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**This is a pre print version of the following article:**

*Original Citation:*

*Availability:*

This version is available <http://hdl.handle.net/2318/1690197> since 2019-02-05T13:52:36Z

*Published version:*

DOI:10.1039/c7ta07381j

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# Ultra-fast spin-mixing in a diketopyrrolopyrrole monomer/fullerene blend charge transfer state

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Keywords: transient absorption spectroscopy, electron paramagnetic resonance,  
diketopyrrolopyrrole, spin-mixing, triplet, fullerene anion, organic photovoltaics

Transient absorption spectroscopy (TAS) and time-resolved electron paramagnetic resonance (TR-EPR) spectroscopy are used to examine a small diketopyrrolopyrrole-based molecule, 2,5-Bis(2-hexyldecyl)-3,6-di(thiophen-2-yl)-2,5-dihydropyrrolo[3,4-c]pyrrole-1,4-dione (TDPP), in bulk heterojunction blend films with the common fullerene derivatives [6,6]-phenyl-C61-butyric acid methyl ester (PC60BM) and [6,6]-phenyl-C71-butyric acid methyl ester (PC70BM). Since diketopyrrolopyrrole (DPP) is now one of the most common building blocks for small molecules and conjugated polymers designed for organic electronic applications, a detailed understanding of the photophysical behaviour for a simple DPP-based system in fullerene blends is required, in order to establish a foundation for spectroscopic investigations into more complex DPP-based molecules. A TDPP:PCBM charge transfer state

is observed to form on ultra-fast timescales following pulsed laser excitation. It subsequently undergoes ultra-fast spin-mixing and geminate recombination to form the TDPP triplet state. The presence of this CT state is directly observed via a clearly identifiable fullerene anion transient absorption spectral signature: one of the first times an intermolecular small molecule/fullerene charge transfer state has been directly observed spectroscopically. The presence of this TDPP:PCBM CT state and overall photophysical mechanism is confirmed using TR-EPR spectroscopy, which unambiguously shows that the TDPP triplet forms via a spin-mixing CT state mechanism.

## 1. Introduction

Diketopyrrolopyrrole (DPP) is one of the most commonly used moieties in organic electronics.<sup>[1]</sup> DPP combines excellent stability, considerable synthetic versatility, high absorption coefficient, and strong fluorescence. Furthermore, it has produced high efficiencies in a number of organic electronic devices, including organic photovoltaics, light emitting diodes, and transistors. A key advantage of DPP is that it can be employed in both small molecule and conjugated polymer applications: in bulk heterojunction polymer solar cells, for example, efficiencies of over 9 %<sup>[2]</sup> have been measured, while efficiencies of over 7 %<sup>[3]</sup> have been achieved with DPP-based small molecules.

Despite its popularity, few reports exist in the literature regarding small molecule DPP's fundamental photophysics in a bulk heterojunction environment with fullerene. The photophysical mechanisms present in an organic solar cell have a direct bearing on its efficiency. The short circuit current is directly proportional to the charge carrier density present in the device, while the open circuit voltage is correlated with the charge transfer (CT) state energy, and the fill factor is strongly affected by bimolecular recombination. Of particular concern is the potential formation of triplet states. If the triplet state energy – of the

donor or the fullerene – is lower than the energy of the CT state, then formation of a triplet state can occur instead of the desired separated charges, which creates an additional loss pathway and reduces the device efficiency. This has been observed in a number of conjugated polymer/fullerene blends, and has been speculated to occur via spin-mixing in the charge transfer state.<sup>[4]</sup>

An experimental technique that has proven extremely useful for investigating the photophysical behaviour of organic photovoltaic blends is transient absorption spectroscopy (TAS). TAS directly monitors the optical absorption of photogenerated transient species: for polymer:PCBM blends, these are usually charge carriers or triplet states. TAS provides information on the identity, yield and recombination of these transient species. It has been applied to numerous polymer:blend systems,<sup>[5]</sup> where the charge carrier decay kinetics typically exhibit a power law decay ( $\Delta OD \propto t^{-\alpha}$ ). This power law behaviour is consistent with models describing bimolecular recombination of dissociated charge carriers in the presence of an exponential distribution of localised (trapped) states,<sup>[6]</sup> and is very different from the behaviour expected of triplet states. Triplet state relaxation back to the ground state is a first order process (at low excited state concentrations), which manifests as a monoexponential decay in the transient absorption data.

An additional spectroscopic technique that is highly complementary to TAS is time-resolved electron paramagnetic resonance (TR-EPR) spectroscopy. Owing to its unique sensitivity to paramagnetic species, TR-EPR can detect and assign triplet states and discriminate them from charge transfer states and radical pairs. The shape of the TR-EPR spectra of organic triplets reports on the mechanism of formation of the paramagnetic species, as it is directly dependent on the difference in populations of the three triplet sublevels. A common feature of all mechanisms leading to triplet formation is spin-selectivity. This translates into distinctive

non-equilibrium population of spin sub-levels (a phenomenon known as spin polarization), which can be used to pinpoint energy transfer pathways and charge transfer processes.

Previous transient spectroscopic studies of DPP-based small molecules have concentrated on the pristine DPP molecule with ancillary conjugated rings (thiophenes, for example),<sup>[7, 8]</sup> or DPP as one component of a covalently bound triad system.<sup>[9, 10]</sup> Remarkable results have already been observed: for example, pristine TDPP (DPP with two ancillary thiophene rings) has been reported to show singlet fission in thin films.<sup>[11]</sup> However, to our knowledge, no time-resolved spectroscopic study has been performed on simple DPP monomers in a bulk heterojunction film blended with fullerene. Furthermore, previous TAS studies have typically examined spectral features in the visible region only.

