aggregates of conjugated molecules (1, 2), and a dominant mechanistic picture is the molecular dimer model (3, 4)

\[ S_0 + S_1 \leftrightarrow ^1(TT) \leftrightarrow 2T_1 \]  

where S0 is the ground state and the intermediate (TT) is the correlated triplet pair with both singlet spin and double-excitation characters. Despite its prevalent use, Monahan et al. (5, 6) pointed out the inadequacy of the dimer model in describing inherently delocalized excitons in the solid state. Exciton delocalization has been cited as an important driving force for singlet fission (7–10). This problem is circumvented in recent demonstrations of efficient singlet fission in single molecules [particularly in dimers of acenes (11–16)] that allow for accurate application of the dimer model and for closely connecting experiment with theory (17). The isolation of the transient (TT) state in a single molecule leads to a much longer lifetime than that in the condensed phase, thus allowing spectroscopic characterization of this ambiguous and poorly understood state. This is exemplified in the detection by electron spin resonance spectroscopy in pentacene dimers of the quintet state, (TT), which is mixed with the (TT) state as predicted by the spin Hamiltonian (18).

The (TT) state is a singlet excited state with double-excitation characters and differs from 2 × T1, not only by the electronic and spin entangled nature of the former but also by the presence of orbital overlap, which changes its excitation energy from the sum of two triplet energies. Scholes (19) recently clarified some persistent confusion on the (TT) state in theoretical descriptions. The energetic difference between the correlated triplet pair state and two individual triplets, that is, the triplet pair binding energy, can be as large as 1 eV, as is known for the excited states of oligoenes (20–22), including carotenoids (23), where the tightly bound triplet pair has been called the "dark" S1 state serving as a sink for nonradiative recombination and a less tightly bound triplet pair (S′) has been associated with singlet fission (15, 24–26). In contrast, in prototypical model systems of pentacene or tetracene dimers (both covalent and van der Waals), computational analysis predicted little, if any, triplet pair binding energy (17, 27–32). However, a recent finding of similar (TT) lifetimes in polypentacene and dipentacene indicates that the triplet pair does not dissociate even in a long conjugated chain (33), suggesting that the correlated triplet pair state is more strongly bound than previously thought.

A major obstacle to a clear understanding of the (TT) state is the lack of spectroscopic signatures from experiments. Zhu and co-workers applied time-resolved two-photon photomission spectroscopy to quantitatively determine the energetic position of the (TT) state from its ionization potential (IP) in crystalline pentacene (34), tetracene (10), and hexacene (6). This approach is unambiguous only for hexacene (6) where the (TT) state is energetically well separated from S1 but is difficult for other singlet fission systems where the (TT) states are in close energetic resonance with S1. The most widely used technique to probe singlet fission has been transient absorption (TA) spectroscopy, but most studies to date have only identified spectral features assigned to S1 and T1 states, and there has been little explanation as to why these TA peaks nearly always overlap (12–16, 35, 36). Exceptions to this prevalent practice can be found in the recent work of Sanders et al. (11) who...
found, in pentacene dimers, an excited-state absorption (ESA) peak at
~690 nm whose magnitude is strongly correlated with the strength of
intertriplet electronic coupling and in the work of Pensack et al. (37)
who observed near-infrared (IR) (1200 to 1400 nm) ESA in pentacene
aggregates assigned to $^1(TT)$ but not to the triplet pair labeled
$^1(T\ldots T)$, which has lost electronic coherence but retained spin coherence. These
two examples reveal the presence of spectroscopic signatures for the
$^1(TT)$ state in TA, but the origins of these transitions and their relationships
to the energetics of the $^1(TT)$ state remain unknown.

