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Two-step charge photogeneration dynamics in polymer/fullerene blends for photovoltaic applications

Sanjeev Singh*⁺, Bill Pandit*, Tek P. Basel, and Z. Valy Vardeny**

Department of Physics & Astronomy, University of Utah, Salt Lake City, Utah 84112

We measured the picoseconds (ps) transient dynamics of photoexcitations in blends of poly(3hexyl-thiophene) [P3HT] (donors-D) and fullerene [PCBM] (acceptor-A) using the transient pump/probe photomodulation technique in an unprecedented broad spectral range from 0.25 to 2.5 eV; and compared the results with organic solar cell performance based on the same blends. In D-A blends with maximum domain separation such as regio-regular P3HT/PCBM with (1.2:1) weight ratio having solar cell power conversion efficiency of ~4%, we found that although the photogenerated intrachain excitons in the polymer nano-domains decay within ~10 ps, no charge polarons are generated on their expense up to ~ 2 ns. Instead, there is a build-up of chargetransfer (CT) excitons at the D-A interfaces having the same kinetics as the exciton decay, which dissociate into separate polarons in the D and A domains at a much later time (>>1 ns). This 'two-step' charge photogeneration process may be typical in organic bulk heterojunction cells. Although the CT excitons are photogenerated on the exciton expense much faster in D-A blends having smaller domain size such as in regio-random P3HT/PCBM, their dissociation is less efficient because of larger binding energy. This explains the poor solar cell power conversion efficiency (<0.1%) based on this blend. Our results support the 'two-step' charge photogeneration mechanism in polymer/fullerene blends, and emphasize the important role of the CT binding energy in generating free charge polarons in organic solar cells.

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* These two authors contributed equally to this work.

- ** To whom correspondence should be addressed.
- ⁺ Present address: School of Electrical and Computer Engineering, Georgia Institute of Technology, Atlanta, GA.

The process of charge photogeneration in organic photovoltaic cells is still a matter of debate. In contrast to the labyrinth photosynthesis process that has evolved in nature [1], the charge photogeneration process in organic photovoltaic cells utilizes one type of heterojunction between two organic semiconductors. The two organic semiconductors, dubbed donor (D-) and acceptor (A-) are cast from solution mixtures to form thin films having nanosize domains of relatively pristine materials and large D-A interface area [2-4]. This type of architecture, dubbed 'bulk heterojunction' usually allows for light absorption in the bulk donor domains that generate excitons, followed by exciton dissociation at the D-A interfaces. However the process by which the excitons reach the D-A interfaces and dissociate to generate separate charge polarons in the D-A nano-domains is only now being the focus of attention [5,6].

It was originally postulated that once the exciton in the bulk donor domain reaches the D-A interface, it undergoes an ultrafast electron transfer to the acceptor forming a hole-polaron (P^+) in the donor and electron-polaron (P^-) in the acceptor, which are free to participate in the subsequent charge transport process towards the device electrodes [7]. However the mutual P^+ - P^- Coulomb attraction should prevent a complete charge separation even if the offset energy of the donor and acceptor active levels is taken into account [8]. On the contrary, the bound P^+ - P^- pair forms a charge transfer (CT) state at the D-A interface deep below the D and A optical gaps. However in spite of ample spectroscopic evidence for the existence of such CT state at the D-A interfaces [9-13], it has been argued that it lies too deep in the gap to have any effect on the charge photogeneration process in the blends [10].

In this work we used the pump/probe transient photomodulation spectroscopy in an unprecedented broad spectral range to elucidate the early stages of the charge photogeneration process in the prototype D-A blend, namely the donor polymer regio-regular (RR-) (3 hexyl thiophene) [P3HT; see Fig. 1(a) inset] and the fullerene acceptor molecule [6,6]-phenyl-C₆₁-butyric acid methyl ester [PCBM; Fig. 1(b) inset]. This blend shows separated D-A domains and consequently has high solar power conversion efficiency, $\eta \sim 4\%$ [13]. Although there have been several studies on the photoexcitation dynamics of the P3HT/PCBM blend, many questions still remain unresolved regarding the role of the CT state on the charge generation processes [14-25]. We present compelling evidence that after the photoexcited excitons in the polymer domains