It is the aim of this paper, therefore, to build on previous literature work by examining TDPP in bulk heterojunction blend films with fullerenes, with particular focus on transient absorption spectral features in the near-infrared region. Importantly, TR-EPR spectroscopy is also used to complement and validate the TAS results, providing insight into the photophysical mechanisms present. The motivation is to establish a baseline of photophysical behaviour that will assist spectroscopic investigations into more complex DPP-based molecules blended with fullerenes, in order to enhance the efficiency of organic electronic applications.

It was found in this paper that a TDPP:fullerene charge transfer state forms on ultrafast timescales (< 100 fs). This CT state undergoes spin-mixing and geminate recombination to form the TDPP triplet state, again on ultra-fast timescales. The CT state was clearly observed via a distinct fullerene anion transient absorption spectral signature with a monoexponential decay lifetime of 300 – 500 ps, depending on the identity of the fullerene. Furthermore, the

TAS shows a rise in the TDPP triplet population, peaking at approximately 30 – 40 ps. This rise in TDPP triplet population with the simultaneous decay of the fullerene anion is further evidence that the fullerene is part of a TDPP:fullerene CT state that is undergoing spin-mixing, followed by recombination to form the TDPP triplet. This interpretation was confirmed through the use of TR-EPR spectroscopy, which unequivocally shows that the TDPP triplet has been formed through a spin-mixing CT state mechanism.

## **2. Results and Discussion**

### **2.1. Steady state absorbance and fluorescence**

The pristine diketopyrrolopyrrole-based monomer (TDPP) film shows a maximum absorbance at 510 nm, with a shoulder observed at 490 and a smaller peak at 565 nm (**Figure 1**). The structure of TDPP is shown in the inset of Figure 1. Addition of fullerene causes an increase in the relative amplitude of the shoulder at 565 nm (**Figure S1a**, supporting information). For polymer/fullerene blends this observation is usually linked to an enhancement in crystallinity.<sup>[12]</sup> If this holds true for the TDPP:PCBM films as well, this implies that the presence of the fullerene molecules imposes some degree of ordering on the TDPP molecules. In turn this could suggest the formation of disordered aggregates in the pristine DPP film. This is supported by the dilute solution state absorbance spectra, which show this lowest energy band (shifted to 550 nm) to be the highest in relative amplitude.

The pristine TDPP films fluoresce strongly, with the fluorescence maximum occurring at 690 nm. The TDPP fluorescence is almost completely quenched by the addition of fullerene, with quenching rates of 97 % observed for both TDPP:PC<sub>60</sub>BM and TDPP:PC<sub>70</sub>BM (**Figure S1b**, supporting information). This typically suggests quenching of the donor exciton via electron or energy transfer to the fullerene.

## 2.2. Nanosecond-microsecond transient absorption spectroscopy

In order to determine the outcome of the fluorescence quenching observed in the TDPP blend films, microsecond TAS was carried out. The pristine TDPP, pumped at 500 nm, showed no signal at any probe wavelength on the timescales available (approximately 500 ns – 10 ms). In contrast, the TDPP:PC<sub>60</sub>BM film produced a strong signal when pumped at the same wavelength. The TA spectrum of TDPP:PC<sub>60</sub>BM is shown in **Figure 2a** with a strong, narrow band appearing at 880 nm and a second distinct peak at 780 nm. The kinetics probed at 880 nm show clear mono-exponential kinetics with a lifetime of  $(14 \pm 1) \mu\text{s}$  (**Figure 2b**). This lifetime was invariant at low excitation densities, but at high excitation densities second order process begin to become evident. Such mono-exponential behaviour indicates a first order process such as triplet decay or charge carrier geminate recombination. The PC<sub>60</sub>BM triplet and radical anion can be discounted as their respective spectral shapes and positions are well known.<sup>[13]</sup> Geminate recombination on timescales encompassing tens of microseconds is unlikely, and thus the most likely candidate for the 880 nm feature observed in the TA spectrum is the TDPP triplet. The loss of mono-exponential behaviour at higher excitation densities could therefore be assigned to triplet-triplet annihilation. However, it should also be pointed out that the DPP radical cation absorbs at 750 nm in solution<sup>[8]</sup> so the possibility that the 870 nm band is related to the DPP cation cannot be entirely ruled out solely on the basis of these data.

Confirmation that the 880 nm peak can be assigned to a TDPP species rather than a fullerene transient species was provided by measuring the TA spectrum of TDPP:PC<sub>70</sub>BM. This spectrum, reported in **Figure 2c**, was virtually identical to that of TDPP:PC<sub>60</sub>BM, clearly showing that both the 880 and 780 nm peaks can be assigned to a TDPP transient species (both the radical anion and triplet state absorbance of PC<sub>70</sub>BM are distinctly red-shifted compared to those of PC<sub>60</sub>BM<sup>[14, 15]</sup>). The TDPP:PC<sub>70</sub>BM decay dynamics, shown in **Figure**

**2d**, could be fitted to a mono-exponential decay with a lifetime of  $(20 \pm 1)$   $\mu\text{s}$ : a slightly longer lifetime than that observed for TDPP:PC<sub>60</sub>BM. Again, this is tentatively assigned to the TDPP triplet, but additional confirmation is required. It should be noted that a prior study by Karsten *et al.*<sup>[10]</sup> used triplet sensitisation to generate DPP triplets in a fullerene-DPP-fullerene triad in solution, where photoinduced absorption showed the resultant DPP triplet to have peaks at 860 and 775 nm – similar to those observed in this work. This close correlation, despite the differences in molecular structure and phase, is further evidence that the TDPP transient species observed here is the triplet state.

