Ultrafast Energy Transfer in Ultrathin Organic Donor/Acceptor Blend

Ajay Ram Srimath Kandada1,2, Giulia Grancini2, Annamaria Petrozza2, Stefano Perissinotto2, Daniele Fazzi2, Sai Santosh Kumar Raavi2 & Guglielmo Lanzani1,2

1 Dipartimento di Fisica, Piazza Leonardo da Vinci 32, 20133 Milano, Italy, 2 Center for Nano Science and Technology @ PoliMi, Istituto Italiano di Tecnologia, via Pascoli 70/3 20133 Milano, Italy.

It is common knowledge that poly(3-hexylthiophene) (P3HT)/[6,6]-phenyl-C61-butyric acid methyl ester (PCBM) blend, a prototype system for bulk heterojunction (BHJ) solar cells, consists of a network of tens of nanometers-large donor-rich and acceptor-rich phases separated by extended finely intermixed border regions where PCBM diffuse into P3HT. Here we specifically address the photo-induced dynamics in a 10 nm thin P3HT/PCBM blend that consists of the intermixed region only. Using the multi-pass transient absorption technique (TrAMP) that enables us to perform ultra high sensitive measurements, we find that the primary process upon photoexcitation is ultrafast energy transfer from P3HT to PCBM. The expected charge separation due to hole transfer from PCBM to P3HT occurs in the 100 ps timescale. The derived picture is much different from the accepted view of ultra-fast electron transfer at the polymer/PCBM interface and provides new directions for the development of efficient devices.

Due to the tremendous surge of interest in organic photovoltaics, the photophysics at the electron donor/acceptor (D/A) organic interface is a highly debated topic in material science. One of the most extensively investigated D/A blend is made of poly(3-hexylthiophene) (P3HT), as the donor component, mixed with phenyl-C61-butyric acid methyl ester (PCBM), as the acceptor. This blend provides solar cells with 85% Internal Quantum Efficiency1 and power conversion efficiency of around 5%2. Such high quantum yield of charge photogeneration with respect to pure organic semiconductors results from blending the two components into a BHJ. The D/A components are deposited from common solution and tend to segregate over tens of nanometers forming PCBM-rich clusters and crystalline domains of P3HT1–6. The two phases however are not separated by sharp interfaces, because substantial intermixing takes place in the border region defined by the extent of diffusion of PCBM molecules into P3HT7. In this intermixed region, the two components are finely dispersed at the molecular level and have been clearly observed in bi-layer structures7. Here we address the photophysical processes happening only at the intermixed border region by probing an ultrathin film with thickness much below the PCBM diffusion length.

By and large, charge transfer is regarded as the fundamental event occurring upon donor photo-excitation in the BHJ8,9 while energy transfer is generally ignored, at least at early times. However the proximity of D and A molecules in the blend may in principle also lead to electronic excitation energy transfer. Indeed this has been reported for a few systems prepared ad hoc, such as D/A bilayers separated by an interlayer that hampers charge separation10–12 or in chemically bound dyads, wherein the D and A are in a fixed positions with respect to each other13. Both these cases are thus modifications of the standard BHJ, where essentially the wavefunction overlap is largely reduced. On the contrary, there are no experimental evidences of electronic excitation energy transfer as primary event in standard BHJ systems. Based on photoluminescence (PL) quenching experiments in D/A blends, it was proposed that energy transfer dominates charge transfer only after 5 ps from the excitation14. The suggested picture is that close pairs undergo charge transfer, while distant pairs may first interact by energy transfer. Förster radius, the measure of the electronic coupling strength between D and A, has been evaluated for the prototypical systems such as P3HT/PCBM in the range between 1 and 5 nm11,13,14. Why electronic excitation energy transfer appears only in special situations and why charge transfer overwhelms it in BHJ, remains an open question.

