High-Symmetry Anthradithiophene Molecular Packing Motifs Promote Thermally Activated Singlet Fission

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ABSTRACT: When considering the optimal molecular packing to realize charge multiplication in organic photovoltaic materials, subtle changes in intermolecular charge transfer (CT) coupling can strongly modulate singlet fission. To understand why certain packing arrangements are more conducive to charge multiplication by triplet pair (TT) formation, we measure the diffraction-limited transient absorption (TA) response from four single-crystal functionalized derivatives of fluorinated anthradithiophene: diF R-ADT (R = TES, TSBS, TDMS, TBDMS). diF TES-ADT and diF TDMS-ADT both exhibit 2D brickwork packing structures, diF TSBS-ADT adopts a 1D sandwich-herringbone packing structure, and diF TBDMS-ADT exhibits a 1D twisted-columnar packing structure. When brickwork or twisted-columnar single crystals are resonantly probed parallel to their charge transfer (CT)-axis projections, the TA signal is dominated by a rising component on the picosecond time scale (rate \( k_{TT} \)), attributed to TT state population. When probed orthogonal to the CT-axis, we instead recover the falling TA kinetics of singlet state depletion at rate \( k_{A} \). The rising to falling rate ratio estimates the TT formation efficiency, \( \epsilon_{TT} = k_{TT}/k_{A} \) relative to exciton self-trapping. \( \epsilon_{TT} \) ranged from near 100% in diF TES-ADT to 84% in diF TDMS-ADT. Interestingly, diF TSBS-ADT crystals only manifest falling kinetics of CT-mediated self-trapping and singlet state depletion. Singlet fission is prohibitive in diF TSBS-ADT crystals owing to its lower symmetry sandwich-herringbone packing that leads to \( S_1 \) to CT-state energy separation that is \( \sim 3 \times \) larger than in other packings. Collectively, these results highlight optimal packing configurations that either enhance or completely suppress CT-mediated TT formation.

INTRODUCTION

Singlet fission, which can generate up to two electron–hole pairs per photon absorbed, has the potential to enhance the efficiency of organic photovoltaics (OPV).\(^1\)–\(^3\) In well-studied singlet fission materials like TIPS-pentacene, the exoergic conversion of one singlet (\( S_1 \)) excitation to a coupled triplet pair (\( TT \)) state is a fast, coherent process that ultimately generates up to two free triplets (\( T_1 \)).\(^3,5,6\) In other systems like tetracene and anthradithiophene (ADT), the \( S_1 \) to \( 2T_1 \) conversion is much slower owing to an endoergic thermal activation barrier. These endoergic singlet fission processes can conserve energy (\( E_{T_1} < 2E_{T_1} \)) and spin only through slower thermal or vibrationally mediated charge transfer (CT) scattering processes.\(^7\)–\(^10\) While the underlying mechanisms of endoergic singlet fission have been studied extensively, there are still open questions about how crystal structure promotes intermolecular interactions and maximizes singlet fission yield.\(^11\) In particular, it remains difficult to predict which packing motifs lead to strongly coupled CT states that can promote TT formation over competing processes, like excimer formation.\(^12\)–\(^14\) To determine which packing motifs are most conducive to endoergic singlet fission, we compare the optical spectra and transient absorption (TA) microscopy responses of four single-crystal functionalized derivatives of fluorinated anthradithiophene: diF R-ADT (R = TES, TSBS, TDMS, TBDMS).

Previous studies have established that diF TES-ADT undergoes singlet fission\(^15\)–\(^18\) but have yet to examine how ADT molecular packing can be modified to promote or...
suppress the formation of TT states. ADT derivatives all share
the same conjugated ring structure that forms the backbone
of the molecule with different side groups that attach off the core
of the molecule (Figure 1a). These different side groups
determine the molecular packing properties of each derivative,
which could further affect the rates of singlet fission by altering
intermolecular electronic and vibrational coupling. The precise
nature of intermolecular interactions has been shown to play a
significant role in singlet fission in many different materi-
als. The derivatives investigated in this study include
diF TES-ADT, diF TSBS-ADT, diF TDMS-ADT, and diF
TBDMS-ADT. As illustrated by the packing motif cartoons in
Figure 1a, diF TES-ADT (purple) and diF TDMS-ADT
(green) both exhibit two-dimensional brickwork packing
structures, diF TSBS-ADT (blue) has a one-dimensional
sandwich-herringbone packing structure, and diF TBDMS-
ADT (red) has a one-dimensional twisted-columnar packing
structure. The effects of molecular packing structure on the
optical properties of ADT crystals have been explored in
detail and are summarized in Figure 1a, which shows the
steady-state absorption and photoluminescence (PL, dashed
lines) spectra of the four different derivatives. Figure 1a shows
that molecular packing leads to distinct intermolecular
coupling as seen by a decreasing spectral Stokes shifts from
0.13 eV in the brickwork packing motifs to 0.058 eV in the
less-ordered sandwich-herringbone packing motif. When
the molecules are dilutely dissolved in dichlorobenzene (<0.001
mM), the intermolecular coupling vanishes and the solvation-
induced spectral Stokes shift is approximately the same for all
ADT derivatives and minimal (typically <5 nm). These
molecular packing properties make ADT derivatives ideally
suited for studying the energetic barrier and proceed to form TT states.31,32 Many
weakly endoergic systems, including anthracene, tetracene, and
diF TES-ADT, still efficiently undergo singlet fission.17,33–35
diF TES-ADT has previously been categorized as a weakly
endoergic system, with literature reporting the 2 × T1 energy,
2E_{TT} being between 25 and 60 meV higher than the S1 energy, E_{S1}.15,6 The Stokes shift, E_{Stokes}, in Figure 1a is used to estimate the
lowest singlet Frenkel energy (E_{S1} = 1/2E_{Stokes}), giving values that range from 2.19 eV in diF TES-ADT to 2.33 eV in
diF TSBS-ADT. Further supporting diF TES-ADT as a
moderately endoergic system, DFT calculations give 2E_{TT} =
2.20 eV, suggesting only a room temperature-range thermal
activation ~20–30 meV is required.

Studies of singlet fission materials pentacene, tetracene,
hexacene, and rubrene have previously used polarization-
dependent TA measurements to identify contributions from
singlet and triplet excitons.27–30 As in many such organic
crystals, the packing motifs of ADT crystals lead to
intermolecular CT states and strongly anisotropic electronic
and optical properties. The different transition dipole moments
within the ADT crystal structures are preferentially aligned
along different axes in real space (see Figure 2b), with the S0S1
transition dipole oriented at 90° relative to the long crystal
axis. Optical absorption in organic crystals is highly anisotropic
not only for the S0S1 transition, but also for transitions from
excited states.19 The anisotropic nature of these crystals can be
utilized to isolate different ESA transitions in TA measure-
ments, including the formation of stable TT states.