The distinct electronic structure of the $^1(TT)$ state should be reflected
not only in its spectroscopic signature but also in its chemical and
physical properties. The oft-cited motivation for nearly every recent pa-
per on this subject has been the potential “usefulness” of singlet fission
to solar energy conversion. The basic argument was put forward initially
by Dexter (38) for the sensitization of conventional solar cells by singlet
fission chromophores in 1979, but a more recent paper by Hanna and
Nozik (39) on using singlet fission to increase the solar cell efficiency
above the Shockley-Queisser limit really energized the field. A number
of research groups have explored the harvesting of triplet pairs from
intermolecular singlet fission using solid interfaces between a singlet fis-
sion material and an electron or triplet acceptor material (34, 40–43).
These efforts have also led to the successful demonstration of singlet
fission–based solar cells with quantum efficiencies exceeding 100%
(44). The recent demonstration of efficient intramolecular singlet fission
in single molecules (11–16, 36) opens the door to new opportunities for
the realization of singlet fission–sensitized solar cells (45). A more
exciting opportunity is the potential for the harvesting of two electron-
hole pairs for photocatalysis, for example, by coupling a singlet fission
molecule to a molecular or cluster-based catalytic center (46) to enable
two-electron redox reactions. Unlike intermolecular singlet fission in
the solid state in which electronic delocalization (5, 47) and entropy
(10) are driving forces to split the $^1(TT)$ state to two electronically de-
coupled triplets (which can nonetheless retain spin coherence) on ultra-
fast time scales (19), the confinement in a molecular dimer or oligomer
traps the two triplets in the $^1(TT)$ state in a single molecule (18, 33).
Thus, instead of individual triplets at solid-state interfaces, the
harvesting of triplets in intramolecular single fission would likely come
from the $^1(TT)$ state. However, the two triplets in the $^1(TT)$ state from
intramolecular singlet fission can be tightly bound, and charge or energy
transfer from each triplet may be inhibited.

Here, we use triisopropylsilylethynyl-functionalized pentacene
(TIPS-Pc) dimers, each coupled at the 2-position without or with a
phenylene spacer, BP0 or BP1 (Fig. 1) (11), as well as pentacene dimers
with different dihedral angles (17), as model systems to quantitatively
probe the nature of the tightly bound $^1(TT)$ state from the ESA spectra.
Molecules of this type allow for the systematic tuning of electronic cou-
pling between the two pentacene units and between the S$_0$ and the $^1(TT)$
states, as reflected in the singlet fission time constants of $\tau_{SF} = 0.76, 20,$

![Fig. 1. The model systems for intramolecular singlet fission and triplet harvesting. (A) Schematics of BP0, BP1, [Fe$_8$O$_4$]-Pc, and [Fe$_8$O$_4$]-BP0. R = (triisopropylsilyl)ethynyl
(TIPS) for [Fe$_8$O$_4$]-Pc and (n-octyldioisopropyl)silyl)ethynyl (NODIPS) for [Fe$_8$O$_4$]-BP0. C = triisopropylsilylethynyl (TIPS) for [Fe$_8$O$_4$]-Pc and [Fe$_8$O$_4$]-BP0. The inset shows estimated IP and EA (electron affinity) from electrochemical
oxidation/reduction potentials of [Fe$_8$O$_4$] and TIPS-pentacene. (B and C) Optical absorption spectra of (B) TIPS-Pc, BP0, and BP1 in toluene and (C) [Fe$_8$O$_4$]-Pc, and
[Fe$_8$O$_4$]-BP0 in dichloromethane solutions.](http://advances.sciencemag.org/)
and 220 ps and triplet recombination time constants of $\tau_{AN} = 0.45$, 16.5, and 270 ns for dimers with zero, one, and two phenylene spacers (BP0, BP1, and BP2), respectively, obtained from an analysis of TA spectra in the visible region (11). Here, we focus on the distinct ESA peak in the near-IR region (hv \sim 1 eV), which is a signature of the $^1(TT)$ state from singlet fission in BP0. Its intensity diminishes as the intertriplet electronic coupling is lowered in BP1 or significantly decreases in bipentacene with different dihedral angles (17). The ESA peak of $^1(TT)$ in BP0 closely resembles that of the $S_1$ state in the near-IR region; both show vibronic progressions of the aromatic ring breathing mode and can be assigned to the $^1(TT)\rightarrow S_n$ and $T_1\rightarrow S_n$ transitions, respectively. This finding unambiguously establishes that $^1(TT)$ is spectroscopically distinct from $2 \times T_1$, and such a spectroscopic signature enables one to quantitatively follow the dynamics of this critical intermediate in singlet fission.