reach the D-A interfaces the charge generation process proceeds via the formation of CT excitons at the interfaces. In RR-P3HT/PCBM with (1.2:1) weight ratio the photogenerated excitons in the polymer domain reach the D-A interfaces forming CT excitons within ~10 ps. In contrast, in regio-random (RRa)-P3HT/PCBM blend where the D and A domain sizes are much smaller, the CT excitons at the D-A interfaces are generated within ~200 fs. However the subsequent exciton dissociation process in this blend is hampered by the large CT binding energy [11], which explains the smaller η -value (<0.1%) of solar cells based on this blend. Our findings support a 'two-step' process for the charge photogeneration in organic D-A blends [5], and emphasize the important role of the CT exciton binding energy in generating free charges in organic solar cells.

The P3HT polymers and PCBM fullerene were supplied by *Plextronics Inc.* [13]. The mixing ratio of the P3HT/PCBM blends was 1.2:1 by weight, which gives the optimal η -value in solar cells [6]. The films were spin cast onto CaF₂ or CsI that are transparent in the mid-IR spectral range. The RR-P3HT/PCBM blend film was thermally annealed at 150° C for 30 minutes to enhance the D-A domains size [6]; whereas the RRa-P3HT/PCBM film was used as deposited.

For the polarized transient photomodulation spectroscopy we used the femtoseconds (fs) twocolor pump/probe correlation technique with two different pulsed laser systems based on a Ti: Sapphire oscillator [26]. These are: a low-power (energy/pulse ~0.1 nJ) high repetition rate (~80 MHz) pulsed laser system for the mid-IR spectral range; and a high power (energy/pulse ~10 μ J) low repetition rate (~1 kHz) pulsed laser system for the near-IR/visible spectral range. The pump excitation photon energy, $\hbar\omega(pump)$ was set at 3.1 eV for above-gap and 1.55 eV for below-gap excitation, respectively. The pulse energy flux on the film was adjusted so that the initial photoexcited exciton density, N(0)≈10¹⁶/cm³ [N(0)≈10¹⁷/cm³] for the mid-IR [near-IR] laser system. For the probe in the mid-IR measurements we used an optical parametric oscillator (OPAL, Spectra Physics) that generates $\hbar\omega(probe)$ ranging from 0.55 to 1.05 eV. In addition, we also used a 'difference-frequency crystal' (AgGaS₂) and the 'signal' and 'idler' beams of the OPAL for generating $\hbar\omega(probe)$ in the spectral interval 0.25 to 0.43 eV [11]. The pump beam was modulated at frequency, f=40 kHz and the changes, ΔT in the probe transmission, T were measured using a phase-sensitive technique. For the transient near-IR/visible spectroscopy measurements we used white light super-continuum as probe, spanning the spectral range from 1.15 to 2.5 eV; the pump modulation frequency here was synchronized with the laser rep. rate. The transient photomodulation signal, $\Delta T/T$ is positive for photoinduced absorption (PA) and negative for photo-bleaching and stimulated emission. $\Delta T(t)/T$ was measured using a computer controlled translation stage up to 2 ns, with time resolution of ~150 fs set by the pump/probe cross-correlation. $\Delta T(0)/T$ spectra from the two laser systems were normalized to each other at several wavelengths in the near-IR/visible spectral range [26].

Since some photoexcitations may live longer than the time interval between successive pulses, then a background PA may be formed. An advantage of the mid-IR laser system is that the background PA spectrum can be readily measured using the same pump/probe set up as for the ultrafast response. This was achieved by measuring the PA signal at t<0 since the probe pulse in this situation arrives before the pump pulse, and therefore is affected by the 'left-over' photoexcitations in the film that survive in between successive pulses [27]. Recall that the background PA is modulated at frequency of 40 kHz, and thus is sensitive to long-lived photoexcitations in the film having lifetime longer than $\sim 1/f$ (=25 µsec).