For singlet fission to occur, materials must meet the
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Compared to the faster, direct singlet fission mechanisms of
exoergic systems like TIPS-pentacene, endoergic systems like
ADT are characterized by slower, CT-mediated singlet fission
dynamics. A representative energy diagram of our system is
shown in Figure 1b (inset), which illustrates how our
experiment pumps the S0 state and then probes both singlet
(S0 → S1) and triplet pair (TT → S1) excited state transitions.
Using ultrafast TA microscopy on single diF TES-ADT
crystals, Figure 1b shows the decaying singlet ground state bleach (aqua line fit), roughly matching the triplet pair rising
ESA (orange line fit). We associate these matching rising and
falling curves as evidence of delayed TT formation that has
been previously reported in this material.15,6 Figure 1b further

Figure 1. Endoergic singlet fission in ADT derivatives. (a) Absorption (solid lines) and photoluminescence (dashed lines) spectra of four diF R-ADT derivatives and solution. The cartoons on right show the top-down crystal packing morphology obtained by functionalizing the diF R-ADT molecule with the R-groups shown. Polarization is 90° relative to the long crystal axis. (b) TA microscopy of a diF TES-ADT crystal gives falling kinetics using a 2.3 eV S0S1 probe (aqua line), but the matching rising kinetics are recovered when probed at 1.26 eV (yellow line), suggesting TT formation. While the solution-phase TA relaxes more slowly, it accelerates as concentration increases from 0.05 mM (gray curve) to 25 mM (black curve), suggestive of aggregation-induced singlet fission. (Inset) A basic energy-level diagram illustrates how the probe pulse energy was varied to resolve either falling singlet ground state bleach or rising TT ESA responses.
plots the $S_1$ relaxation dynamics of a diF TES-ADT crystal alongside diF TES-ADT solutions in dichlorobenzene at various concentrations. In the solution phase, the TA kinetic decays shown for diF TES-ADT are approximately invariant for all four chemical diF R-ADT functionalizations (see Supporting Information, Figure S1). The TA response of diF R-ADT in the most dilute solution decays most slowly without undergoing singlet fission because this system lacks the necessary intermolecular interactions. However, as the diF R-ADT concentration increases, the decay rate accelerates, suggesting that the singlet state is depleted faster by dimer-mediated TT state formation. This introductory result demonstrates the importance of intermolecular coupling to initiate the singlet fission process, which we will explore further by comparing diF R-ADT crystals of different packing motifs.

**METHODS**

Thin crystals of different crystal packing motifs were grown from diF R-ADT functionalized derivatives through a drop-cast method with controlled evaporation. Each functional substituent group on the ADT molecule is shown in Figure 1a and is synthesized to yield one of three distinct packing motifs that can best be described as brickwork (TES and TDMS groups), twisted-columnar (TBDMS group), and sandwich-herringbone (TSBS group). In molecular crystals, these substituents critically control packing and intermolecular coupling, as indicated by their differing Stokes shifts relative to solution, as highlighted in Figure 1a (dashed lines). Crystals were grown on fused substrates upon which a 45 nm layer of silver was deposited to enhance optical signals collected in a reflective geometry. A droplet of a dichlorobenzene solution of each ADT derivative was deposited on the substrate and grown under a slow solvent evaporation method at 3 °C in a covered flask. This procedure yielded well-defined crystals that have been characterized using XRD methods (see Supporting Information, Figure S2). Coupled with the observation of clear crystal boundaries seen in polarization-dependent microscopy (see Supporting Information, Figure S3), nearly single-crystal regions could be selectively excited using the diffraction-limited optical probes used throughout this study. A spatially resolved optical characterization of absorption, PL, and PLE spectra was done via an all-reflection optical microscope system with a 300–800 nm scanable Xe arc lamp source (Horiba Fluorolog). Polarization-dependent hyperspectral imaging was accomplished on an electron-multiplying CCD camera (Princeton Instruments ProEM). For all measurements throughout, the diF R-ADT crystals were kept under high-vacuum in an optical cryostat (Advanced Cryonics LT-003 M or Quantum Design OptiCool) to prevent photodegradation.

TA dynamics were investigated in these crystals using diffraction-limited TA microscopy. Ultrafast laser pump pulses were tuned to be resonant with the diF R-ADT $S_0$–$S_1$ transitions using either (1) ~180 fs pulses at 80 MHz from a second-harmonic generation system (SHG, APE Harmonix) pumped by a Coherent Ultra Chameleon or (2) 35–95 fs pulses at 10 kHz from a noncollinear optical parametric amplifier (NOPA, ORPHEUS-N, LightConversion). The high repetition rate measurements enabled measurement collection at the lowest photon fluxes, typically <$5 \times 10^{12}$ photons/cm$^2$, where contributions from effects like triplet–triplet annihilation were undetectable. Tunable optical probe pulses were generated from either (1) a broadly tunable Ti:sapphire output (NOPA, ORPHEUS-N, LightConversion) or (2) a second optical parametric amplifier (LightConversion, ORPHEUS-F). Furthermore, white light supercontinuum probes were used for collecting optical bleach signals near the $S_1$ transition and for all spectrally resolved TA microscopy.
To identify prominent ESA peaks, the probe energy was scanned from 1 to 1.45 eV. This near-IR energy probe range was selected to be approximately half the S$_0$–S$_1$ transition energy. This less-congested region of the spectrum enables clear identification of the polarization-sensitive singlet fission kinetics. Specifically, the primary ESA signals accessible with these probe energies were either singlet (S$_1$→S$_n$) or triplet pair-dominated (TT→S$_2$).

The TA microscopy method uses collinear pump–probe beamlines raster-scanned by a piezo-scanning mirror in a 4f-confocal geometry to direct the beamlines to the diF R-ADT samples held under a high-vacuum environment. Nearly diffraction-limited excitation is achieved with high-NA (>0.4), NIR-compatible objectives with cover glass correction (Olympus 50×IR and Beck 15× reflective objective). Pulse widths are optimized using cross-correlation measured by photoconduction at the sample position. The piezo-scanning beam and sample stages are used to align to a target crystal whose polarization-dependent spectra were measured by hyperspectral microscopy. The pump and probe polarizations are controlled via two λ/2 plates. All polarization angles are measured with respect to the long crystal axis, which corresponds to the b-axis in diF TES-ADT and diF TDMS-ADT, and the a-axis in diF TBDMS-ADT and diF TSBS-ADT. Unless specified, the pump polarization was selected to align to the S$_0$S$_1$ dipole projection along the single-crystal axis of each diF R-ADT crystal. The transient reflection TA microscopy signal is retrieved by TEM-cooled InGaAs photodetector and lock-in detection (Zurich HF-LI) at selected pump beam modulation rates between 1 and 500 kHz. High modulation rates were achieved with an acousto-optical beam deflection with a TeO$_2$ crystal. Since each transition has a different transition dipole orientation, transitions were isolated in TA microscopy measurements by varying the polarization of the probe beam. Solution-based TA measurements were taken in a separate conventional transmissive pump–probe setup with 1 MHz modulation.