To establish the distinct chemical properties of the $^1(TT)$ state, we use the redox-active molecular cluster $\text{Fe}_8\text{O}_4\text{pz}_{12}\text{Cl}_4$ (pz, pyrazolate), which we label $[\text{Fe}_8\text{O}_4]$, as an electron acceptor (48, 49) and tether BP0 to $[\text{Fe}_8\text{O}_4]$ through a Fe-phenoxide bond (schematically illustrated in Fig. 1A). As a control, we replace the pentacene dimer BP0 with a pentacene monomer (Pc). Note that the formation of the Fe-phenoxide bond in both $[\text{Fe}_8\text{O}_4]$-Pc and $[\text{Fe}_8\text{O}_4]$-BP0 introduces a low-energy absorption tail (~1.3 to 1.8 eV; Fig. 1C). This spectral feature has been assigned to the phenolate-to-Fe(III) ligand-to-metal charge transfer (LMCT) transition (50) but may also have contributions from pentacene-to-$[\text{Fe}_8\text{O}_4]$ charge transfer (CT) transitions. We show that electron transfer from pentacene to $[\text{Fe}_8\text{O}_4]$ occurs efficiently from an individual $T_1$ state in pentacene ($[\text{Fe}_8\text{O}_4]$-Pc), but not from the tightly bound triplet pair state in $[\text{Fe}_8\text{O}_4]$-BP0. This finding establishes that the chemical property of the $^1(TT)$ state is distinctively different from that of an individual triplet and suggests that reducing intertriplet electronic coupling in $^1(TT)$ might be needed for the harvesting of triplets from intramolecular singlet fission.

RESULTS AND DISCUSSION

Spectroscopic signature of the $^1(TT)$ state

We use TA spectroscopy to probe singlet fission in BP0 and BP1 (11). We excite the $S_1$ state of each pentacene dimer at hv = 2.1 eV and probe the subsequent dynamics from the TA of a white-light continuum (Fig. 2). Figure 2 (A and B) shows TA spectra at selected pump-probe delays; $\Delta t = 0.1$ ps (red), 10 ps (purple), and 100 ps (blue) for BP0. The visible parts of the TA spectra have been discussed extensively before, and the broad positive TA features at hv ~ 1 eV, similar to the vibronic progressions of $S_0\rightarrow S_1$, $T_1\rightarrow T_2$, and $^1(T_1T_1)\rightarrow^1(T_1T_3)$ discussed above. This ESA is assigned to an $S_1\rightarrow S_0$ transition, with transition energy close to the $S_1\rightarrow S_2$ transition for a single pentacene molecule. In the absorption spectrum of TIPS-Pc in Fig. 1B (blue), there is a weakly allowed $S_0\rightarrow S_2$ peak at 2.82 eV, in agreement with the two-photon absorption spectrum of the same molecule (52). Given the $S_0\rightarrow S_1$ peak at 1.93 eV (blue spectrum in Fig. 1B), we obtain the $S_1\rightarrow S_2$ transition of 0.89 eV. In conjugated bipentacene dimers, the singlet states are described by linear combinations of two localized states on each pentacene chromophore (17). Although both symmetric and antisymmetric linear combinations are possible, the optically bright $S_1$ state in BP0 is of odd parity (u). Therefore, excited state transitions must occur to $S_0$, states of even parity (g). We assign the 0.92-eV peak to a transition from $S_1$ to the symmetric linear combination of the monomer $S_0$ states.

At longer pump-probe delays, for example, $\Delta t = 10$ ps (purple) or 100 ps (blue), when there is only the triplet pair state, the ESA spectrum blue-shifts to 1.01 ± 0.005 eV and the vibronic signature becomes better shorter than the 30-µs lifetime of an individual triplet (11). Here, we focus on the near-IR region, which provides key spectroscopic insight into the triplet pair state.

There is a distinct ESA peak at 0.922 ± 0.005 eV when the singlet dominates at $\Delta t = 0.1$ ps (red) (Fig. 2A); this peak is also accompanied by a vibronic feature on the higher energy side, with hv_vib ~ 0.17 eV, similar to the vibronic progresses of $S_0\rightarrow S_1$, $T_1\rightarrow T_2$, and $^1(T_1T_1)\rightarrow^1(T_1T_3)$ discussed above. This ESA is assigned to an $S_1\rightarrow S_0$ transition, with transition energy close to the $S_1\rightarrow S_2$ transition for a single pentacene molecule. In the absorption spectrum of TIPS-Pc in Fig. 1B (blue), there is a weakly allowed $S_0\rightarrow S_2$ peak at 2.82 eV, in agreement with the two-photon absorption spectrum of the same molecule (52). Given the $S_0\rightarrow S_1$ peak at 1.93 eV (blue spectrum in Fig. 1B), we obtain the $S_1\rightarrow S_2$ transition of 0.89 eV. In conjugated bipentacene dimers, the singlet states are described by linear combinations of two localized states on each pentacene chromophore (17). Although both symmetric and antisymmetric linear combinations are possible, the optically bright $S_1$ state in BP0 is of odd parity (u). Therefore, excited state transitions must occur to $S_0$, states of even parity (g). We assign the 0.92-eV peak to a transition from $S_1$ to the symmetric linear combination of the monomer $S_0$ states.