In this work we study the photoexcitation dynamics of the intermixed P3HT/PCBM region obtained from a nominal 5/5 nm diffuse bi-layer. Contrary to the expectations, we demonstrate that ultrafast electronic energy transfer between donor and acceptor is the primary process occurring upon photoexcitation. Subsequent hole transfer from PCBM to the polymer takes place on the 100 ps time scale, resulting in the expected charge separation.

Received 17 April 2013
Accepted 7 June 2013
Published 25 June 2013

Correspondence and requests for materials should be addressed to G.L. (guglielmo.lanzani@iit.it)
A thin P3HT layer of about 5 nm has an optical density of about $2 \times 10^{-2}$ (see Supplementary Information (S.2) for thin film preparation and characterization). This value is far too low to have detectable signal above the shot noise limit in a standard pump-probe system. To overcome such a limitation, we propose a novel Transient Absorption by Multiple Pass (TrAMP) technique, that enables us to perform measurements on samples with optical densities as low as $10^{-5}$. This technique draws its inspiration from absorption spectroscopy of gases where high sensitivities are achieved by increasing the optical path length by multiple reflections of the beam in the gas chamber. The same principle is extended to transient absorption spectroscopy by passing the pump and probe pulses multiple times, keeping the time delay between them constant. In this way, the pump-probe signals can be amplified by a factor proportional to the number of passes through the sample (see Supplementary Information (S.1) for more details).

Figure 1a shows the schematic lay-out of the TrAMP setup while Figure 1b demonstrates the concept and validity of our approach. It shows the TrAMP kinetics collected at various numbers of passes in a 15 nm thin film of P3HT on a 250 μm fused silica substrate. The pump is at 400 nm and the probe wavelength tuned at 680 nm.

Standard pump probe (single pass) cannot detect any differential transmission changes in such a thin film. However, by the multiple pass scheme, the TrAMP technique provides an efficient amplification of the pump-probe signal, that is shown growing at each pass, while keeping the kinetic itself intact. This also demonstrates that possible effects on the time resolution, such as group velocity delay between pump and probe and pulse broadening, are negligible in our experiment, even in the worst condition of large spectral separation between the pump and the probe wavelength. By appropriate group velocity mismatch calculations (see Supplementary Information for details), it can be shown that the expected time resolution of the experiment is within 200 to 400 fs after 15 passes, depending on the pump-probe wavelengths. After certain number of passes, signal amplification reaches a saturation level that can be attributed to the almost complete absorption of the pump. The threshold number of passes depends on the optical density of the sample and for our bilayers this number is between 10 and 15 and all the experiments were conducted within this range. The pump wavelength for all the experiments reported here was tuned to 520 nm. An input energy of 10 nJ per pulse was employed. The spot-size varies for each pass in TrAMP between 100 μm and 200 μm. This will correspond to a maximum fluence of about 30 μJ/cm².

In order to investigate the intermixed D/A system avoiding large phase segregation, we first deposit a 5 nm P3HT layer on fused-silica substrate and then we spin coat the PCBM on top from an orthogonal solvent with a nominal thickness of 5 nm. The absorption spectrum of the sample is a linear combination of the absorption spectrum of the two components with an OD of 0.02 at the absorption peak (Figure 2a). Because the P3HT film thickness is smaller than the PCBM molecule diffusion length, PCBM almost fully penetrate the polymer and we obtain a layer of P3HT doped with PCBM (see cartoon in Figure 2b). This is also suggested by simple AFM analysis performed on the bilayer (see AFM characterization in figure S3 of the Supplementary Information).