### RESULTS

Figure 2a plots the PL and PLE spectra of diF TES-ADT single crystals. At lower temperatures (80 K), the PL emission is dominated by emission from a lower-lying peak at an energy of $2E_{T_1} - E_{vib}$. As shown in previous studies, this $2E_{T_1} - E_{vib}$ energy corresponds to emission from the TT state that is allowed through a Herzberg–Teller vibrational coupling mechanism. The polarization-dependent PLE spectra of diF TES-ADT crystals in Figure 2a are obtained using transmissive hyperspectral microscopy imaging for 1.93±0.04 eV emission. Interestingly, the S$_0$S$_2$ peak in the PLE spectra is the most strongly anisotropic, being minimized to almost extinction at 0°. However, the monomer S$_0$S$_2$ selection rules for both anti- and syn- configurations of diF R-ADT predict maximum absorbance occurs closer to 90°, conflicting with the observed maximal absorption at 0°. To resolve, Figure 2a inset shows the DFT-calculated CT-axis dipole is at nearly 90° in a top-down view of the brickwork structure of diF TES-ADT. Rather than following the monomer selection rules, the S$_0$S$_2$ transition polarization aligns to the CT-axis, suggesting...
absorption to an excited state with CT character. Strong CT-character of the $S_0S_2$ transition is further supported by the near-perfect alignment of the 2.99 eV $S_0S_2$ absorption peak across all four derivatives shown in Figure 1a. The CT-character of the $S_0$ absorption will be critical to understanding the polarization-dependent TT to $S_1$ ESA responses of diF R-ADT single crystallites shown later in Figure 3.

The hyperspectral microscopy images in Figure 2b show the strong $S_0S_2$ linear absorption anisotropy of nearly single crystals of diF TES-ADT. Figure 2b further plots time-resolved movie snapshots of a $\sim20$ $\mu$m wide crystal blow-up of a single crystal region with probe polarizations at $90^\circ$ (blue arrow) and $0^\circ$ (orange arrow). These spatially resolved TA microscopy maps highlight the uniform nature of the rising and falling relaxation kinetics of a diF TES-ADT crystal region. Comparing the 1 and 100 ps TA microscopy snapshots of the dynamics, we observe a spatially uniformly rising ESA response for $90^\circ$ probe polarization and a matching falling ESA response at $0^\circ$ probe polarization. Such dynamics are difficult to isolate in regions with overlapping domains or larger laser spot sizes because the rising and falling kinetics become convolved due to different dipole orientations. See our Supporting Information, Video V1 (0–800 ps range) for compiled TA microscopy maps that animate the matching rising and falling dynamics summarized in Figure 2b.

Consistently, the most distinct separation of rising and falling TA microscopy kinetics are observed for probe polarizations aligned either parallel or perpendicular to the crystal axes. Figure 2c further includes diF TES-ADT kinetics taken at $30^\circ$ and $60^\circ$ intermediate probe polarizations to demonstrate these dynamics are composed of weighted contributions from the kinetic rate processes shown for $0^\circ$ and $90^\circ$. As a result, all other TA pump–probe dynamics are plotted only for probe beams that are polarized either parallel ($0^\circ$) or perpendicular ($90^\circ$) to the long axis of the crystals.

For each of the four diF R-ADT packings motifs shown, Figure 3a compares the normalized TA signals taken at $0^\circ$ (blue lines) and $90^\circ$ (orange lines) probe polarizations. In Figure 3ai and ii, the brickwork-packed structures (diF TES-ADT) show prominent exponential rises for $90^\circ$ probe polarizations, oriented perpendicular to the long axis of the crystal. Conversely, for $0^\circ$ probe polarization, the rising kinetics are replaced by matching falling kinetics. This $\sim76$ ps matching rising and falling time constant is associated with the depletion of the $S_1$ exciton to form a correlated triplet pair state, TT. The linear absorption spectrum for the twisted-columnar packing structure, diF TDMS-ADT, in Figure 3b shows that the maximum $S_0S_2$ resonance is orthogonal to that seen in brickwork packings. Consequently, the TT formation rise was strongest when the probe beam was polarized along the long axis of the crystal ($0^\circ$, blue line). Unlike the brickwork packings, Figure 3a shows that the $\sim37$ ps rise appears much faster than the $62$ ps fall. The fitting parameters in Table 1 account for this by using a biexponential rise to allow for contributions from both coherent vibrational and thermal excitations to overcome the endoergic activation barrier for TT formation. The polarization-dependent absorption spectra shown in Figure 3b support that the distinctive ESA rises shown in Figure 3a occur in diF TES-ADT, diF TDMS-ADT, and diF TDDBS-ADT at whichever polarizations match the dipole of the final $S_1$ absorption.

Most interestingly, the kinetics of the sandwich-herringbone packing structure, diF TSBS-ADT, in Figure 3a(iii) are entirely different. In particular, over a wide range of tunable probe energies (1.15–1.4 eV), only falling relaxation kinetics are extracted that are nearly invariant to probe polarization. Moreover, the diF TSBS-ADT relaxation kinetics are consistently faster than solution-phase kinetics (see Figure 1b). Collectively, this is consistent with a related work that suggest this packing favors exciton self-trapping and Frenkel emission over TT formation.

Table 1 gives the corresponding least-squares deconvolution fitting rates of the plots in Figure 3a for each probe polarization in these diffraction-limited TA microscopy measurements. This preliminary analysis explicitly ignores any subpicosecond dynamics beyond the instrument response function (IRF), and uses a least-squares biexponential ($\tau_1, \tau_2$) fit to capture the two dominant time scales over the 1.2 ns scanning range shown. These measurements employed low pump fluences of $3.5 \times 10^{12}$ photons/cm$^2$ and were in the strongly linear region of pump-power dependence. Later in Figure 6, further data will show how the faster dynamics ($<200$ fs) contribute to morphology-dependent singlet fission and exciton–exciton annihilation in diF R-ADT crystals. Kinetic analysis (see Thermally Activated Kinetics: Self-Trapped Exciton versus TT Formation) of each time constant permits assignment to the following polarization-dependent ESA processes: (i) $\tau_1$ rise: the TT $\rightarrow S_1$ ESA is associated with TT formation at the rate $k_{TT}^r$, (ii) $\tau_1$ fall: the $S_1 \rightarrow S_0$ ESA is primarily associated with the $S_1$ depletion to the TT state and self-trapped excitons (STE) at rate $k_{SS}$, (iii) $\tau_2$ fall: primarily associated with the $S_1$ and TT recombination rates. The rise and fall times of $\tau_1$ match between the Table 1 columns taken for opposite probe polarizations within the stated error. Singlet fission on very similar time scales has been observed in other endoergic systems including tetracene, where the roughly matching rising/falling rates are consistent with TT state formation through a thermally activated endoergic singlet fission barrier.