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Fig. 2. TA in the near-IR and visible regions reveal singlet and triplet characters of $^1(TT)$. TA spectra in (A) the near-IR and (B) the visible regions for BP0 at different pump-probe delays, $\Delta t = 0.1$ ps (red), 10 ps (purple), and 100 ps (blue), following excitation at time zero by hv = 2.1 eV. The triplet TA spectrum from sensitization (black) is also shown in (A) and (B). (C) Kinetic profiles from TA spectra for BP0 at the indicated probe photon energies. (D) TA spectra at $\Delta t = 1$ ps (red) and 100 ps (blue) for BP1 following excitation at time zero by hv = 2.1 eV. The corresponding triplet spectrum (black) from sensitization is also shown.
resolved. This ESA peak does not originate from a \( T_1 \rightarrow T_n \) transition as it is completely absent in the triplet absorption spectrum (black) from sensitization. On the basis of the similarity of this ESA peak to that of the \( S_1 \rightarrow S_n \) transition at early times, we assign the former to a \( ^1(TT) \rightarrow S_n \) transition. The \( ^1(TT) \) state in BP0 is expected to correspond to the near-IR ESA assigned to \( 1(TT) \) in the pentacene dimer, the near-IR ESA assigned to \( 1(TT) \) in a single electronic state, which can be approximately viewed as the \( S_1 \rightarrow S_n \) and \( 1(TT) \rightarrow S_n \) transition energies, \( \Delta E = 90 \text{ meV} \), is close to the predicted exoergicity of \( \sim 100 \) to \( 150 \text{ meV} \) for singlet fission in bientacene (17). The small energy spacing implies that both \( S_n \) and \( S_n' \) likely originate from different linear combinations of the \( S_2 \) monomer state of different parity. Note that, unlike the results shown here for the pentacene dimer, the near-IR ESA assigned to \( ^1(TT) \) in pentacene aggregates does not show vibronic features (37).

The ESA spectrum of the \( ^1(\text{TT}) \) state reveals its delocalized singlet and localized triplet characters in the near-IR and the visible regions, respectively. We use “delocalized singlet” or “delocalization” to emphasize \( ^1(\text{TT}) \) in a single electronic state, which can be approximately viewed as two \( T_1 \) states (on two pentacene units) that are electronically coupled and coherent. Likewise, the term “localized triplet” or “localization” refers to a \( T_1 \) state on an individual pentacene unit with physical properties that are insensitive to the presence or absence of electronic coupling and coherence with a neighboring \( T_1 \) state. Spectroscopically, delocalization and localization are reflected in the transitions \( ^1(\text{TT}) \rightarrow S_n \) and \( ^1(T_1T_1) \rightarrow ^1(T_1T_n) \), respectively. Note that the two notations, \( ^1(\text{TT}) \) and \( ^1(T_1T_1) \), describe the same triplet pair state. The kinetic profiles (Fig. 2C) for the \( ^1(\text{TT}) \rightarrow S_n \) (orange) and \( ^1(T_1T_1) \rightarrow ^1(T_1T_n) \) (green) transitions are similar; the difference at short time scales (<1 ps) can be attributed to the different overlapping contributions from ESA of \( S_1 \). Note that transitions to \( S_n \) are allowed from \( ^1(\text{TT}) \) but spin-forbidden from \( ^3(\text{TT}) \) or \( ^3(\text{TT}) \). The perfect agreement between the decays of \( ^1(\text{TT}) \rightarrow S_n \) and the \( ^1(T_1T_1) \rightarrow ^1(T_1T_n) \) signals suggests that there are negligible transitions within the triplet pair manifold, for example, \( ^1(\text{TT}) \rightarrow ^3(\text{TT}) \) or \( ^1(\text{TT}) \rightarrow ^3(\text{TT}) \), during the lifetime (450 ps) of the triplet pair state. Transitions within the triplet pair manifold are expected to occur on much longer time scales (18, 53).

