Interfacial Morphology and Effects on Device Performance of Organic Bilayer Heterojunction Solar Cells

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Supporting Information

ABSTRACT: The effects of interface roughness between donor and acceptor in a bilayer heterojunction solar cell were investigated on a polymer–polymer system based on poly(3-hexylthiophene) (P3HT) and poly(diocylfluorene-alt-benzothiadiazole) (F8BT). Both polymers are known to reorganize into semicrystalline structures when heated above their glass transition temperature. Here, the bilayers were thermally annealed below glass transition of the bulk polymers (≥140 °C) at temperatures of 90, 100, and 110 °C for time periods from 2 min up to 250 min. No change of crystallinity could be observed at those temperatures. However, X-ray reflectivity and device characteristics reveal a coherent trend upon heat treatment. In X-ray reflectivity investigations, an increasing interface roughness between the two polymers is observed as a function of temperature and annealing time, up to a value of 1 nm. Simultaneously, according bilayer devices show an up to 80% increase of power conversion efficiency (PCE) for short annealing periods at any of the mentioned temperatures. Together, this is in agreement with the expectations for enlargement of the interfacial area. However, for longer annealing times, a decrease of PCE is observed, despite the ongoing increase of interface roughness. The onset of decreasing PCE shifts to shorter durations the higher the annealing temperature. Both, X-ray reflectivity and device characteristics display a significant change at temperatures below the glass transition temperatures of P3HT and F8BT.

KEYWORDS: all-polymer, solar cells, bilayer, floated, interface formation, low temperature annealing

INTRODUCTION

Organic solar cells are a promising alternative to inorganic solar cells, because production costs can be reduced through different large area deposition techniques, like roll-to-roll.¹ Power conversion efficiencies around 10% are achieved right now with blends² or tandem cells.³ Performance of an organic solar cell depends on several parameters; the most important ones are absorption of light to form an exciton, charge separation, charge carrier mobility, and charge collection at the electrodes. Absorption and mobility can be directly influenced by the choice of organic material, but the two other parameters are mainly determined by the device architecture itself. Charge collection can be improved with a better built-in potential or additional hole/electron conduction layers at the anode/cathode side. The charge separation is highly influenced by properties of an interfacial area between the donor and acceptor. This circumstance is related to the short lifetime of the Frenkel excitons, which corresponds to a short diffusion length of about 10 nm. This fact makes the distribution of donor and acceptor domains in the solar cell extremely vital. In the past, plenty of effort has been spent on the control of domain size and distribution in bulk-heterojunction blends by annealing of the organic semiconductors.⁴⁵ Thereby it is generally an accepted opinion that temperatures at or slightly above glass transition of the polymer are required to allow significant motion of polymer chain segments, allowing diffusion or reorientation.⁶⁷ Thereby it has been shown recently that the effective glass transition temperature (Tg) of polymer thin films can occasionally be slightly reduced compared to the bulk induced by free interface effects.⁸

The aim of this work is to determine the effects of low-temperature annealing (far below glass transition) on the interfacial morphology of a polymer–polymer bilayer and their consequences on the characteristics of an according organic solar cell. Investigated was a bilayer system of two semicrystalline polymers, the widely used P3HT as donor (D) and F8BT as acceptor (A). Various studies have already been performed on polymer bilayers to characterize the interfacial morphology,⁹–¹³ thereby solely amorphous systems, annealing above glass transition or the effects of sequential spin-coating with orthogonal solvents have been investigated, but only seldomly these are directly compared to the performance of organic devices.¹¹,⁴⁰ In this work, a sharp bilayer interface is generated by floating technique,³⁹ and the interfacial morphology altered by thermal annealing below glass transition. This is characterized by X-ray reflectivity (XRR) and correlated with

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the device physics of according solar cells. Please note that XRR probes the contrast of electron densities across the interface of different layer materials. Therefore, a distinction between (i) an interdiffusion of polymer chains from one layer into the other layer or (ii) an interface broadening of phase separated polymers cannot be made. As it is common practice for XRR, the interface morphology is given as an interface roughness reporting the average penetration of the two polymer layers into each other.

**EXPERIMENTAL SECTION**

**Sample Preparation.** The substrates (Spectrosil quartz glass, indium–tin-oxide (ITO) coated sodium silicate glass, and plain sodium silicate glass) were cleaned by sonication in methanol, acetone, and isopropanol alcohol (10 min each), followed by oxygen plasma etching (100 W, 10 min). The silicon substrates were treated in the same way except for methanol rinsing and plasma etching. The acceptor polymer poly[9,9-di-n-octylfluorenyl-2,7-diyl-alt-(benzo-[2,1,5]thiadiazole-4,8-diyl)] (F8BT) was provided by Cambridge Display Technology Ltd. with a molecular weight (Mw) of 150 kg mol⁻¹. Poly(3-hexylthiophene-2,5-diyl) (P3HT) was supplied by Rieke Metals Inc. with Mw of 70 kg mol⁻¹ (regioregularity 93%, electronic grade). Chemical structures of the organic semiconductors are shown in Figure 1.