The TA microscopy kinetic traces plotted in Figure 4a show all four possible pump and probe polarization permutations labeled as $\theta_{pump}/\theta_{probe}$. While the probe-beam polarization impact is strong, the ESA kinetic rates are only weakly impacted by changing the pump polarization. The ESA polarization-dependent dynamics are determined by the probe polarization and the coupling strength between its electric field and the available transition dipole moments. The transition dipole moments and orientations in diF R-ADT crystals were investigated using DFT dimer calculations summarized in Figure 4b and are further described in Supporting Information, Table S2 and in J. Van Schenck et al. The DFT-calculated Frenkel (purple dashed arrow) and

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| Table 1. Summary of the Falling (↓) and Rising (↑) $\tau_1$ and $\tau_2$ Time Constants of Probe-Polarization-Dependent TA$^a$ |
|---------------------------------------------------|
| diF R-ADT; R = 0° | $\tau_1$ (ps) | $\tau_2$ (ps) | $\tau_1$ (ns) | $\tau_2$ (ns) |
| TES brickwork 1 | 76 ± 5 | 74 ± 3 | 3.1 ± 1.4 | 7.3 ± 1.2 |
| TDMS brickwork 2 | 78 ± 9 | 93 ± 9 | 3.5 ± 1.3 | 10 ± 2 |
| TDDBS twisted-columnar | 91, 65 | 62 ± 5 | 3.0 ± 0.3 | 2.7 ± 0.9 |
| TSBS sandwich-herringbone | 64 ± 21 | 49 ± 31 | 3.5 ± 0.6 | 0.47 ± 0.07 |

$^a$The nearly matching rise/fall times in three of the four diF R-ADT crystal packings are associated with CT-mediated TT state formation, a precursor for singlet fission.
the strongest CT (red dashed arrow) dipole orientations are plotted in Figure 4b relative to the crystal plane of light incidence. DFT calculations for diF TBDMS-ADT give two similar CT-axis orientations due to its twisted-columnar structure. To compare the calculated dipole orientations against experimental predictions, Figure 4b shows that the solid green arrow aligns to the polarization that maximizes the S0S2 absorption. This calculated and observed transition dipole moment is always along the axis of the strongest intermolecular coupling in each ADT packing morphology. The approximate agreement of the calculated strongest CT-state dipole orientation and that of the S0S2 absorption peak suggests that this ∼2.99 eV transition is borrowing oscillator strength from the CT-state transition. We further observe the CT-mixed S0S2 transition dipole orientations (obtained from the polarization-dependent optical absorption spectra) and the probe polarizations of the rising TT-formation dynamics (obtained from polarization-dependent ESA measurements) match across all packing morphologies. The observed probe-beam polarization dependencies also align very closely with the calculated transition dipole moments predicted by TD-DFT.

Further analysis of the ESA dynamics shown in Figure 4a is possible by noting that falling TA kinetics at one polarization likely still contribute somewhat to the rising kinetics taken at the opposite probe polarization. As described in the Supporting Information, section S7, the weaker falling contribution to strongly rising kinetics may be approximately removed by curve subtraction. Figure 4a shows a dashed line that fits the falling kinetic data multiplied by a probe-beam anisotropy factor, Aπ, and subtracted from the larger rising kinetic curve. Nonetheless, fitting the resulting rises of these subtracted curves produces lifetimes in Figure 4a that are only ∼3 ps longer than the as-measured exponential rise for diF TES-ADT. Except for diF TSBS-ADT, the rising formation kinetics overwhelm any convolved falling kinetics such that subtractive analysis shown by the dashed lines in Figure 4a is unnecessary. In Figure 4a(iii), the same procedure for diF TSBS-ADT shows the 90° probe dynamics decay faster than the 0° probe data. After 66 ps, the diF TSBS-ADT electrons and lattice are fully thermalized with the surroundings, and so the residual S1 relaxation is seen only for the dipole-aligned 0° probe. The ideal probe energies for each packing structure were selected using spectrally resolved TA microscopy. Specifically, the peaks of maximum ESA response were identified by taking a white-light supercontinuum probe over a 1.0 to 1.56 eV range, as shown in Figure 5b.

In summary, three of the four diF R-ADT derivatives examined gave strongly rising TA microscopy dynamics.
suggesting TT formation with an average rate, \( k_{TT} = (79 \pm 4 \text{ ps}^{-1}) \) or 12.7 ns\(^{-1}\) for a constant pump fluence of \( 3.5 \times 10^{12} \) photons/cm\(^2\). When the probe polarization is changed by 90°, complementary falling dynamics with an average rate of \( k_A = (74 \pm 6 \text{ ps}^{-1}) \) or 13.5 ns\(^{-1}\) are observed. Showing both the forward and reverse rates strongly contribute to \( k_{TT} \). Supporting Information, Figure S6 shows the observed diF R-ADT rise times varied from 12 to 174 ps as the laser fluence increased \( \sim 50x \). Such stable, long-lived TT states are consistent with lifetimes previously reported in other diF R-ADT systems.\(^{15,16,24}\) Depending on the time scale, this TT notation may refer to coupled dipole states (\( T_1T_1 \)) or a \( T_1-T_1 \) state, signifying diffusion of the triplet pair.\(^{8,36}\) As this work primarily focuses on which packing motifs facilitate vibrationally mediated TT state formation, it does not attempt to distinguish between these various forms of triplet pair correlations.
Table 2. Summary of Relevant Energy States Also Diagrammed in Figure 5c

| (diF R-ADT) R = | TES (brickwork 1) | TDMS (brickwork 2) | TDBMS (twisted-columnar) | TSBS (sandwich-herringbone) |
|----------------|------------------|------------------|---------------------------|-----------------------------|
| S0S1 Abs (eV)  | 2.25             | 2.26             | 2.25                      | 2.36                        |
| S0S2 Abs (eV)  | 2.98             | 2.98             | 2.98                      | 2.98                        |
| Stokes shift (eV) | 0.13             | 0.13             | 0.14                      | 0.05                        |
| S1 Frenkel (eV) | 2.19             | 2.20             | 2.18                      | 2.33                        |
| E<sub>CT</sub> TT [0-1] peak (eV) | 2.03             | 2.04             | 2.04                      | 2.19                        |
| E<sub>CT</sub> [eV] [dimer index] | 2.032 [1–2]      | -                | 1.954 [1–3]               | 2.04 [2–3]               |
| ΔE = E<sub>CT</sub> - E<sub>ES</sub> (eV) | 0.16             | -                | 0.22                      | 0.29                        |
| E<sub>TT</sub> - E<sub>CT</sub> (eV) | 0.02             | -                | 0.07                      | 0.16                        |

Values are experimentally determined, except for CT energies, which were DFT calculated for the optimal crystal dimer index labeled.24

### DISCUSSION

I. Isolating CT-Mediated Singlet Fission Dynamics Using Probe Polarization and Energy Selectivity.

Isolating the rising kinetics associated with TT state formation requires careful probe energy and polarization selection. Across the visible probe energy range, diF R-ADT TA kinetics are dominated by overlapping ESA transitions, making any quantitative rate analysis challenging. While this effect is mitigated by using near-IR probes, Figure 5b shows the singlet-character of the TT state means that the TT → S1 is often nearly degenerate with S1 → S0 ESA peaks. The TA microscopy spectra in Figure 5b plot probe energies from 1.05 to 1.35 eV and show that 1.26 eV is the optimal spectral energy to isolate the rising TT formation in diF TES-ADT.

Comparing the TA microscopy spectral peak widths shown in Figure 5b, the diF TES-ADT ESA peaks are ~1.8x broader than the diF TSBS-ADT ESA peaks. The dashed Gaussian lines in Figure 5b suggest diF TES-ADT has S1 → S0 transitions that are convoluted with the TT → S1 ESA peaks. The diF TSBS-ADT packing ESA peak widths are narrower, as TT state formation is not expected. The kinetic trace of diF TES-ADT probed at the E<sub>pr</sub> = ~1.15 eV ESA resonance is shown in Figure 7b. Here, the relaxation at a 90° probe polarization is slower than at 0° because the falling singlet and rising triplet pair kinetics strongly overlap at the E<sub>pr</sub> = ~1.15 eV ESA peak in Figure 5b. Upon subtracting the two curves, the red-trace in Figure 7b shows the rising dynamics successfully recover the same 77 ps rise originally observed at the optimal 1.26 eV probe energy. Contributions from singlet depletion kinetics are minimal near the E<sub>pr</sub> = 1.26 eV ESA peak, as it requires an unlikely S1 to S2 (ν = 3) ESA transition. Thus, the strongly rising triplet pair ESA dynamics are best resolved at the 1.26 eV probe energy.