In order to confirm the assignment of the 880 nm peak to TDPP triplets, the decay kinetics were also measured under a pure oxygen atmosphere. It is expected that any triplets would be quenched by the triplet ground state of oxygen, thereby showing a shorter lifetime and smaller signal amplitude compared to the inert atmosphere. However, neither of these effects were observed: the kinetics were virtually identical under both argon and oxygen atmospheres for both blend films (**Figure 2e** and **2f**). This would then suggest that the 880 nm peak does indeed belong to the TDPP radical cation. However, it must be noted that the oxygen quenching mechanism can only take place if the TDPP triplet state is higher in energy than the first excited singlet state of oxygen. Since TDPP has been considered for singlet fission applications, this implies a relatively low-lying T<sub>1</sub> state. Hence, TD-DFT calculations were performed to estimate the energy of the T<sub>1</sub> state of TDPP, to assess whether or not triplet quenching by oxygen is in fact feasible in this system. B3LYP/6-31G(d) calculations do indeed predict a very low-lying T<sub>1</sub> state, at approximately 1 eV. This is close to the oxygen quenching threshold of 0.98 eV, and thus it can be concluded that oxygen quenching would be very difficult in this system. However, previous results for a fullerene-DPP-fullerene triad in solution show DPP triplets to be sensitive to oxygen.<sup>[9]</sup> It is possible, therefore, that the film morphology stabilises the triplet energy below that of the oxygen quenching threshold. It is

thus not possible to conclude from these TAS results whether the transient species being formed is the TDPP radical cation or triplet state. The microsecond lifetime of the decay dynamics and close correlation with related molecules in the literature,<sup>[10]</sup> however, are strongly suggestive of a triplet state.

If the 880 and 780 nm features observed in the TDPP/fullerene blend spectra are indeed from the TDPP triplet state, whereas no TDPP triplet is observed in a pristine TDPP film, the implication is that a simple intersystem crossing mechanism cannot be responsible for the formation of the TDPP triplet. In order to explore this further, a picosecond TAS study was performed.

### **2.3. Picosecond transient absorption spectroscopy**

The ps-TA spectrum of the pristine TDPP film shows two strong peaks at approximately 800 nm and 1600 nm (**Figure 3a**). At low excitation densities, both peaks decay mono-exponentially with the same lifetime of  $(150 \pm 10)$  ps, suggesting they arise from the same transient species (**Figure 3b**). The ultra-fast generation of these peaks, first order behaviour, and lifetime are consistent with a singlet state, and both bands can therefore be attributed to the TDPP singlet exciton. The loss of mono-exponential behaviour at higher excitation densities can then be assigned to singlet-singlet annihilation. The 800 nm band (or the edge of it, in visible region only studies) has been observed before in TAS studies of related DPP-based molecules and was also assigned to the  $S_1$  state.<sup>[7, 9, 10]</sup> Interestingly, there is also evidence of very weak, broad, longer-lived bands at 850 – 1100 nm that remain after the strong singlet peaks have decayed. Extrapolating the decay of these additional peaks to longer times shows that these signals will have completely decayed by 500 ns, and this explains their absence in the  $\mu$ s-TAS data. The identity of this feature is unclear: it is possible

that it can be assigned to short-lived triplets that form inefficiently via direct intersystem crossing.

The ps-TA spectrum of TDPP:PC<sub>60</sub>BM is shown in **Figure 3c**. The two TDPP singlet exciton peaks are evident at the earliest times, but they are accompanied by additional features at 870 nm and 1030 nm. The 1030 nm band has completely decayed by the end time resolution (6 ns), but the 870 nm band persists beyond this time and is likely the same feature observed in the  $\mu$ s-TAS: the TDPP triplet. Although the 870 nm band is immediately apparent at the earliest times (< 100 fs), at low excitation densities a growth in the signal amplitude is observed (see **Figure 3d**), with the maximum occurring at approximately 40 ps. This growth is not observed at higher excitation densities, suggesting that the precursor transient species may be quenched by higher order processes. The 1030 nm band position is consistent with the PC<sub>60</sub>BM anion, and at low excitation densities it decays with mono-exponential behaviour and a lifetime of  $(230 \pm 40)$  ps. This lifetime was determined using a probe wavelength of 1120 nm, on the red edge of the PCBM anion peak in order to avoid contamination from the longer lived 870 nm band. The mono-exponential decay of the PC<sub>60</sub>BM anion and hundreds of picosecond lifetime indicates that it is decaying via geminate recombination.

The assignment of the 1030 nm band to the PC<sub>60</sub>BM anion received further support when the ps-TA spectrum of TDPP:PC<sub>70</sub>BM was measured (**Figure 3e**). In the latter case, the 1030 nm band was absent: instead, a band was observed at approximately 1320 nm. The position of this band is consistent with literature values for the PC<sub>70</sub>BM anion.<sup>[15]</sup> Mono-exponential decay dynamics were observed again, with a lifetime of  $(420 \pm 80)$  ps (measured from the blue edge, at 1200 nm). This lifetime is longer than that observed for the PC<sub>60</sub>BM anion. As with the TDPP:PC<sub>60</sub>BM blend, the long lived TDPP triplet state was observed at 870 nm in the

TDPP:PC<sub>70</sub>BM film. The rise in population of the TDPP triplets is more apparent for TDPP:PC<sub>70</sub>BM (**Figure 3f**), with the peak population occurring at approximately 30 ps.