Supporting the conclusion that delocalization or intertriplet electronic coupling in the \( ^1(\text{TT}) \) state is reflected in the \( ^1(\text{TT}) \rightarrow S_n \) transition strength, we find that, in BP1, the weakening of the inter-

\( T_1 \) electronic coupling diminishes its delocalized character as reflected in the \( ^1(\text{TT}) \rightarrow S_n \) transition strength (Fig. 2D) where the near-IR peak for \( ^1(\text{TT}) \) at long times, for example, \( \Delta t = 100 \text{ ps} \) (blue), becomes non-resolvable from the broad background, in distinct contrast to the \( S_1 \rightarrow S_n \) peak at \( \Delta t = 1 \text{ ps} \). In contrast, the localized character represented by the \( ^1(T_1T_1) \rightarrow ^1(T_1T_n) \) transition in the visible region remains (11).

To more quantitatively isolate the \( S_1 \) spectrum from that of the \( ^1(\text{TT}) \), we carry out global analysis based on a sequential kinetic model, \( S_1 \rightarrow ^1(\text{TT}) \rightarrow S_0 \) (11). The resulting \( S_1 \) (red) and \( ^1(\text{TT}) \) (blue) spectra are shown in Fig. 3. The global analysis yields time constants of \( 0.75 \pm 0.05 \text{ ps} \) and \( 460 \pm 10 \text{ ps} \) for singlet fission and triplet pair annihilation, respectively, in agreement with the previous report (11). Similar to the \( S_1 \rightarrow S_n \) transition, the \( ^1(T_1T_n) \rightarrow S_n \) transition is also characterized by vibronic peaks assigned to 0-0 and 0-1 transitions, with a vibrational energy spacing of 0.16 to 0.17 eV, which corresponds to the ring breathing mode of pentacene along the short molecular axis (54). In addition to the near-IR peak, the \( ^1(\text{TT}) \) state in BP0 also features a distinct peak at \( 1.810 \pm 0.005 \text{ eV} \). Similar to the transition at \( 1.012 \pm 0.003 \text{ eV} \), the peak at \( 1.810 \pm 0.005 \text{ eV} \) diminishes as the intertriplet coupling weakens in BP1 and BP2 (11). Thus, the peak at \( 1.810 \pm 0.005 \text{ eV} \) also reflects the singlet character of the \( ^1(\text{TT}) \) state and can be assigned to a \( ^1(\text{TT}) \rightarrow S_n \) transition. Because of the overlapping bleaching feature (\( S_0 \rightarrow S_1 \)), we are not able to resolve vibronic progression for this transition.

Further supporting the conclusion that the near-IR \( ^1(\text{TT}) \rightarrow S_n \) transition is a spectroscopic signature of the tightly bound triplet pair state, we turn to modified BP0 molecules with different dihedral angles. In this approach, we control the dihedral angle twist by steric hindrance from the phenyl group attached to the 1-position of one or both pentacene units in the bipentacene molecule, as shown schematically in the insets in Fig. 4 (17). Computational analysis gives dihedral angles between the two pentacene molecules of 42° and 57°, and these two

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**Fig. 3.** TA spectra of BP0 for the \( S_1 \) and \( ^1(\text{TT}) \) states from global analysis. Red: Singlet state. Blue: Triplet pair state. Inset: 2D pseudocolor (intensity) plot of TA spectra following excitation at zero time by hv = 2.1 eV. The transitions, along with vibronic progressions, are shown on each spectrum.
molecules are therefore labeled as BP-42 and BP-57, respectively (17). For comparison, the dihedral angle in BP0 is 37°; thus, BP0 ≡ BP-37. Theoretical analysis showed that the intertriplet electronic coupling decreases with increasing dihedral angle (17). The singlet fission time constants are $\tau_{SF} = 0.76$, 1.69, and 3.38 ps, and the corresponding intertriplet annihilation time constants are $\tau_I = 0.45, 1.6$, and 5.2 ns for BP-37, BP-42, and BP-57, respectively (17). Figure 4 shows the near-IR region of the $S_1$ (red) and $1^T(TT)$ (blue) ESA spectra for BP-42 (top) and BP-57 (bottom). We multiply the $1^T(TT)$ spectra by factors of 2.5 and 4.6 for BP-42 and BP-57, respectively, to normalize the $1^T(TT)$-$S_n$ peak intensity to the $S_1$-$S_n$ intensity in each case. For comparison, the normalization factor would be 1.25 for BP-37 in Fig. 3. Thus, relative to the $S_1$-$S_n$ transition, the $1^T(TT)$-$S_n$ transition strength is 80, 40, and 22% for BP-37, BP-42, and BP-57, respectively. This confirms the correlation between the $1^T(TT)$-$S_n$ ESA transition strength and the intertriplet electronic coupling in the $1^T(TT)$ state.