For molecular crystals of diF R-ADT, probe polarization is also critical for optimal isolation of the TT formation kinetics. Specifically, the diffraction-limited TA microscopy measurements use probe-beam polarizations either parallel or orthogonal to the CT-dipole axis projections. The linear absorption and PLE spectra in Figures 1a, 3b, and 5a demonstrate the significance of polarization selection. Figure 5a compares the polarization-dependent absorption spectra of brickwork-packed diF TES-ADT (top) and sandwich-herringbone-packed diF TSBS-ADT (bottom). In brickwork-packed diF TES-ADT, Figure 5a shows both the S0S1 and S0S2 dipole transitions are roughly maximized at 90°. By contrast, in diF TSBS-ADT, the S0S2 dipole transition is maximized at 0° (blue line), while the S0S1 absorption peak is maximal near 90° (orange line). This suggests the S0S2 peak is maximized for polarizations matching the calculated CT-axis dipole projection shown in Figure 4b,c. To best observe the strongly rising TT formation kinetics in diF TES-ADT, a probe energy of 1.26 eV and 90° polarization along the short-axis of the crystal is desired.

Table 2 and Figure 1a show that while Frenkel exciton energies and Stokes shifts differ significantly between derivatives, all four derivatives have an identical S0S1 energy at 2.99 eV. This suggests that the S0 and S1 bands are nearly vertically aligned, as the transition is strongly mixed with CT-states in all packings. The S0S2 transition drawn in Figure 5c also has a much larger polarization anisotropy factor than the S0S1 transition. Specifically, Figures 3b and 5a show that the S0S2 peak disappears almost entirely whenever the polarization is orthogonal to the molecular crystal CT-axis projection. As illustrated in Figure 5c with the TT → S2 ESA transition arrow, this CT-mixing property of the S2 state is critical to understanding why single-crystal diF R-ADT ESA kinetics are dominated by a rising ESA signal when resonantly probed along the CT-axis projection.

The S<sub>1</sub> depletion kinetics in all four diF R-ADT derivatives relax at a similar rate (k<sub>A</sub>), which is far faster than observed in solution-phase (see Figure 1b). TA microscopy on concentrated solutions and amorphous regions generally cannot clearly isolate TT formation kinetics, as the rising TT kinetics cannot be deconvoluted from competing falling S<sub>1</sub> depletion kinetics. All four derivatives share a similar energetic CT-barrier depicted in Figure 5c, even if TT formation is negligible, as seen for diF TSBS-ADT. This CT-barrier is surmounted with the help of thermal and optically excited vibrational modes to ultimately generate TT states and self-trapped exciton (STE) states. The energetic placement of the TT state relative to the lowest CT-state helps predict the branching ratio between STE versus TT formation using the rate conservation equation, k<sub>A</sub> = k<sub>TT</sub> + k<sub>STE</sub>. The efficiency for TT formation over STE trapping is then calculated by the rate ratio, E<sub>TT</sub> = k<sub>TT</sub>/k<sub>STE</sub>.

Using the energies in Table 2, Figure 5c postulates how the excited state energy landscapes may differ between each diF R-ADT packing motif. The relaxed TT state is generally accepted as lying approximately one vibronic mode below the 2E<sub>T1</sub> energy. For example, in diF TES-ADT, our DFT calculation suggests 2E<sub>T1</sub> = 2.20 eV, and so E<sub>TT</sub> ≈ 2E<sub>T1</sub> - E<sub>esb</sub> or 2.03 eV. Experimentally, this TT energy is supported by the temperature-dependent Herzberg–Teller brightened emission peaks highlighted earlier in Figure 2a and documented in recent studies.15,16,30 In Figure 5c, the CT-state energies are drawn in red to represent an electron–hole pair self-trapped between two diF R-ADT crystal sites. These CT-state values are DFT calculated for the dimer-index that gives the coupling of the S0S2 peak. Using Supporting Information, Table S1 for full details).24 Table 2 further estimates the energy barrier, ∆E as
$E_{S_1} - E_{CT}$ germane to the competition between TT and STE formation. Interestingly, $\Delta E$ increases as the crystal symmetry and $\pi$-stacking character diminishes, that is, $\Delta E$ scales with packing as $R = \text{TES} < \text{TDMS} < \text{TBDMS} \ll \text{TSBS}$. This pattern qualitatively agrees with the kinetics in Figures 3 and 4, showing the rising TT formation kinetic component diminishes as $\Delta E$ increases and STE formation becomes the more favorable energetic pathway.

II. Thermally Activated Kinetics: Self-Trapped Exciton versus TT Formation. While anthradithiophene is best classified as an endoergic singlet fission material, the energetic barrier between $S_1$ and $2 \times T_1$ can be surmounted by either thermal activation or faster vibrationally mediated routes. Figure 6a and b compare the ultrafast TA relaxation kinetics of diF TES-ADT to diF TSBS-ADT crystals at $0^\circ$ (blue circles) and $90^\circ$ (orange circles) probe polarizations. The instrument response (dashed line) results from a convolved resonant optical parametric amplifier (OPA) pump at 2.35 eV, and a TT $\rightarrow S_2$ resonant noncollinear optical parametric amplifier (NOPA) probe at $E_{pr} = 1.26$ eV. Both Figure 6a and b show dynamics on the order of 12 ps that are faster than the $\sim 76$ ps time scales reported in Table 1 because the pump photon flux is $\sim 15 \times$ larger at $9.4 \times 10^{13}$ photons/cm$^2$. This rate acceleration is induced by laser heating and the net reverse rate processes (e.g., singlet fusion and triplet–triplet annihilation), which have been previously observed in diF R-ADT materials.15

Over the 0 to 1.2 ps interval plotted in Figure 6a (before break), only the $90^\circ$ (orange circles), CT-axis-probed data shows a fast initial decay spike, $k_{\text{fast}} = (0.102\text{ ps})^{-1}$. This fast decay is largely absent at $0^\circ$ probe polarization (blue circles). As the observation of this fast initial spike requires selective probe-beam polarization along the CT-axis, we connect it with fast TT formation processes before the Frenkel exciton state has thermalized. Similar fast decays are prominent in other endoergic singlet fission materials, such as tetracene.37 Consistent with singlet fission being prohibited in the sandwich-herringbone packing, Figure 6b shows this $k_{\text{fast}}$ component is also absent for the diF TSBS-ADT crystal (blue circles).