The ps-TAS results therefore suggest the following mechanism. Initial excitation of the TDPP produces TDPP singlet excitons, which then undergo an ultra-fast electron transfer to the fullerene, forming a TDPP/fullerene charge transfer state. Spin-mixing (which also must occur on ultra-fast timescales) then follows, producing the triplet CT state. The triplet CT state then undergoes geminate recombination to form the TDPP triplet (the lowest energy triplet state in the system). The fullerene anion peak can therefore most likely be attributed to the CT state rather than free fullerene anions, since the increase in the triplet population is occurring simultaneously with the fullerene anion decay. Furthermore, the rise in triplet population on the picosecond timescale is additional evidence that a spin-mixing CT state mechanism is active, since simple intersystem crossing is typically a much slower process.

#### 2.4. Time-resolved electron paramagnetic resonance spectroscopy

In order to provide further support for this mechanism, TR-EPR experiments were performed for TDPP:PC<sub>70</sub>BM and its constituent components. PC<sub>70</sub>BM in solution produces a signal consistent with triplets formed via ISC that fully decay in 2  $\mu$ s (**Figure 4a**). The signal presents a polarization pattern *EEAA* (where *E* is emission and *A* is enhanced absorption) and can be simulated with *ZFS* parameters  $|D| = 337$  MHz and  $E = 0$  MHz and zero-field relative populations  $P_x:P_y:P_z = 0:1:1$ , consistent with previously reported data for C<sub>70</sub> monoadducts.<sup>[16]</sup> The pristine TDPP film produces virtually no signal on the timescale of the technique, indicating a lack of either ISC triplets or photo-induced radicals (**Figure 4b**).

In contrast, TDPP:PC<sub>70</sub>BM (1:2 by weight, **Figure 4c and 4d**) produces a more complex, long-lived EPR signal. At early times (400-600 ns from the laser flash, **Figure 4c**), the EPR

signal for the blend film can be assigned to two components. On the basis of the control experiments on PC<sub>70</sub>BM in solution discussed above, the central features can be attributed to PC<sub>70</sub>BM triplets populated via ISC and it reflects the high molar ratio of PC<sub>70</sub>BM in the blend. On the basis of the ZFS and polarisation pattern respectively, the second broader component can be attributed to a triplet state formed on TDPP via radical pair recombination in high magnetic field. This is particularly apparent at long times of over  $\sim 2 \mu\text{s}$ , when the PC<sub>70</sub>BM triplet has decayed (Figure 4d). **Figure S2** (supporting information) reports the comparison of the two TR-EPR surfaces for PC<sub>70</sub>BM and TDPP:PC<sub>70</sub>BM (1:2 ratio) to illustrate the time dependence of the measured signals. The TR-EPR spectrum at later times presents polarization pattern *AEEAAE*, which is characteristic of selective population of the T<sub>0</sub> sublevel (S-T<sub>0</sub> mixing) by charge recombination in the singlet-born weakly coupled radical pair at all orientations due to the conservation of the spin angular momentum in the recombination process.<sup>[17]</sup> The polarization pattern is indicative of a positive ZFS parameter *D*, which from simulation can be estimated to be  $D = + 1550 \text{ MHz}$ , together with a  $|E| = 340 \text{ MHz}$ . The EPR results are therefore entirely consistent with the proposed mechanism involving a TDPP:PC<sub>70</sub>BM charge transfer state that undergoes ultra-fast spin-mixing and recombination to yield the TDPP triplet state.

### 3. Conclusions

Small molecule TDPP:fullerene bulk heterojunction blend films were examined with TAS and TR-EPR spectroscopy and transient absorption spectroscopy, with specific focus on transient absorption spectral features in the near-infrared region. This spectroscopic study establishes a baseline of photophysical behaviour that will create a foundation for spectroscopic investigations into more complex DPP-based molecules, commonly used in organic electronic applications.

It was directly observed that a TDPP:fullerene charge transfer state forms on ultrafast timescales, shorter than the instrument resolution ( $< 100$  fs). This CT state undergoes ultrafast spin-mixing and geminate recombination to form the TDPP triplet state, the lowest energy excited state in the system. The presence of the CT state was directly observed via a clearly identifiable fullerene anion transient absorption spectral signature with a monoexponential lifetime of approximately 300 – 500 ps (depending on the identity of the fullerene). This is one of the first times an intermolecular small molecule/fullerene charge transfer state has been observed spectroscopically. This presence of the TDPP:fullerene CT state and overall photophysical mechanism was confirmed using time-resolved EPR spectroscopy, which explicitly shows that the TDPP triplet has been formed through a spin-mixing CT state mechanism.

#### **4. Experimental section**

The TDPP was synthesised using a known literature procedure.<sup>[18]</sup> Thin films of TDPP, TDPP:PC<sub>60</sub>BM, and TDPP:PC<sub>70</sub>BM (1:2 by weight) were prepared via spin-coating chloroform/o-dichlorobenzene (47:3) solutions onto glass substrates.

*Nanosecond-microsecond TAS:* Microsecond TAS on the above film samples was recorded using laser pulses (6 ns, 532 nm, repetition rate 10 Hz) from a Nd:YAG laser (Spectra-Physics, INDI-40-10) with a pump wavelength of 510 nm, using pump intensities of 0.5 – 36  $\mu\text{J cm}^{-2}$ . The light output of a tungsten lamp (Bentham) was used as a probe and signals were recorded with Si and InGaAs photodiodes, housed in a preamplifier and an electronic filter (Costronics Electronics) connected to an oscilloscope and PC. Probe wavelengths were selected with a monochromator. The films were kept under an argon atmosphere to prevent film degradation, while an oxygen atmosphere was trialled to check for triplet quenching.