In all the pentacene dimers investigated here, the decay rate of the triplet pair state is also found to be strongly correlated with the extent of delocalization in the $1^T(TT)$ state, which is reflected in the $1^T(TT)$-$S_n$ transition strength. This is understood as the rate of $1^T(TT)$-$T_1$ annihilation is determined by the inter-pentacene electronic coupling strength, as addressed in detail elsewhere (11, 17).

The relative amplitudes of the 0-0 and 0-1 transitions allow us to estimate the Huang-Rhys factor ($S$) in each case and, thus, the relative shifts in the potential energy surfaces (PESs) involved. The Huang-Rhys factor is related to the offset ($\Delta Q_e$) in the equilibrium positions of the two PESs in an optical transition: $S = 0.5\alpha (\Delta Q_e)^2$, where $\alpha = \mu_0/h \mu$ is the reduced mass, and $\omega$ is the angular frequency of the vibration (55). In the harmonic oscillator and low-temperature approximation appropriate for the pentacene ring breathing mode at room temperature, the ratio in the Franck-Condon factors (and the ratio in peak intensities) between the 0-1 and 0-0 transition is equal to the Huang-Rhys factor (55). Thus, we obtain $S = 0.36 \pm 0.05$ and $0.45 \pm 0.05$ for the $S_1$-$S_3$ and $1^T(TT)$-$S_n$ transitions, respectively, from the near-IR ESA spectra for BP0 in Fig. 3. For comparison, we obtain from the optical absorption spectrum a value of $S = 0.55 \pm 0.05$ for the $S_2$-$S_n$ transition (11). For the pentacene ring breathing mode, we neglect the difference in equilibrium geometries between $S_n$ and $S_n'$ because they both likely come from the linear combination of $S_1$ in each pentacene chromophore. The spectroscopic results obtained above allow us to construct PESs for singlet fission in BP0. Although there are four possible arrangements of the PES from experimental $\Delta Q$ values, Fig. 5 shows the scenario that is more consistent with the expectation of increasing nuclear displacement with excitation energy. The offset in equilibrium positions of the $S_1$ PES and the $1^T(TT)$ PES ($\Delta Q_e \sim 0.081 \text{Å}$) is also consistent with theoretical results on the covalent dimer (11, 56). Although Fig. 5 is only an approximation given the uncertainties in the spectroscopic determination of Huang-Rhys factors, it represents the first estimation of $S$ as an average from experimental data.

**Distinct chemical property of the $1^T(TT)$ state**

The above spectroscopic analysis of singlet fission in BP0 provides evidence for a strong coupled triplet pair state, $1^T(TT)$, whose delocalized and localized characters are revealed in ESA in the near-IR and visible regions, respectively. Here, we show that the tightly bound triplet pair state exhibits chemical properties that are different from those of an individual triplet.

The inset in Fig. 1A shows the estimated values for the IP and EA for TIPS-Pc and [Fe8O4], respectively. These values are obtained from the
either the T1 or the S1 in pentacene to the [Fe8O4] cluster. Figure 1C approximates energy level diagram, we expect efficient electron transfer from the estimated IPs of S1 and T1 states from the excitation energies of TIPS-Pc.

Direct photoexcitation of the CT state, electron transfer can occur from T1 in pentacene to [Fe8O4]-BP0 (gray); see the Supplementary Materials for T1 spectra of the other molecules. Note that neither the isolated [Fe8O4]-nor the uncoupled pentacene molecules absorb light below ~1.75 eV. Excitation of isolated [Fe8O4] at higher photon energies results in completely different TA spectra (fig. S2). The ultrafast formation of T1 within experimental time resolution (~100 fs) from the selective excitation of CT indicates that the cluster and pentacene ligands are strongly electronically coupled. Supporting this conclusion, we found in a triplet sensitization experiment that the observable T1 signal from [Fe8O4]-Pc is an order of magnitude lower than that from TIPS-Pc (fig. S3).