The bulk heterojunction OPV solar cell devices were composed of a transparent indium tin oxide (ITO) anode; a spin-cast polyethylenedioxythiophene/polystyrene sulphonate (PEDOT/PSS) hole transport layer; an active material layer spin-cast from a blend of P3HT donor and PCBM acceptor; and capped with LiF/Al cathode. The ITO-coated glass substrates (Delta Technology, CB-50IN) were cleaned by ultrasonic treatment and oxygen plasma treatment. The PEDOT/PSS (Clevios, P VP AI 4083) layer was spin-cast at 5000 RPM for 20 sec at ambient conditions, and transferred to a nitrogen-filled glovebox ($O_2 < 1$ ppm) for annealing at 120 °C for 30 min. The organic blend comprised of P3HT (Plextronics, Plexcore OS 2100) and PCBM (purity>99.9%) that were prepared at weight ratio of 1.2:1 in 1,2-dichlorobenzene solution that was heated at 50 °C for 30 min. and stirred overnight. The device active layer was spin-cast from the solution blend at 400 RPM for 6 min. and annealed at 150 °C for 30 min.; the device active area was 2 mm×2.5 mm. The fabrication was completed by thermally evaporating a 1 nm thick film of LiF

layer followed by a 100 nm thick film of Al. Finally the device was encapsulated under a cover glass using UV-curable optical adhesive (Norland, NOA 61). The device I-V characteristics under illumination were measured using a Keithley 236 Source-Measure unit. The light intensity of the solar simulator composed of a xenon lamp and an AM1.5G filter was calibrated at 100 mW/cm² using a pre-calibrated silicon photovoltaic cell. The OPV device output current was measured using phase sensitive lock-in technique.

In order to identify photogenerated polarons in the photomodulation spectrum we also measured the doping induced absorption spectrum in P3HT, where a thin polymer film was exposed to low pressure iodine gas for ~10 seconds. The doping induced absorption spectrum is then obtained by subtracting the optical density of the pristine polymer film from that of the doped film, as shown in Fig. 3(a) inset. It is seen that the doping induced absorption spectrum in RR-P3HT (and RRa-P3HT) is dominated by two absorption bands, namely P_1 and P_2 that peak at 0.1 and 1.8 eV, respectively [28], which are due to delocalized hole-polarons on the polymer chains.

We first measured $\Delta T(t)/T$ spectra of *pristine* polymer and fullerene films (Fig. 1). $\Delta T(0)/T$ spectrum of pristine RR-P3HT film (Fig 1(a)) is dominated by a single PA₁ band at 1 eV followed by photo-bleaching above 1.97 eV and a small stimulated emission band at 1.75 eV, which attests to the excellent quality of the RR-P3HT polymer used here [13]. These three spectral features originate from photogenerated excitons since they decay together (Fig. 1(a) right inset) with an exponential time constant, τ_0 =70 ps. No photogenerated polarons which peak at 0.1 and 1.8 eV (see Fig. 3(a) inset) are observed here. Figure 1(b) shows $\Delta T(0)/T$ spectrum of pristine PCBM film. It is dominated by two PA bands, namely EX₁ and EX₂ at 1.0 and 2.25 eV, respectively that are due to photogenerated excitons. A third PA band, CT at 1.75 eV originates from charge-transfer excitons in the fullerene film, since it does not exist in the photomodulation spectrum of isolated PCBM molecules in polystyrene (Fig. 1(b) right inset). No photogenerated polarons which peak at 1.15 eV [29,30] are discerned.

To better understand the transient PA spectra in the polymer/fullerene blends we measured the x-ray diffraction (XRD) pattern from the RRa- and RR-P3HT/PCBM blend films (Fig. 2) using the Cu K_{α} X-ray line at λ =0.154 nm. The XRD pattern of RR-P3HT/PCBM contains a prominent