### Results

Figure 3a shows the transient absorption ($\Delta T/T$) spectra of 5 nm thin P3HT film as a control experiment. The TrAMP spectrum and its temporal evolution are consistent with the one reported in literature. At wavelengths shorter than 620 nm it exhibits a positive signal due to ground state bleaching (GSB). At probe wavelengths greater than 620 nm it shows a negative photo-induced absorption (PA) signal. A comparison with dynamics, as reported in Figure 3c, and a rather extended set of experimental and computational data reported in literature helps in disentangling the contributions beyond spectral overlap. Two species can be identified at early time: singlet excitons and polaron pairs (PP) as assigned by Koryyanke et al. The fast decaying PA band appearing at 650 nm is assigned to geminate recombination of PP. The longer lived signal is due to singlet states, with some contribution from polarons escaping the geminate recombination as reported in earlier TA studies on P3HT.

The $\Delta T/T$ spectrum of the P3HT/PCBM intermixed layer is shown in Figure 3b, and selected time traces are reported in Figure 3d. At a first glance, the early spectrum resembles the pure P3HT spectrum, though with a remarkable reduction in the amplitude of GSB with respect to the PA. Within 2 ps, both the P3HT GSB and the PA band centered at 650 nm decay to zero with similar kinetics, giving rise to a negative signal extending on the whole experimental range. Notably the 580–620 nm spectral region that corresponds to P3HT ground state absorption shows null or negative signal. This indicates that P3HT is weakly or not excited anymore. The broad PA should correspond to an excited state population that is not perceived by the ground state of the polymer. The charge transfer process cannot explain such an observation: upon charge transfer (CT) and hole-polaron formation, the polymer GSB...
stays constant due to the persistent depletion of the ground state population.

Our observation can however be explained by electronic energy transfer from P3HT to PCBM. Accordingly, P3HT GSB would decay and a new PA due to \( S_n \leftarrow S_1 \) in PCBM would appear (see Supplementary Information (S.4) for quantum chemically computed \( S_n \leftarrow S_1 \) transitions in PCBM). As a reference, \( \Delta T/T \) spectrum of pristine PCBM, measured upon resonant excitation (400 nm), is

Figure 2 | Bilayer characterization. (a) Illustration of the morphology of the 5 nm bilayer of P3HT/PCBM used in present experiment. (b) Absorption spectrum of 5 nm films of P3HT and PCBM and the bilayer.

Figure 3 | Ultrafast dynamics in the ultrathin P3HT/PCBM bilayers. (a) TrAMP spectra and (c) the associated kinetics of a control film of P3HT with a thickness of 5 nm. (b) TrAMP spectra of P3HT/PCBM (5 nm/5 nm) bilayer at pump-probe delays of 1 ps, 5 ps and 500 ps. The TA spectrum of PCBM pumped at 400 nm is shown in the 1 ps window. The absorption of P3HT is also shown. (d) The associated TrAMP kinetics of the bilayer taken at \( \lambda_{\text{probe}} = 580 \) nm (GSB) and at 650 nm and 750 nm (PA). Also plotted is the kinetic at \( \lambda_{\text{probe}} = 750 \) nm of a 5 nm PCBM film. The Pump pulse for all the measurements is tuned to 520 nm.
reported in Figure 3b. It is a broad photoinduced absorption band, extending over the whole experimental range. This fits reasonably well with our observation, supporting our explanation. It is fair to note a discrepancy between the lineshape of the pure PCBM PA band and our data, in the 580–620 nm spectral region. This can be ascribed to a residual GSB of the polymer, indicating that following polymer photoexcitation there is a branching between electronic energy transfer and charge transfer towards PCBM, here in favor of the former.

Concomitant excitation of PCBM, when we excite the P3HT, should also be evaluated as a possible explanation for the reported experimental results. At this purpose we studied a 5 nm thick PCBM film, with a similar optical density of the intermixed sample, pumping at 520 nm. The plot in Figure 3d contains the time trace at 750 nm, assigned to PCBM S\textsubscript{1} ← S\textsubscript{0} PA peak (see Supplementary Information Figure S4 and S5). The signal amplitude, without any normalization, clearly shows that the role of direct PCBM excitation is negligible. We infer that direct photo-excitation of PCBM contributes less than 10% of the signal measured in the P3HT/PCBM sample.