The coupled T1-CT state features first-order decay kinetics well described by single-exponential decays (solid curves in Fig. 6B), with time constants of $\tau_{CT:T1} = 28 \pm 3$ ps and $16 \pm 2$ ps for [Fe8O4]-BP0 and [Fe8O4]-Pc, respectively. The simple first-order kinetics is reflected in both the decay in T1-like ESA signal (red dots) and the recovery in ground-state bleaching (blue dots) for [Fe8O4]-BP0 in Fig. 6B. The T1-CT decay constant is five orders of magnitude shorter than that of an individual T1 state in pentacene or bipentacene molecules (11). Because no fluorescence emission is observed for any of the cluster-pentacene complexes, we assign the fast decay in the T1-CT state to nonradiative recombination. Both CT across the pentacene-cluster interface and the presence of paramagnetic Fe atoms can couple to electron spins, thus facilitating recombination (61).

**Fig. 6.** TA reveals the strong coupling of CT state to T1. (A) TA spectra at 1 ps for [Fe8O4]-Pc (green) and [Fe8O4]-BP0 (red) upon CT excitation of 1.65 eV. The gray curve is the triplet spectrum of [Fe8O4]-BP0 from triplet sensitization. (B) Triplet decay dynamics for [Fe8O4]-Pc (green) and [Fe8O4]-BP0 (red and blue for ESA and ground-state bleaching, respectively). The solid curves are single-exponential fits with the indicated lifetimes ($\tau = 16 \pm 2$ ps for [Fe8O4]-Pc and $28 \pm 3$ ps for [Fe8O4]-BP0).

**Fig. 7.** TA spectra and dynamics of [Fe8O4]-BP0 under 2.1 eV excitation. (A) 2D pseudocolor plot of TA ($-\Delta T/T$, T, transmission) as a function of pump-probe delay ($\Delta t$) and probe photon energy. (B) TA spectra at $\Delta t = 0$ ps (red), 10 ps (blue), and 100 ps (green), along with T1 spectrum from sensitization (gray). (C) Singlet fission dynamics, as represented by S1 decay at 2.07 eV (red) or (TT) buildup at 2.36 eV (blue). (D) Comparison of $^{1}\text{TT}$ decay dynamics for [Fe8O4]-BP0 and BP0.
Unlike the strong coupling of individual T1 in pentacene or bipentacene to the CT state at their interfaces to [Fe₈O₄], we find that the triplet state in the tightly bound ¹(TT) in BP0 does not undergo CT to the electron accepting cluster. Figure 7A shows TA spectra for [Fe₈O₄]-BP0 as a function of pump-probe delay, following initial photoexcitation at hν = 2.1 eV. Figure 7B shows horizontal cuts at selected pump-probe delays (Δt = 0, 10, and 100 ps), along with a T1 spectrum obtained from sensitization of [Fe₈O₄]-BP0. At this excitation photon energy, BP0 is known to undergo efficient singlet fission (11), and the results for [Fe₈O₄]-BP0 are nearly identical to those in BP0. Initially (Δt = 0 ps; red spectrum in Fig. 7B), the TA spectrum is that of S1 characterized by the broad ESA in the visible region and a vibronically resolved ESA in the near-IR region. The singlet exciton decay and triplet rise in [Fe₈O₄]-BP0 are both characterized by a single-exponential lifetime of τ_S = 0.55 ± 0.02 ps, which is slightly shorter than the corresponding process in BP0 (τ_SF = 0.76 ps) (11). Figure 7D compares the ¹(TT) decay dynamics in [Fe₈O₄]-BP0, as monitored by the decays of ESA signals attributed to both triplet (2.36 eV, red) and singlet (0.97 eV, green) characters. For comparison, we also show in Fig. 7D the ¹(TT) decay dynamics in BP0 (2.36 eV, blue). The three decay traces are superimposable. The data for [Fe₈O₄]-BP0 are well described by a single-exponential decay with a time constant of τTT = 0.42 ± 0.03 ns, which is, within experimental uncertainty, identical to that of BP0. In stark contrast to the efficient CT from an individual T1 state in [Fe₈O₄]-Pc, there is no measurable CT from the tightly bound ¹(TT) state in [Fe₈O₄]-BP0.