Bragg band at $2\theta_{[100]}=5.3^{\circ}$ and its harmonics at $2\theta=10.7^{\circ}$ and 16° , respectively that originate from the P3HT nanocrystalline domains in the film [31]; as well as a smaller Bragg band at $2\theta_{[311]}=19.3^{\circ}$ that originates from the PCBM nanocrystalline domains in the blend [32]. We therefore conclude that the annealed RR-P3HT/PCBM blend film contains separate donor and acceptor crystalline domains. We may estimate the average nanocrystalline domain size, *D* from the full width at half maximum, $\Delta_{2\theta}$ of the respective Bragg bands using the Scherrer relation: *D* = $0.9\lambda/\Delta_{2\theta}\cdot\cos\theta$; we obtain $D\approx16$ nm (20 nm) for the polymer (PCBM) nano-domains. In contrast, the XRD pattern of the RRa-P3HT/PCBM blend does not show prominent P3HT band harmonics, and in addition the PCBM band is missing (Fig. 2). These indicate that the PCBM molecules do not form well-separated domains here; instead, they penetrate into the P3HT lamellae and consequently are much closer on average to the polymer chains.

Figure 3(a) shows $\Delta T(t)/T$ spectra of RR-P3HT/PCBM blend. $\Delta T(0)/T$ spectrum is very similar to that of pristine RR-P3HT (Fig. 1(a)), indicating that excitons are initially photogenerated within the polymer domains. At t>0 the excitons decay; however *no polarons are generated* at the expense of the exciton decay up to 300 ps since there is no PA build-up at low $\hbar\omega$ (probe), where

the polaron P_1 band dominates the absorption spectrum (as seen in Fig. 3(a) inset). We thus conclude that the photogenerated excitons in the polymer domains decay into a *new state* that is not separated free polarons. This new state must be related with the D-A interfaces in the films since the excitons do not form such a state in the pristine polymer, and we thus propose that is a CT exciton at the D-A interface. In contrast, the background PA spectrum in the mid-IR (Fig. 3(a)) is very similar to the P_1 band in the polaron doping induced absorption spectrum (Fig. 3(a) inset), showing that charge polarons are indeed photogenerated in this RR-P3HT/PCBM film but at later time; in agreement with the high solar cells efficiency based on this blend [13]. We thus conclude that the charge photogeneration process in the blend proceeds in two stages [5,16]. The first stage is exciton trapping in CT states at the D-A interfaces, followed by a much slower exciton dissociation into free polarons in the D and A domains at t>>2 ns (= the time limit of our translation stage).

The exciton decay dynamics in the blend is much faster than in the pristine polymer (see PA₁ decays in Figs. 3(a) and 1(a)). The shorter lifetime in the blend is related to the exciton dynamics towards the D-A interfaces, and therefore we studied the PA₁ decay kinetics in more detail. PA₁ decay cannot be fit with a single or few exponential decay functions; nor can it be fit using a diffusion model [\sim (1+t/ τ)⁻¹]. Alternatively, PA₁ decay can be fit using multiple power-law decays that originate from a Förester resonant energy transfer (FRET) into the CT exciton [33], averaged over the exciton initial distance from the D-A interface (see Appendix A). This model yields the following time dependent surviving exciton density N(t) in the polymer nano-grains;

$$N(t)/N(0) = \exp(-t/\tau_0)[m_1 + m_2(C_1t^{1/2} - C_2t^{1/3} + C_3t^{1/6})],$$
(1)

where $\tau_0=70$ ps is the natural exciton lifetime in RR-P3HT that is obtained from the PA₁ dynamics of Fig. 1; m₁ and m₂ are fitting parameters; and the C constants are given by the relations: C₁=0.2u⁻³, C₂=0.66u⁻² and C₃=0.54/u, where u=*D*/2*R*₀ is the parameter ratio of the grain size, *D* to twice the FRET radius, *R*₀, which was measured before to be between 3 and 9 nm [34]. Using *R*₀=6 nm and *D*=16 nm from the XRD studies, we obtain u=1.3. Subsequently the excellent fit to the PA₁ decay seen in Fig. 3(b) was obtained using m₁=0.14 and m₂=7.