The electronic energy transfer occurs on a time scale faster than our resolution (∼200 fs). The broad PA band assigned to PCBM is formed during pump excitation and it is present from the very beginning in our transient spectra. This explains the different ratio between GSB and PA in the P3HT/PCBM sample with respect to the one in pristine P3HT.

On the longer timescale, we observe a build-up of the GSB with a time constant of 130 ps along with a correlated increase in the PA band centered at 750 nm. The return of GSB in transient spectra is a time constant of 130 ps along with a correlated increase in the PA the one in pristine P3HT.

Discussion

The phenomenon of ultrafast energy transfer before charge transfer in systems commonly utilized in organic BHJ solar cells is puzzling and contradicting common beliefs. In order to further test our interpretation we performed experiments on tri-layered samples where charge transfer can be effectively blocked by inserting a large band-gap thick interlayer of Poly-Vinyl Acetate (PVA) between P3HT and PCBM. Measurements on the tri-layer films are reported with thin (<5 nm) and thick (>10 nm) PVA layers (Figures 5b and 5d respectively). When the PVA layer is thin the GSB of P3HT fully decays to zero in 2 ps, again leaving a PA without GSB for longer time delays. The lack of GSB is consistent with ultrafast electronic energy transfer even in presence of the thin PVA interlayer. We can set an upper limit for the time scale of the process that is again within the excitation process, about 200–300 fs. The steady PA is assigned to S\textsubscript{0} ← S\textsubscript{1} PCBM electronic transition. In presence of the interlayer, however, there is neither delayed build up of GSB nor PA at longer time. We suppose that because of the very large surface roughness, there is no substantial reduction in P3HT-PCBM average distance when a thin interlayer is deposited, preserving the dipole-dipole electronic energy transfer, while the charge transfer is suppressed by the lack of wavefunction overlap. When the interlayer is thick (Figure 5d), pure P3HT dynamics, similar to Figure 3c can be observed. In particular there is a longer lived GSB due to the singlet

![Figure 4](https://www.nature.com/scientificreports/)

**Figure 4** | Thickness dependent TrAMP dynamics. (a) TrAMP kinetic at λ\textsubscript{probe} = 900 nm for 5 nm/5 nm bilayer of P3HT/PCBM. A multieponential fit (A(t)) is also shown. (b) Similar kinetic and the fit (B(t)) for a 5 nm film of P3HT. TrAMP kinetics for the bilayers with 10 nm (c), 20 nm (d) and 50 nm (e) thicknesses of P3HT with a constant 5 nm layer of PCBM on the top. Also shown are the fits obtained by a linear superposition of A(t) and B(t) in each of the three cases.
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Considering the P3HT/PCBM interface as molecular in nature and, as Density Functional Theory (DFT) calculations provide optimized intermolecular distance between the P3HT and PCBM centers around 6–9 Å, we focus the transfer rate analysis on the 6 Å to 1 nm spatial range. As reported in Figure 6, in this spatial range EET is faster than CT mechanism and the time scales obtained are in good agreement with those observed in the experiments, with EET in the ultrafast time regime (from few fs to hundreds of fs, Figure 3) and the CT, or the charge separation process, in the picosecond time scales. The short range EET is due to Dexter coupling that out-competes both Förster and CT. According to the theory developed by Scholes and Ghiogino, the Dexter exchange integral can be well represented as a through-configuration excitation resonance interaction term where the overlap integrals between neutral-excited (Ψ(D/A), Ψ(D/A*)), and anionic-like (Ψ(D/A), Ψ(D/A*)) electronic wavefunctions play a major role, giving the short range dependence contribution. Due to interface level alignment of type II, amongst ionic-like configurations, only Ψ2 is however expected to play a major role in the overall wavefunction superposition.