In summary, using covalently linked pentacene dimers as model systems, we show evidence for a tightly bound triplet pair state, which reveals its delocalized ¹(TT) and localized T1 characters in the near-IR and visible ESA spectra, respectively. The near-IR ESA spectra can be assigned the ¹(TT)→S₄ transition, which is similar to the S₁→S₃ transition, with vibrational progression corresponding to the well-known aromatic ring breathing mode. The ¹(TT)→S₄ transition is an indicator of the intertriplet coupling strength; when a phenylene spacer is inserted between the pentacene moieties (BP1) or varies the angle between the pentacene moieties (BP45, BP90, and 1,2-BP) to decrease this coupling, we find that the ¹(TT)→S₄ ESA peak decreases. This is in contrast to the spectrum in the visible region, assigned to the ¹(T₁T₃)→¹(T₁T₃) transition present with similar intensities for all bipentacene molecules. Using an electron-accepting iron oxide molecular cluster [Fe₈O₄] linked to pentacene and bipentacene (BP0), we find that electron transfer to the cluster occurs efficiently from an individual T1 but not from the ¹(TT) state. Thus, the tightly bound ¹(TT) state exhibits a distinctly different chemical reactivity from that of an individual T1 state. A viable strategy to efficiently harvest triplets from intramolecular singlet fission is to control the intertriplet electronic coupling via molecular design.

MATERIALS AND METHODS

Synthesis

The synthesis of TIPS-Pc, BP0, and BP1 molecules (11) , BP0 with different dihedral angles (17), and the [Fe₈O₄] cluster (49) has been previously described. To install the pentacene-based ligands on [Fe₈O₄], we first deprotonated the pendent phenol group with an excess of sodium hydride in tetrahydrofuran (THF). The reaction mixture was filtered through a 0.2-μm syringe filter and added dropwise to a solution of [Fe₈O₄] in THF. We used a 1:1 stoichiometric ratio of the ligand to [Fe₈O₄] to prepare the monosubstituted clusters, which were purified by reversed-phase chromatography. Additional synthetic details and characterization data can be found in the Supplementary Materials.

Optical absorption

The TIPS-Pc, BP0, and BP1 samples were dissolved in dry toluene (with a concentration of <100 μM) and kept free from oxygen and moisture for optical measurements on a Shimadzu UV 1800 UV-Vis Spectrophotometer. Ultraviolet-visible (UV-Vis) absorption spectra of BP0 and BP1 (Fig. 1B) showed a slight red shift from that of TIPS-Pc but contained otherwise nearly identical vibronic features near the absorption threshold (S₀→S₁) (11). Solutions of [Fe₈O₄], [Fe₈O₄]-Pc, or [Fe₈O₄]-BP0 in chloroform were used for absorption measurements. Optical absorption spectra of [Fe₈O₄], [Fe₈O₄]-Pc, and [Fe₈O₄]-BP0 in Fig. 1C will be discussed later.

Transient absorption

To investigate singlet fission and triplet transfer, we used femtosecond TA (fs-TA) spectroscopy. The samples were dissolved in dry toluene and kept free from oxygen and moisture. The pump pulse came from an optical parametric amplifier (tunable from UV to the near-IR, 100-fs pulse width, 1 kHz rep rate). The probe pulse was a white-light supercontinuum (from 450 to 850 nm and from 850 to 1600 nm for the visible and near-IR range, respectively). The delay between pump and probe pulses was controlled by a translational stage with a delay time up to 3 ns. The detection consisted of a pair of multichannel detector arrays coupled to a high-speed data acquisition system (HELIOS, Ultrafast System Inc.). The sample solution was at room temperature during measurement. The nanosecond-microsecond TA measurements were carried out on the same setup as fs-TA with the same pump pulse. The probe pulse was a white-light supercontinuum (from 400 to 1600 nm) generated by a supercontinuum laser (Leukos). The laser pulse width was ≤1 ns at 2 kHz. The pump-probe delay was controlled electrically.

The triplet-sensitizing experiment was carried out on the same setup except for the fact that the white-light probe beams were generated by a picosecond laser and the pump-probe delay was controlled electrically. A mixture of a (bi)pentacene compound and an excess of anthracene was dissolved in toluene with the concentration of anthracene ~100× that of (bi)pentacene. Photoexcitation at 3.35 eV created singlets in anthracene, which underwent intersystem crossing to form triplets. The triplets in anthracene subsequently transferred to (bi)pentacene molecules via diffusional collisions on a time scale of 1 to 2 μs (see the Supplementary Materials).

SUPPLEMENTARY MATERIALS

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/3/7/e1700241/DC1

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