On the picosecond time scale, the $90^\circ$ pump polarization kinetics (dark orange) in Figure 6a rise faster than the $0^\circ$ pump polarization kinetics (light orange). A monoexponential initial rise and fall is needed to fit the $0^\circ$ pump case, giving $k_{\text{TT}} = (12 \text{ ps})^{-1}$ and $k_\Lambda = (11 \text{ ps})^{-1}$. When pumped at $90^\circ$ along the CT-axis, the rising and falling kinetics of diF TES-ADT in Figure 6a require a biexponential fit with an additional 4.8 ps rise component that comprises $\sim 15\%$ of the overall rising amplitude. The modulated second rising rate, $k_\Lambda = (4.8 \text{ ps})^{-1}$ is further supported by the weak beating fitted in the $90^\circ$ pump TA microscopy kinetics plotted in Figure 6a (red trace). While our 72 fs instrument response could not fully resolve the $\sim 33 \text{ fs}$ phonon beating mode shown, this beat period corresponds with the $E_{\text{ vib}} = 0.14 \text{ eV}$ C–C vibronic mode of diF R-ADT. As the mode is observed in the rising TT formation kinetics, it suggests that optically excited vibrational modes are mediating TT formation at a faster rate, $k_\Lambda$. Interestingly, when the pump is orthogonal to the CT-axis projection of diF TES-ADT, this faster rise is no longer observed (see $0^\circ$ pump, $90^\circ$ probe kinetics), suggesting that these coherent vibrational modes only help mediate TT formation along the CT-axis.

Compared to dilute solution measurements with negligible CT-mediated processes, the $S_1$ state populations ($n_{S_1}$) shown

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**Figure 6.** Ultrafast thermally and vibrationally CT-mediated dynamics. (a) TA microscopy of diF TES-ADT (1.26 eV probe) at $\theta_{\text{pump}}/\theta_{\text{probe}}$ polarizations labeled. When pumped at $0/90^\circ$, the signal rises with a $12.3 \text{ ps}$ time constant; however, when pumped along the CT-axis at $90^\circ/90^\circ$, the rise is instead biexponential. Weak vibrational beating at a $33 \text{ ps}$ period is best seen in the subtracted data [90°/90°] fitted in red. (b) diF TSBS-ADT dynamics (1.28 eV resonant probe) fall at similar rates $k_\Lambda$, but there are no rising kinetic contributions at any probe energy (1.15–1.38 eV) or polarization. (c) (top) Schematic of endoergic singlet fission in brickwork-packed diF TES-ADT. The equivalent STE process observed for diF TSBS-ADT packing. (bottom) shows TT formation kinetics in brickwork-packe diF TES-ADT. The equivalent STE process observed for diF TSBS-ADT packing.
in Figure 6a,b are depleted rapidly by vibrationally assisted ($k_p$) and thermally activated rates, $k_{th}$. The depletion of the S1 state on intermediate time scales of $\approx 0.5$ ps (post-thermalization) to several nanoseconds is approximated by the rate equation $rac{dn_{S1}}{dt} = \delta(t)n_0 - (k_{TT} + k_{STE})n_{S1}$, where $n_0$ is the thermalized singlet initial population. At higher photon flux, the contribution from optical vibrational modes becomes increasingly significant, and a second faster rate contribution, namely, $k_p = k_{TT}^{S1} + k_{STE}^{S1}$, results in a biexponential rise. The contribution of this faster singlet depletion rate process $k_p$ is small enough to be neglected, with its rising amplitude not exceeding 15%. On a longer nanosecond time scale, the radiative and nonradiative rates, $k_{S1} = k_t + k_{nr}$, deplete the S1 Frenkel exciton state. These dynamics are studied extensively elsewhere.\(^{24}\)

Since strongly rising kinetics in Figures 3–6 suggests the probe beam predominantly resolves the TT-state formation, a different kinetic model is required when the probe beam is polarized along the CT molecular crystal-axis projection. The triplet pair state population ($n_{TT}$) is approximated by the S1 coupled rate law, $\frac{dn_{TT}}{dt} = k_{TT}n_{S1} - (k_{HT} + k_{TT})n_{TT}$. $k_{HT}$ refers to the radiative relaxation of the S1 state in Br-TES-ADT by Herzberg–Teller PL emission.\(^{15,16,24,36}\) Full details and solutions to the coupled kinetic rate laws are found in Supporting Information, section S6. Applying this rate model of CT-mediated processes to all four derivatives gives the rates summarized in Table 3. The rate ratios also give the TT-formation efficiency, $\epsilon_{TT}$, relative to the competing self-trapped exciton (STE) pathways.

### III. Impact of diF R-ADT Packing Structure on Singlet Fission Efficiency

As diagrammed in Figure 6c, endoergic triplet pair formation can proceed by a concerted spin-conserved superexchange\(^{39,40}\) of electrons (solid) and holes (dashed) to generate the final triplet pair state drawn. In brickwork and twisted-columnar packings, the lowest CT-state and the relaxed TT states are approximately within $k_{3}T_{3}$ of each other, making the concerted exchange of charges to form TT states the most favorable scattering route. The opposite is true for the sandwich-herringbone CT-mediated processes that preferentially trap carriers.

Interestingly, Table 2 shows the calculated CT states of diF TES-ADT and diF TSBS-ADT packing are approximately degenerate, and so the higher Frenkel exciton energy of the sandwich-herringbone structure becomes a critical factor. Specifically, Figure 6c summarizes how diF TSBS-ADT preferentially scatters to the CT-state calculated to be $\Delta E \approx 110$ meV lower than the TT state. Large $\Delta E$ values should be more common in crystals and crystallites with mixed packing symmetries, like the sandwich-herringbone molecular packing, and serve to preferentially trap carriers that would otherwise proceed to form stable, long-lived triplet pairs.
Table 3 above summarizes the rate constants obtained for the various packing morphologies and lists the probe polarization conditions that maximize (or minimize) the rising or falling kinetic contributions. The rate constants associated with TT state formation were separated from the S$_1$ decay rates by simply rotating the probe-beam polarization 90°. The longer rate constants ($k_{S1}$, $k_{HT}$) in Table 3 are further compared with recently published room-temperature rates ($k_{PL,2,3}$) obtained from time-resolved single-photon counting (TCSPC) photoluminescence.24 Specifically, we observe rate equivalence between the two methods if $k_{PL} \approx k_{S1} + k_{HT}$.24 Unlike the TA microscopy rates, the TCSPC-measured rates cannot be separated based on polarization, and so the second TCSPC rate $k_{PL}$ reflects the sum of TT-relaxation and S$_1$-relaxation rates ($k_{HT} + k_{S1}$) obtained by polarization-dependent TA microscopy.

The rates in Table 3 were all taken at a low pump fluence of $3.5 \times 10^{13}$ photon/cm$^2$ and are approximated well by a monoexponential rise and fall. Only the kinetics of diF TBDMS-ADT had a % amplitude large enough (at ~40%) that both $k_{TT}$ and $k_p$ rates were necessary. As the rates associated with thermally activated singlet fission accelerate with pump fluence, the rate ratio or efficiency ($\varepsilon_{TT} = k_{TT}/k_p$) provides a more meaningful quantitative metric. $\varepsilon_{TT}$ is the efficiency of TT state formation relative to STE formation. Table 3 suggests the TT formation process in the diF TES-ADT brickwork packing is the most efficient at 95–100%. Compared to diF TES-ADT, the R-substituent side groups in Figure 1a show that diF TDMAS-ADT contains a comparatively bulkier side group. This suggests that the bulkier TDMAS substituent makes self-trapping more favorable, lowering $\varepsilon_{TT}$ to 84%, as summarized in Figure 7a.