*Picosecond TAS:* Film samples under inert nitrogen atmosphere were excited with 510 nm, 2 – 12  $\mu\text{J cm}^{-2}$  pulses, generated by a commercially available optical parametric amplifier TOPAS (Light conversion) pumped by a Solstice Ti:Sapphire regenerative amplifier (Newport Ltd). Changes in the optical density of the films induced by the laser excitation were probed with a second broadband pulse (830–1450 nm) generated in a sapphire crystal. A HELIOS (Ultrafast systems) transient absorption spectrometer was used for recording the dynamics of the transient absorption spectra up to 6.5 ns with an average 200 fs instrument response function.

*Time-resolved EPR:* A Bruker E580 pulsed EPR spectrometer operating at  $\sim 9.6$  GHz and equipped with a Bruker ER4118X-MD5 dielectric resonator was used to record the TR-EPR spectra solution ( $\text{PC}_{70}\text{BM}$ , 100 $\mu\text{M}$ ) and thin film (DPP and  $\text{PC}_{70}\text{BM}/\text{DPP}$  blend). Spectra were recorded in direct detection mode without magnetic field modulation. Hence they show characteristic enhanced absorptive (A) and emissive (E) features, as indicated in the reported spectra. A Surelite broadband OPO system within the operating range 410–680 nm, pumped by a Surelite I-20 Q-switched Nd:YAG laser with 2nd and 3rd harmonic generators (20 Hz, pulse length: 5 ns) was used to achieve a pulsed laser excitation at an appropriate wavelength optimized on the signal intensity, with the energy at the sample approximately 10 mJ per pulse. A cryogen-free cryostat from Cryogenic ltd and a Lake Shore temperature controller (model 350) were used to cool the sample and maintain the temperature at 50 K. EPR samples were prepared as described above and sealed under vacuum in quartz tubes (outer diameter 5mm). TR-EPR spectra were simulated using the EasySpin toolbox<sup>[19]</sup> in MATLAB™ to extract ZFS parameters –  $D$  and  $E$  - and sublevel populations ( $P_x, P_y, P_z$ ). The simulation of the TR-EPR spectrum of  $\text{PC}_{70}\text{BM}$  in solution took as initial parameters literature data for  $\text{C}_{70}$  monoadducts.<sup>[16]</sup> The simulation of the DPP: $\text{PC}_{70}\text{BM}$  thin film considered the ZFS parameters

$D$  and  $E$  to be both positive. An isotropic  $g$  value equal to the free electron  $g$  value ( $g_x = g_y = g_z = 2.0023$ ) was used in all simulations.

## 5. Author contributions.

The TDPP was synthesised by AL, supervised by HB. Film samples were fabricated by NL and JS under the supervision of TMC. TAS was performed by NL, JS, and TMC, with the data analysed by NL and TMC. EPR was performed by ES and the data analysed by ES and CWMK. The paper was written by TMC with assistance from ES and CWMK. The study was conceived by TMC and supervised by TC, HB, and CWMK.

## Supporting Information

Supporting Information is available from the Wiley Online Library or from the author.

## Acknowledgements

TMC would like to acknowledge support from EPSRC project EP/N026411/1.

Received: ((will be filled in by the editorial staff))

Revised: ((will be filled in by the editorial staff))