In support of the CT intermediate role in the charge photogeneration process in the blend, Fig. 3(a) also shows that PA build-up indeed occurs in both mid-IR and near-IR [35] spectral ranges. In fact there are two PA bands, namely CT₁ in the mid-IR and CT₂ in the near-IR that are generated at longer time *at the expense of the exciton PA*₁ *decay*. Fig. 3(b) shows that the CT₂ build-up dynamics in the near-IR closely matches the exciton PA₁ decay, since the same function of time fits both PA₁ decay and CT₂ build-up dynamics (measured at 1.75 eV probe). Fig. 4(a) more clearly shows the two PA bands that are generated at the expense of the exciton PA₁ decay. To obtain the full photogenerated CT spectrum we subtracted the photomodulation spectrum at t=30 ps from that at t=0, after normalizing the two PA bands at 1 eV and 2 eV for the CT₁ and CT₂ bands, respectively. It is seen that the CT spectrum contains two prominent PA bands that peak at 0.6 (CT₁) and 1.75 eV (CT₂), respectively, which are very different than the bands P₁ and P₂ of polarons (Fig. 3(a) inset). Consequently we propose that these two PA bands are due to optical transitions within the CT manifold at the D-A interfaces [5,8].

To support this assumption we measured the transient photomodulation spectrum using $\hbar\omega(\text{pump})=1.55 \text{ eV}$, which is *below the optical gap* of the polymer and fullerene constituents. Such low $\hbar\omega(\text{pump})$ can resonantly excite the CT state at the D-A interface since its energy was measured to be between 1.2 to 1.6 eV [13], without first photogeneration of excitons in the polymer domains. Fig. 4(b) shows that under these conditions the two CT PA bands are *instantaneously* photogenerated; which is compelling evidence that they originate from the CT states at the interfaces. This supports our assignment for the CT bands in the transient photomodulation spectrum of this blend.

Interestingly the background PA spectrum excited with below-gap pump excitation (Fig. 4(b)) is very similar to that generated using above-gap pump excitation (Fig. 3(a)), which we identified as due to long-lived charge polarons [13]. This shows that there exists a mechanism where thermalized CT excitons at the D-A interfaces are able to separate into free polarons in the donor and acceptor domains, *regardless of the initial* $\hbar\omega$ (pump) [5]. This finding is very important,

since it can refute the notion that the CT state in the blend lies too deep in the gap to have any influence over the charge photogeneration process. Apparently the exciton kinetic energy when reaching the CT state plays a minor role in the charge photogeneration process; this may explain the flat spectral response of the photocurrent action spectrum in organic solar cells [13].

For comparison, we also study the charge photogeneration mechanism in RRa-P3HT/PCBM blend where the fullerene molecules are closer to the polymer chains on average. Fig. 5(a) shows $\Delta T(t)/T$ spectra of *pristine* RRa-P3HT. It also contains a single PA₁ exciton band at ~1 eV followed by stimulated emission above ~1.75 eV that shares the same dynamics (Fig. 5(a) inset). However $\Delta T(0)/T$ spectrum in RRa-P3HT/PCBM blend (Fig. 6(a)) is very different from that in RR-P3HT/PCBM blend (Fig. 3(a)). The former spectrum shows the same two CT PA bands (namely CT₁ and CT₂ discussed above) that are generated within ~ 500 fs (Fig. 6(b)), in concert with the ultrafast decay of the exciton PA₁ band. The fast exciton decay here is consistent with the proximity of the PCBM molecules to the P3HT polymer chains in RRa-P3HT/PCBM blend. Interestingly, the background PA spectrum here does not show long-lived polaron

photogeneration; in fact the background PA spectrum is the *same* as the transient PA spectrum. This shows that the photogenerated CT excitons in this blend cannot easily dissociate into free polarons, in agreement with the poor solar cell efficiency (<0.1%) based on this blend (Fig. 7). It also shows that the CT exciton dissociation is related to the D-A domain size. The larger is the D-A interface area the smaller is the CT exciton binding energy, and consequently the more efficient is the CT exciton dissociation into separate charge polarons in the D and A domains [36].

This conclusion is further supported by the I-V characteristics under solar-like illumination of photovoltaic devices based on RR-P3HT/PCBM and RRa-P3HT/PCBM blends using AM1.5 filter as shown in Fig. 7. It is seen that the power conversion efficiency and short circuit current of RR-P3HT/PCBM blend are more than an order of magnitude higher than that of RRa-P3HT/PCBM blend.