Despite its favorable π-stacking configuration, with an $\varepsilon_{TT}$ of 88%, twisted-columnar-packed diF TBDMS-ADT is more likely to trap excitons than the brickwork-packed diF TES-ADT. This is likely owing to its lower-lying CT$_1$ state energy shown in Figure 5sci and Table 2. As one caveat, diF TBDMS-ADT consistently gave the fastest kinetic rises owing to an enhanced contribution from vibrationally mediated TT formation with rate $k_p$. This $k_p$ rate process is observed most strongly when both the exciton and vibrational modes are excited along the CT-axis (0° pump polarization). This suggests that vibrationally coherent exciton–phonon processes may mediate TT formation better in the twisted-columnar packing than in brickwork.

No clear evidence of TT state formation could be resolved in diF TSBS-ADT crystals at any energy or polarization investigated. The subtractive analysis shown in Figure 4c suggests that, if the contribution exists at all, it must be very small in amplitude compared to the self-trapping efficiency implied by the falling singlet dynamics. As one last consideration, the CT character of a material, or how much the CT and Frenkel excitons mix, can enhance singlet fission.41,42 diF TSBS-ADT retains the most Frenkel character, making it unfavorable for TT formation. diF TSBS-ADT has only two nearest neighbors, with a large slip distance between adjacent molecules in the “sandwich” part of the “sandwich-herringbone” packing. DFT predicts that diF TSBS-ADT has the smallest total exciton coupling of the three systems. As a result, in contrast to the other derivatives, the exciton delocalization in diF TSBS-ADT is suppressed, and so, rather than forming a TT state, the formation of highly localized Frenkel excitons and stable excimers are favored instead.24 In diF TES-ADT, large exciton couplings that lead to Frenkel-CT mixing have been attributed to its 2D nature, which provides more nearest-neighbor molecules. diF TBMDS-ADT also has a pronounced CT character, making it favorable to undergo singlet fission. However, rather than the number of nearest neighbors, its large π–π overlap between adjacent molecules that results in a smaller slip distance between molecules in the column is what leads to high interactions. This may help explain why the optically driven phonon rate, $k_p$, that mediates TT formation was ~8× stronger in amplitude than seen in the brickwork packing.

In summary, unlike brickwork or twisted-columnar packing motifs, the sandwich-herringbone packing shows little evidence of singlet fission or TT formation. This is supported by (i) the absence of any ultrafast $k_{list}$ dynamic rates in diF TSBS-ADT kinetics, suggesting any direct singlet fission pathways are inhibited by more localized excitations, (ii) the $S_1$ to lowest CT$_1$ state relaxation energy, $\Delta E$ is ~2× larger in diF TSBS-ADT, suppressing any CT-mediated TT state formation, (iii) brickwork packing geometries have both multidirectional translational symmetry in the incident plane of polarization and more nearest-neighbors than sandwich-herringbone packing (four in diF TES-ADT vs two in diF TSBS-ADT), making the latter packing less favorable for singlet fission.

## CONCLUSIONS

While diF R-ADT is best classified as an endoergonic singlet fission material, the energetic barrier between the $S_1$ and 2 × $T_1$ states is sufficiently small that both vibrational and thermally activated populations of the TT state are possible.37,43 The TT formation rate is limited by an $S_1$–CT energy barrier that depletes the singlet state with a thermally activated net bidirectional rate, $k_s$. Using polarization-dependent TA microscopy, the corresponding rate of TT state formation, $k_{TT}$, was isolated in three of the four single-crystal functionalized derivatives of diF R-ADT (R = TES, TDMS, TDMAS, and TSBS). Certain packing morphologies preferentially form self-trap excitons (STE) in excimer-like states at a rate calculated by $k_{STE} = k_s - k_{TT}$. For each molecular packing, the competing rising rate, $k_{TT}$, and falling rate, $k_s$, were separated using probe beam polarizations that were either parallel or orthogonal to the single-crystal CT-axis projection. As was shown in Figure 5b, the resonant probe window was further carefully selected to find a TT $\rightarrow$ $S_1$ ESA peak that was not convolved with $S_1$ $\rightarrow$ $S_1$ peaks. By combining careful probe-energy and probe-polarization selectivity, TA microscopy successfully isolated falling singlet and rising TT dynamics from the raw TA traces, without any need for error-prone postanalysis (e.g., by subtracting kinetics curves or pump–probe anisotropy). The ratio between rising and falling rates $\varepsilon_{TT} = k_{TT}/k_s = k_{TT}/(k_{TT} + k_{STE})$ is equivalent to the efficiency of TT formation over CT-mediated self-trapped excitation formation.

In three of the four molecular crystals studied, the TT formation efficiency was large ($\varepsilon_{TT} \approx 84$% to nearly 100%). Brickwork and twisted-columnar packings give efficiencies that decrease with the energy separation $\Delta E = E_{S1} - E_{CT}$ and the relative size of the $-R$ group substituent. diF TES-ADT and diF TDMAS-ADT both exhibit 2D brickwork packing structures and are dominated by a strongly rising TA response that accelerates from 153 to 12 ps with increasing photon flux. Likewise, when resonantly probed parallel to the charge transfer (CT)-axis, similar rising time scales were extracted
with the 1D twisted-columnar packing structure of diF TBDMS-ADT. The probe polarization-dependent dynamics suggest that the excited-state TT transition dipole orientation is perpendicular to the long axis of the diF TES-ADT and diF TDMS-ADT crystals and the short axis of the diF TBDMS-ADT crystal. At higher pump photon fluxes, the net TT formation could not be described by a single rate and required a biexponential rising fit. This second fast rate, $k$, is associated with coherent vibrationally assisted TT formation and was shown in Figure 6 to be most prominent when the vibrationally excited mode was pumped parallel to the crystal CT-axis. In diF TES-ADT, the $k_p$ rate contribution to TT formation was always small ($<\sim15\%$ of rising amplitude). Interestingly, in diF TBDMS-ADT, this coherent vibrational contribution to overall rising kinetics was at least 2× larger.

Lastly, for diF R-ADT molecules with a sandwich-herringbone motif, the dynamic signatures of singlet fission are almost completely suppressed. The geometry of diF TSBS-ADT leads to more localized excitons, which inhibits a coherent formation of the TT state. It also leads to a greater energetic barrier between $S_1$ and TT, favoring other relaxation channels. These attributes together suggest that the sandwich-herringbone packing motif is prohibitive for endoergic singlet fission. While diF TES-ADT remains a promising singlet fission organic semiconductor, we have shown that small changes to the crystal packing result in preferential exciton self-trapping rather than charge multiplication. Thus, the barrier for TT-formation is ordinately sensitive to subtle tweaks in the molecular packing. This motivates further work exploring how external temperature and applied fields may further maximize both TT formation and triplet extraction yields for diF R-ADT derivatives and related endoergic singlet fission systems.

### ASSOCIATED CONTENT

#### Supporting Information

The Supporting Information is available free of charge at https://pubs.acs.org/doi/10.1021/acs.jpcc.1c10977.

TA measurements of diF R-ADT derivatives in solution, XRD measurements of diF R-ADT crystals, polarization-dependent microscopy images of a diF TES-ADT crystal, TA microscopy experimental setup, relevant transition energy tables kinetic model details and schematic, and flux-dependent measurements (PDF).