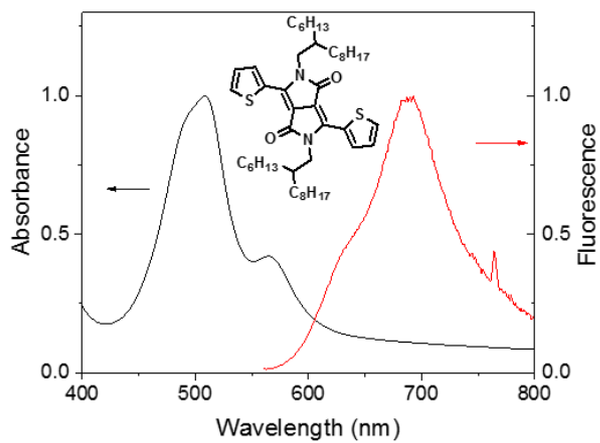
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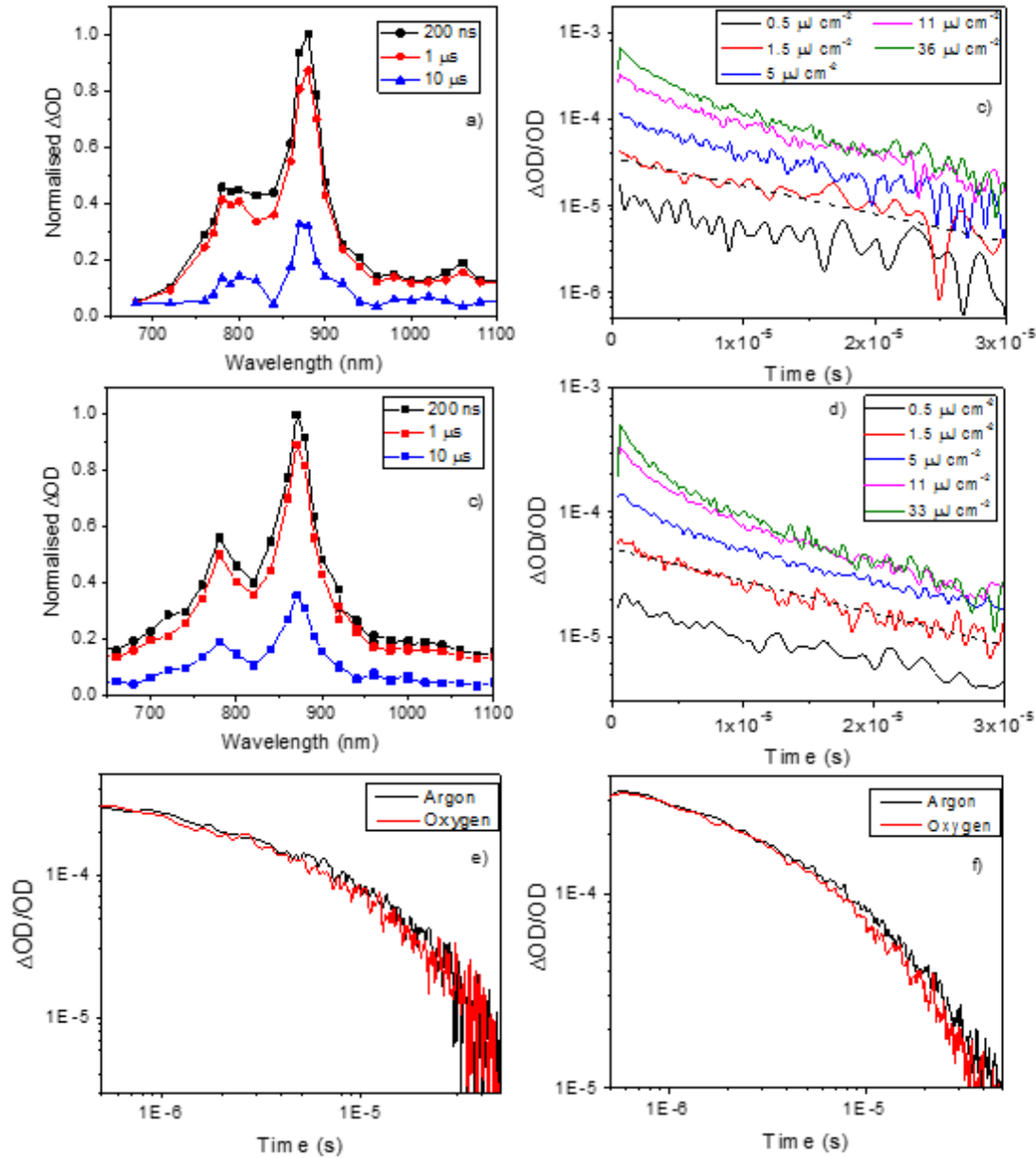
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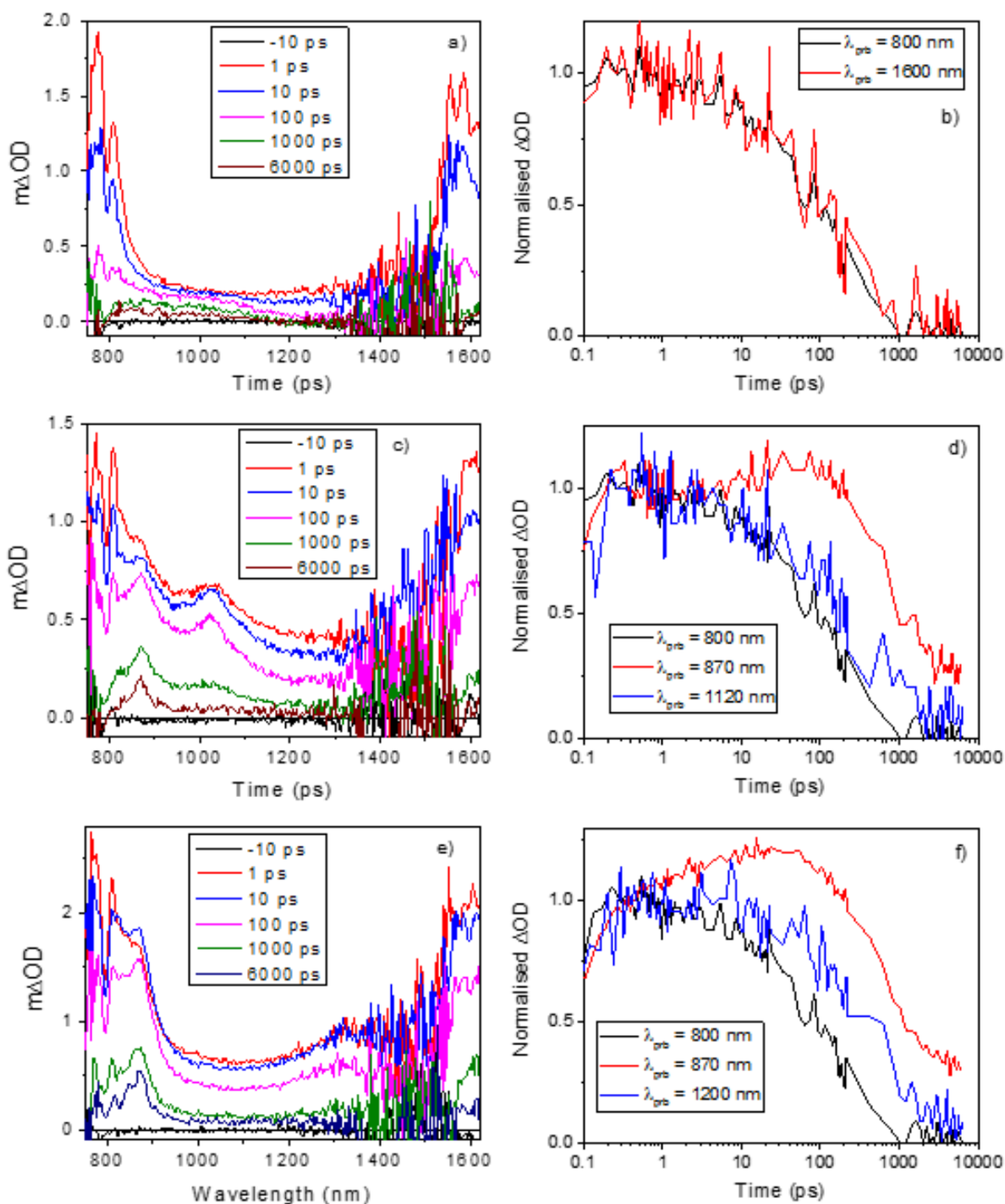
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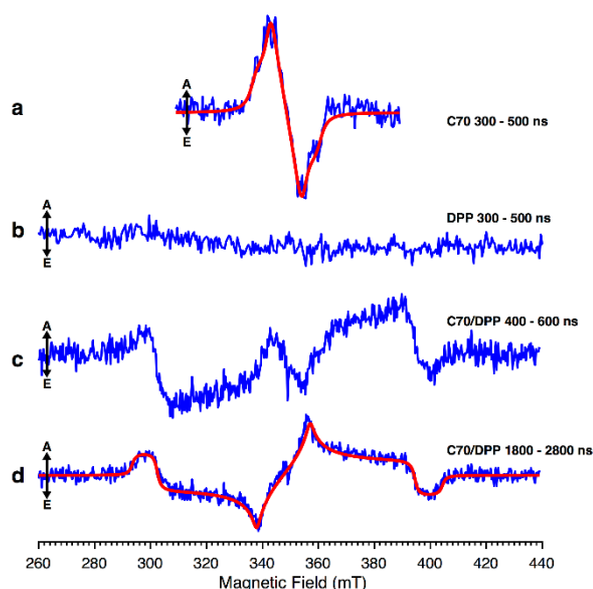
**Figure 1.** The normalised steady state absorbance and fluorescence spectra of pristine TDPP films, with the molecular structure shown in the inset.