In summary, using the ps transient pump/probe photomodulation technique in P3HT/PCBM blends with typical D-A bulk heterojunction morphology we demonstrated that the charge photogeneration mechanism in organic solar cells occurs in two-steps. First the photogenerated excitons in the polymer domains reach the D-A interfaces within few ps time depending on the domain size, where they form CT excitons. This process is followed by CT exciton dissociation into free charged polarons in the D and A domains in the ns-µsec time scale, which remains to be observed. The CT exciton dissociation depends on the CT binding energy, which is significantly lower for larger D-A interface area. Our results emphasize the importance of the D-A domain size in organic solar cells based on bulk heterojunction morphology.

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Appendix A

Derivation of Förster energy transfer dynamics for excitons in the polymer grains

We assume that the polymer grains are spherical having radius R (Fig. A1(a)). Also we hypothesize that the Förester energy transfer (FRET) kinetics from a point "r" inside the polymer grain to its surface is exponential in nature [33]. Therefore the surviving exciton density inside the polymer grain at point r can be written as follow:

$$n(t) = n(0)\exp\left(-v(r)t\right) \tag{A1}$$

where $v(r) = v_o \frac{R_o^6}{(R-r)^6}$ and R_o is the FRET radius. In order to calculate the total surviving exciton density, N(t) inside the polymer grain we integrate over the distance, *r*, normalized by the sphere volume:

$$N(t) = N(0) \frac{\int_0^{R_{min}} exp(-v(r)t) 4\pi r^2 dr}{4\pi R^3/3}$$
(A2)

Here R_{min} is the distance from the center where v(r) reaches its maximum rate, v_{max} , and thus does not change with r for $r > R_{min}$. Equation (A2) can also be written in terms of the exciton lifetime, τ

$$N(t) = N(0) \frac{\int_{\tau_{max}}^{\tau_{min}} exp\left(-\frac{t}{\tau}\right)g(\tau)d\tau}{4\pi R^{3}/3}$$
(A3)

Where $\frac{1}{\tau} = \frac{1}{\tau_o} \frac{R_o^6}{(R-r)^6}$ and τ_o is the exciton natural decay lifetime in the bulk polymer. Writing for simplicity $R = \mu Ro$, where μ is a constant ($\mu < 1$), we can substitute the distance r and lifetime τ using μ : $r = R_o (\mu - (\frac{\tau}{\tau_o})^{\frac{1}{6}})$ and $dr = -\frac{R_o}{6\tau_o^{1/6}}\tau^{-5/6}d\tau$. Substituting these relations in Eq. (A2) we obtain the following equation for N(t):

$$\frac{N(t)}{N(0)} = \frac{\exp\left(-\frac{t}{\tau_0}\right)}{2\mu^3 \tau_0^{1/6}} \int_{\tau_{min}}^{\tau_{max}} \frac{\exp\left(-\frac{t}{\tau}\right)}{\tau^{5/6}} \left[\mu^2 + \left(\frac{\tau}{\tau_0}\right)^{\frac{1}{3}} - 2\mu \left(\frac{\tau}{\tau_0}\right)^{\frac{1}{6}}\right] d\tau \tag{A4}$$

Replacing $X = \frac{t}{\tau}$ and assuming $X_{min} = 0.1$ and $X_{max} = \infty$ we then obtain:

$$N(t)/N(0) = \exp\left(-\frac{t}{\tau_0}\right) \left[m1 + m2\left[\left(\frac{0.542}{\mu}\right)t^{\frac{1}{6}} - \left(\frac{0.66}{\mu^2}\right)t^{\frac{1}{3}} + \left(\frac{0.203}{\mu^3}\right)t^{\frac{1}{2}}\right]\right]$$
(A5)

The above expression is derived for a single grain size R. For simplicity we assume that most polymer grains are of size R_{avg} where $R_{avg} = \mu_{avg}R_o$. Including μ_{avg} into Eq. (A5) we finally get Eq. (A6) [or Eq. (1) in the text] for the decay of excitons in the polymer blend.