Calculations based on such equations support the experimental observation that EET is faster than CT (at the molecular level). Here the equations 1 and 2 are used just to support the experimental observation that EET is faster than CT mechanism and the time scales obtained are in good agreement with those observed in the experiments, with EET in the ultrafast time regime (from few fs to hundreds of fs, Figure 3) and the CT, or the charge separation process, in the picosecond time scales. The short range EET is due to Dexter coupling that out-competes both Förster and CT.

In conclusion we report here the dynamics occurring in the finely intermixed phase of P3HT and PCBM. We show that the expected charge generation process is mediated by an initial ultrafast electronic energy transfer from the polymer to the acceptor molecule. The initial energy transfer (D* + A → D + A*) takes place in less than 200 fs. Charge separation is a secondary process occurring between the excited acceptor and the polymer domain on the 100 ps time scale. Such intermixed domains are the actual border regions between D and A clusters in the photovoltaic active layer based on BHJ. In this frame electronic energy transfer should be an important event in the primarily step for charge generation, usually underestimated in the description of the photovoltaic event. This, first of all, brings to some considerations on the role and actual values assigned so far to the exciton diffusion length. The critical length scale of 10 nm we found here is actually the convolution among the exciton diffusion, the molecular inter-diffusion and the average distance which allow for electronic energy transfer at the interface. Finally we note that Ward et al. have recently shown a possible energy transfer pathway from copolymers such as PCDTBDT to PCBM. Thus, the process we highlight here is not necessarily specific to P3HT/PCBM, but it may be considered as an unavoidable paradigm for the concept of organic photovoltaic.

Methods
Sample preparation. The poly(3-hexylthiophene) (P3HT) (Merk, >99% regioregular) was diluted in a dichlorobenzene (DCB) solution at different weighting ratio (1 mg/mL, 5 mg/mL) and heated at 70 °C degrees for 1 hour. The solution was spin coated on a clean 250 µm thick fused silica substrate at 4000 rpm or 1000 rpm for 45 seconds, respectively, in order to get a film thickness of 5 or 50 nm. Note that the freshly spun P3HT films have been dried on a hotplate at 70 °C for 10 min prior to spin-coating the top layer. For the production of the bilayer we adopt the spin-coating sequential approach using an orthogonal solvent combination (dichlorobenzene and dichloromethane) to process the [6,6]-phenyl C61-butyric acid methyl ester (PCBM) layer on top of P3HT preventing the re-dissolution of the P3HT underneath. The PCBM was dissolved in a dichloromethane solution (20 mg/mL), briefly heated at 40 °C to ensure maximal dissolution. The PCBM solution was casted at 4000 rpm for 45 seconds onto the P3HT layer to obtain a thickness of 5 nm. In order to increase the order of the P3HT layer before PCBM deposition we perform a thermal annealing, heating the sample to 140 degrees for 5 minutes.

TrAMP. For the ultrafast experiments, laser pulses (40 fs, 250 kHz) at 800 nm were generated using Coherent Micra Ti:Saphire oscillator in conjunction with Coherent RegA 9040 amplifier. Part of the laser output was used to pump Coherent OPA in order to generate pump pulse at 520 nm. The other part of the pulse was used to pump a sapphire plate (2 mm) to generate white light continuum as the probe pulse. The delay between the pump and probe pulses was controlled by a computer controlled translation stage. Both the pulses were made collinear using a beam splitter before launching into the TrAMP setup. The TrAMP system can be thought to be a misaligned concentric optical cavity; as shown in figure 1, one of the mirrors focuses the beam and the second mirror collimates the beam and sends it back. At proper incidence conditions, a multi-reflection can be achieved between the two mirrors with the ray tracing an ellipse on the sample (as well as on the mirrors). The number of reflections can be controlled by tuning the angle of incidence and the mirror alignment or alternatively, the beam can be extracted from the cavity using another smaller mirror. A high pass filter was used to remove the pump pulse; the pulse probe was dispersed using a monochromator and detected using a lock-in, photo-diode system.
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