**Figure 2.** The  $\mu$ s transient absorption spectra films of TDPP:PC<sub>60</sub>BM (a) and TDPP:PC<sub>70</sub>BM (b), both 1:2 by weight, using 510 nm and 36  $\mu$ J cm<sup>-2</sup> excitation. The corresponding decay kinetics as a function of excitation density for TDPP:PC<sub>60</sub>BM (c) and TDPP:PC<sub>70</sub>BM (d), both 1:2 by weight, using 510 nm excitation and a probe wavelength of 875 nm. The dependence on oxygen is shown for TDPP:PC<sub>60</sub>BM (e) and TDPP:PC<sub>70</sub>BM (f), both 1:2 by weight, using 510 nm, 11  $\mu$ J cm<sup>-2</sup> excitation and a probe wavelength of 875 nm.



**Figure 3.** The picosecond transient absorption spectra and kinetics as a function of probe wavelength for a pristine TDPP film (a, b), a TDPP:PC<sub>60</sub>BM film (1:2 by weight) (c, d), and a TDPP:PC<sub>70</sub>BM film (1:2 by weight) (e, f). A pump wavelength of 510 nm was used, and excitation densities of 2.7 and 7.3  $\mu\text{J cm}^{-2}$  were used for the kinetics and spectra respectively.



**Figure 4.** 50 K TR-EPR spectrum of PC<sub>70</sub>BM (blue line) in solution recorded after 400-600 ns from the laser pulse (a). The PC<sub>70</sub>BM spectrum can be simulated with ZFS parameters  $D = 337$  MHz,  $|E| = 0$  and relative populations  $P_x:P_y:P_z = 0:1:1$  (red line). 50 K TR-EPR spectrum of a pristine TDPP film recorded after 400-600 ns from the laser pulse (b). 50 K TR-EPR spectrum for the TDPP:PC<sub>70</sub>BM 1:2 blend film at early times (400-600 ns from the laser flash), where two contributions are distinguishable: a broad triplet and a narrower feature reminiscent of the PC<sub>70</sub>BM triplet (c). 50 K TR-EPR spectrum for the TDPP:PC<sub>70</sub>BM 1:2 blend film at later times (1800-2800 ns from the laser flash); only the recombination triplet on TDPP is visible (d). The TDPP:PC<sub>70</sub>BM spectrum can be simulated with ZFS parameters  $D = +1550$  MHz,  $|E| = 340$  MHz and assuming that only the  $T_0$  sublevel in high-field is populated by the charge transfer mechanism.

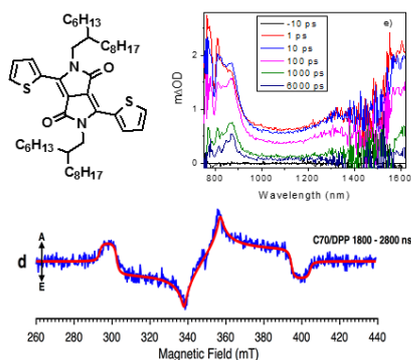
## Table of contents entry

Transient absorption spectroscopy and time-resolved electron paramagnetic resonance spectroscopy show a spin-mixing CT state in a small diketopyrrolopyrrole-based molecule blended with fullerene. This CT state is directly observed via a clear fullerene anion transient absorption band – a very rare observation. The resulting triplet states and mechanism were confirmed via EPR.