$$N(t)/N(0) = \exp\left(-\frac{t}{\tau_o}\right) \left[m1 + m2\left[\left(\frac{0.542}{\mu_{avg}}\right)t^{\frac{1}{6}} - \left(\frac{0.66}{\mu_{avg}^2}\right)t^{\frac{1}{3}} + \left(\frac{0.203}{\mu_{avg}^3}\right)t^{\frac{1}{2}}\right]\right]$$
(A6)

We found that the best fit to PA₁ decay was obtained using m2 = 7 and $\mu_{avg} = 1.3$, as seen in Fig. A1(b). Using this μ_{avg} and $R_0 = 6$ nm, the average grain diameter, D of the polymer in RR-P3HT/PCBM [1.2:1] blend ~16 nm, which is in good agreement with our XRD measurements described in Fig. 2.

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Figure Captions

FIG. 1. (color on line) (a) The transient photomodulation spectrum of pristine RR-P3HT film at t=0 and t=100 ps, respectively; the exciton bands PA₁, SE and PB are indicated. The right inset shows the transient decay of PA₁ and PB bands up to t=200 ps; the left inset shows the polymer backbone chain. (b) The transient photomodulation spectrum of a PCBM film at t=0; the exciton PA bands EX₁ and EX₂, and CT exciton band are indicated. The right inset shows $\Delta T/T(0)$ spectrum of isolated PCBM molecules in polystyrene (weight ratio of 1:100) that lacks the CT exciton band. The left inset shows the PCBM molecular structure.

FIG.2. (color on line) The XRD pattern from RR-P3HT/PCBM (red) and RRa-P3HT/PCBM (blue), where the P3HT bands [100], [200] and [300] and PCBM band [311] are assigned; the inset focuses on the PCBM band.

FIG. 3. (color on line) (a) The transient photomodulation spectrum of RR-P3HT/PCBM blend film at t=0 and t=300 ps, respectively; the exciton band PA₁, and CT exciton bands CT_1 and CT_2 are indicated. The green circles and line represent the background (BG) PA spectrum measured at t=-5 ps. The inset shows the doping induced absorption of pristine RR-P3HT film, where the polaron bands P₁ and P₂ are assigned. (b) The transient decay of PA₁, build-up dynamics of CT₂, and the PB decay up to 180 ps. The line through the data points is a fit using the FRET mechanism (see text); the same function also fits the CT₂ build-up dynamics.

FIG. 4. (color on line) (a) The transient photomodulation spectrum of RR-P3HT/PCBM blend film at t=30 ps excited at 3.1 eV, normalized and subtracted from the spectrum at t=0, that shows the two newly formed CT_1 and CT_2 bands. (b) Same as in (a) but at t=0 and excited at 1.55 eV, which is below the gap of both polymer and fullerene constituents.

FIG. 5. (color on line) The transient photomodulation spectrum of pristine RRa-P3HT film at t=0 and t=200 ps, respectively; the exciton bands PA_1 and SE are indicated. The right inset shows the transient decay of PA_1 and SE bands up to t=200 ps; the left inset shows the polymer backbone chain.

FIG. 6. (color on line) (a) The transient photomodulation spectrum of RRa-P3HT/PCBM blend at t=0 and t=10 ps, respectively excited at 3.1 eV; the PA bands PA₁, CT_1 and CT_2 are assigned. (b) The transient decay of PA₁, and build-up dynamics of CT_1 and CT_2 up to 15 ps.

FIG. 7. (color on line) The *I-V* characteristic of two solar cells based on PCBM blend with RR-P3HT (black) and RRa-P3HT (red) donor polymers under solar-like illumination of AM 1.5. The inserted Table gives the device photovoltaic characteristic parameters such as short circuit current density, J_{sc} ; open-circuit voltage, V_{oc} ; fill-factor, FF; and the power conversion efficiency, PCE in %.

FIG.A1: (color on line) (a) Schematic diagram of the polymer grain of radius R, where r is the exciton distance from the center. (b) The decay of PA₁ band in the RR-P3HT/PCBM blend, measured at 1 eV probe photon energy. The decay is fit using Eq.(A6) with the parameters given in Appendix A1.



FIG.1



FIG. 2







FIG.4



FIG.5



FIG. 6





FIG. A1