A supplementary video of compiled polarization-dependent ultrafast microscopy maps of a diF TES-ADT crystal over time (MP4).

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#### Notes

The authors declare no competing financial interest.

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### REFERENCES

1. Smith, M. B.; Michl, J. Singlet Fission. Chem. Rev. 2010, 110 (11), 6891–6936.
2. Hanna, M. C.; Nozik, A. J. Solar Conversion Efficiency of Photovoltaic and Photoelectrolysis Cells with Carrier Multiplication Absorbers. J. Appl. Phys. 2006, 100 (7), 074510.
3. Xia, J.; Sanders, S. N.; Cheng, W.; Low, J. Z.; Liu, J.; Campos, L. M.; Sun, T. Singlet Fission: Progress and Prospects in Solar Cells. Adv. Mater. 2017, 29 (20), 1601652.
4. Rao, A.; Friend, R. H. Harnessing Singlet Exciton Fission to Break the Shockley–Queisser Limit. Nat. Rev. Mater. 2017, 2 (11), 1–12.
5. Bakulin, A. A.; Morgan, S. E.; Kehoe, T. B.; Wilson, M. W. B.; Chin, A. W.; Zigmantas, D.; Egorova, D. J.; Rao, A. Real-Time Observation of Multie excitonic States in Ultrastat Singlet Fission Using Coherent 2D Electronic Spectroscopy. Nat. Chem. 2016, 8 (1), 16–23.
6. Musser, A. J.; Liebel, M.; Schnedermann, C.; Wende, T.; Kehoe, T. B.; Rao, A.; Kukura, P. Evidence for Conical Intersection Dynamics Mediating Ultrastat Singlet Exciton Fission. Nat. Phys. 2015, 11 (4), 352–357.
7. Monahan, N.; Zhu, X. Y. Charge Transfer-Mediated Singlet Fission. Annu. Rev. Phys. Chem. 2015, 66, 601–618.
8. Miyata, K.; Conrad-Burton, F. S.; Geyer, F. L.; Zhu, X.-Y. Triplet Pair States in Singlet Fission. Chem. Rev. 2019, 119 (6), 4261–4292.
9. Ulrich, T.; Munz, D.; Guldi, D. M. Unconventional Singlet Fission Materials. Chem. Soc. Rev. 2021, 50 (5), 3485–3518.
10. Casanova, D. Theoretical Modeling of Singlet Fission. Chem. Rev. 2018, 118 (15), 7164–7207.
11. Johnson, J. C. Open Questions on the Photophysics of Ultrastat Singlet Fission. Commun. Chem. 2021, 4 (1), 1–3.
12. Huang, Y.; Buyanova, I. A.; Phansa, C.; Sandoval-Salinas, M. E.; Casanova, D.; Myers, W. K.; Greenham, N. C.; Rao, A.; Chen, W. M.; Puttisong, Y. Competition between Triplet Pair Formation and Excimer-like Recombination Controls Singlet Fission Yield. Cell Reports Phys. Sci. 2021, 2 (2), 100339.
13. Dover, C. B.; Gallaher, J. K.; Frazer, L.; Tapping, P. C.; Pettry, A. J.; Crossley, M. J.; Anthony, J. E.; Kee, T. W.; Schmidt, T. W. Endothermic Singlet Fission Is Hindered by Excimer Formation. Nat. Chem. 2018, 10 (3), 305–310.
14. Young, R. M.; Wasilewski, M. R. Mixed Electronic States in Molecular Dimers: Connecting Singlet Fission, Excimer Formation,
(32) Casillas, R.; Papadopoulos, I.; Ullrich, T.; Thiel, D.; Kunzmann, A.; Guldí, D. M. Molecular Insights and Concepts to Engineer Singlet Fission Energy Conversion Devices. Energy Environ. Sci. 2020, 13 (9), 2741−2804.

(33) Bae, Y. J.; Kang, G.; Mallikas, C. D.; Nelson, J. N.; Zhou, J.; Young, R. M.; Wu, Y.-L.; Van Duyne, R. P.; Schatz, G. C.; Wasielewski, M. R. Singlet Fission in 9,10-Bis(Phenylethynyl)-Anthracene Thin Films. J. Am. Chem. Soc. 2018, 140 (45), 15140−15144.

(34) Cruz, C. D.; Chronister, E. L.; Bardeen, C. J. Using Temperature Dependent Fluorescence to Evaluate Singlet Fission Pathways in Tetracene Single Crystals. J. Chem. Phys. 2020, 153 (23), 234504.

(35) Wu, T. C.; Thompson, N. J.; Congreve, D. N.; Hentsz, Y.; Yost, S. R.; Van Voorhis, T.; Baldo, M. A. Singlet Fission Efficiency in Tetracene-Based Organic Solar Cells. Appl. Phys. Lett. 2014, 104 (19), 193901.

(36) Musser, A. J.; Clark, J. Triplet-Pair States in Organic Semiconductors. Annu. Rev. Phys. Chem. 2019, 70, 323−351.

(37) Stern, H. L.; Cheminal, A.; Yost, S. R.; Broch, K.; Bayliss, S. L.; Chen, K.; Tabachynyk, M.; Thorley, K.; Greenham, N.; Hodgkiss, J. M.; Anthony, J.; Head-Gordon, M.; Musser, A. J.; Rao, A.; Friend, R. H. Vibrionically Coherent Ultrafast Triplet-Pair Formation and Subsequent Thermally Activated Dissociation Control Efficient Endothermic Singlet Fission. Nat. Chem. 2017, 9 (12), 1205−1212.

(38) Wong, C. Y.; Penwell, S. B.; Cotts, B. L.; Noriega, R. J.; Wu, H. G.; Ginsberg, N. S. Revealing Exciton Dynamics in a Small-Molecule Organic Semiconductor Film with Subdomain Transient Absorption Microscopy. J. Phys. Chem. C 2013, 117 (42), 22111−22122.

(39) Berkelbach, T. C.; Hybertsen, M. S.; Reichman, D. R. Microscopic Theory of Singlet Exciton Fission. I. General Formulation. J. Phys. Chem. 2013, 118 (11), 114102.

(40) Berkelbach, T. C.; Hybertsen, M. S.; Reichman, D. R. Microscopic Theory of Singlet Exciton Fission. II. Application to Pentacene Dimers and the Role of Supercrystallization. J. Phys. Chem. 2013, 118 (11), 114103.

(41) Wang, X.; Liu, X.; Cook, C.; Schatschneider, B.; Marom, N. On the Possibility of Singlet Fission in Crystalline Quatertrylene. J. Chem. Phys. 2019, 123 (10), 5890−5899.

(42) Wang, X.; Liu, X.; Tom, R.; Cook, C.; Schatschneider, B.; Marom, N. Phenylated Acene Derivatives as Candidates for Intermolecular Singlet Fission. J. Phys. Chem. C 2019, 123 (10), 5890−5899.

(43) Wang, G.; Zhang, C.; Liu, Z.; Wang, R.; Ma, H.; Wang, X.; Xiao, M. Singlet Fission Dynamics in Tetracene Single Crystals Probed by Polarization-Dependent Two-Dimensional Electronic Spectroscopy. J. Phys. Chem. A 2020, 124 (50), 10447−10456.