**Keywords:** transient absorption spectroscopy, electron paramagnetic resonance, diketopyrrolopyrrole, spin-mixing, triplet, fullerene anion, organic photovoltaics

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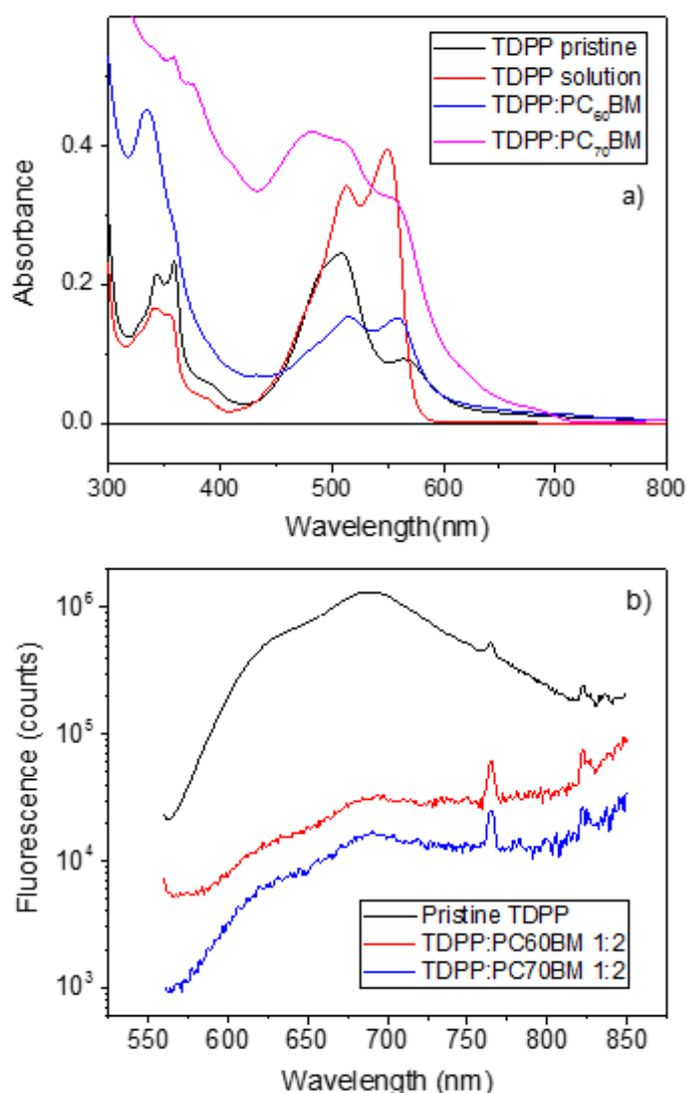
**Spin-mixing in a diketopyrrolopyrrole monomer / fullerene blend charge transfer state: a transient absorption spectroscopy and time-resolved electron paramagnetic resonance study**



## Supporting Information

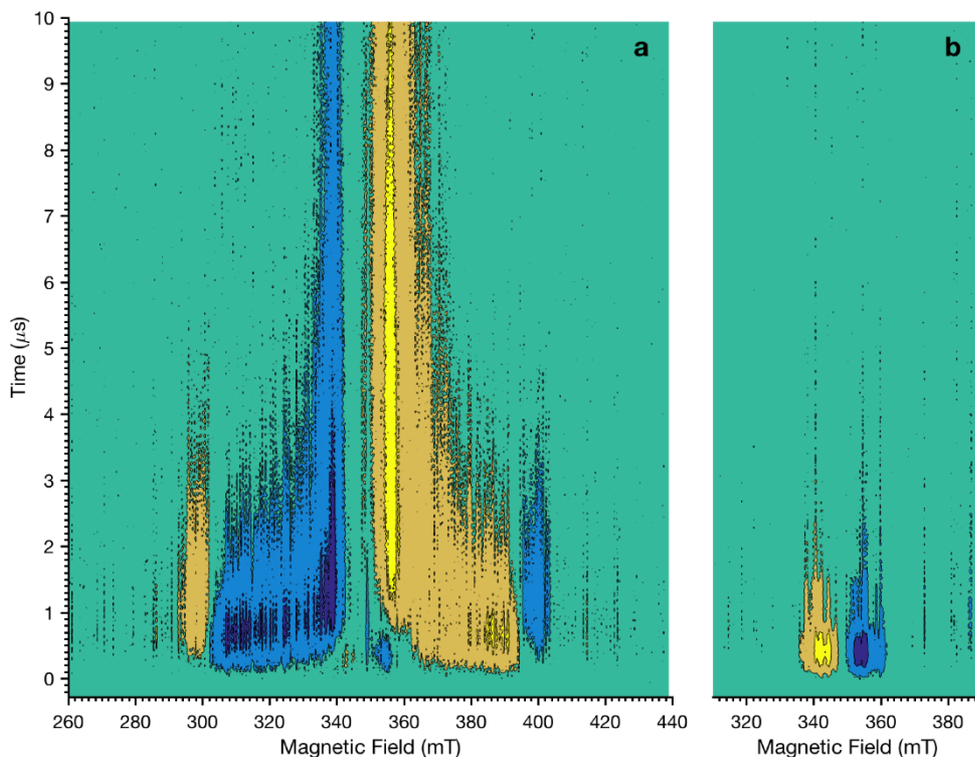
### Spin-mixing in a diketopyrrolopyrrole monomer / fullerene blend charge transfer state: a transient absorption spectroscopy and time-resolved electron paramagnetic resonance study

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**Figure S1.** The ground state absorbance spectra of the pristine TDPP film compared to solution (6 % o-dichlorobenzene, 94 % chloroform by volume), along with the two blend

films of TDPP:PC<sub>60</sub>BM and TDPP:PC<sub>70</sub>BM (both 1:2 by weight) (a). The fluorescence quenching from pristine TDPP to the blend films with fullerene is shown in (b).



**Figure S2.** Contour representations of the 2D TREPR surfaces correlating the signal intensity as a function of magnetic field and time. (a) TDPP:PC<sub>70</sub>BM 1:2 blend and (b) PC<sub>70</sub>BM film. Visual comparison suffices to attribute the central feature in the TDPP:PC<sub>70</sub>BM 1:2 blend spectrum to a PC<sub>70</sub>BM triplet generated via ISC.