**Figure 1.** Chemical structures of F8BT (left) and P3HT (right).

All solutions of P3HT were prepared from anhydrous chlorobenzene (Sigma-Aldrich) at 100 °C for 20 min with concentrations of 8.5 g/L. F8BT was less soluble than P3HT and therefore stirred for 90 min at 100 °C with a concentration of 7.8 g/L. Films were deposited on Spectrosil quartz substrates for photophysical characterization and microscopy, on native silicon substrates for X-ray characterization, and on Spectrosil quartz substrates for photovoltaic characterization. Single layer samples of P3HT or F8BT were spin-coated from hot solution directly onto the supporting substrate. For bilayer samples, P3HT was always spin-coated as the first layer. Subsequently, the F8BT as the second layer was first spin-coated onto a sodium glass substrate, lift-off in demineralized water and floated onto the P3HT layer to avoid potential material intermixing, which can occur even with an orthogonal solvent. A bilayer prepared with floating guarantees a sharp planar heterojunction interface and is ideal for the investigation concerning the interface morphology. Control samples of single layer F8BT by floating were also prepared to account for potential effects caused by the floating technique. Thickness of the layers was kept constant at 40 nm for P3HT and 60 nm for F8BT (facilitating XRR analysis). Photovoltaic devices were completed by evaporation of a 100 nm silver cathode and 60 nm for P3HT, and of P3HT/F8BT bilayers, pristine, and after annealing at 100 °C for 5 min to investigate changing surface features. Surface roughness was deduced from AFM and was used to correlate with surface roughness of the films determined by XRR. Optical absorbance was measured on single layers and bilayer films deposited on Spectrosil quartz glass with a UV/vis spectrophotometer (Hewlett-Packard HP 8453). Photoluminescence (PL) spectra were measured in a nitrogen-purged integrating sphere under argon ion laser excitation at 515 nm at room temperature. Light was detected with an Oriel InstaSpec IV spectrophotograph. Photoluminescence quantum efficiencies (PLQE) were calculated for the P3HT emission, as described by de Mello et al. Further, PL spectra were also recorded with a Shimadzu RF-5301PC spectrophotometer under incoherent monochromatic light excitation at 400 nm (150 W xenon lamp) at room temperature. For photovoltaic device characterization, current–voltage characteristics were acquired under solar simulated conditions (intensity equivalent to 100 mW/cm², AM 1.5G) using a solar simulator light source (ABET Technologies 10500, specification ABB). Photovoltaic characterization was done for bilayer devices annealed at 90, 100, and 110 °C with an annealing time from 2 to 250 min. For every temperature/annealing step at least 12 devices were investigated.

**RESULTS**

**Interface Characterization.** Figure 2 shows results of XRR for P3HT/F8BT bilayers, pristine, and after heat-treatment at temperatures of 140, 110, 100, and 90 °C. Thickness of the polymer single layers varied for P3HT from 39–43 nm and for F8BT from 61–64 nm, as extracted from XRR fits on single layers (compare the Supporting Information), which led to a predicted total bilayer thickness of 100 to 107 nm. This value was confirmed in XRR by the bilayer characteristic Kiessig-fringes below qz = 0.20 Å⁻¹ as seen in Figure 2, which correspond to the total thickness of the bilayer. Unfortunately, the lack of contrast in electron density between P3HT and F8BT of ρP3HT = 0.347 Å⁻³ and ρF8BT = 0.371 Å⁻³ result in a relatively low reflection coefficient at the P3HT/F8BT polymer interface; therefore, the X-ray beam was not able to differentiate the two layers from another at low qz-values. However, above qz = 0.20 Å⁻¹ fringes from a thinner layer appear, which could be clearly assigned to the P3HT layer in-between the substrate and the F8BT layer. The emergence of P3HT-fringes at higher qz-values is directly connected to the quality of the polymer interface and correlated to the roughness of the interface. This conclusion can be drawn from the fitting parameters obtained by modeling of the XRR data (compare Figure 3). Though the
used P3HT had relatively high molecular weight, lamellar ordering was not entirely suppressed, as indicated by an arising Bragg-peak at \( q_z = 0.38 \text{ Å}^{-1} \), seen more or less distinct in all films of Figure 2.

The fits for pristine as-made floated bilayers indeed indicate a sharp interface, with the determined interface roughness between P3HT and F8BT found negligible with \( \sigma_{\text{RMS}} \approx 0 \text{ nm} \). In the next step, the interface morphology was modified by thermal annealing. A polymer’s glass transition temperature marks an important region of considerable change in its chain mobility. Therefore, a strong impact on the development of a polymer/polymer interface can be expected by annealing around this point. Glass transition is found for bulk-P3HT at \( T_{g,\text{P3HT}} = 141 \text{ °C} \)\textsuperscript{17} and for bulk-F8BT depending on the molecular weight, between \( T_{g,\text{F8BT}} = 125 \text{ °C} \) (for \( M_W = 101 \text{ kg mol}^{-1} \)) and \( T_{g,\text{F8BT}} = 133 \text{ °C} \) (for \( M_W = 220 \text{ kg mol}^{-1} \))\textsuperscript{18} A heat treatment of a P3HT/F8BT bilayer at 140 °C for 15 min results already after this short period in a complete disappearance of the P3HT fringes (Figure 2). This indicates that at temperatures near \( T_g \) the interface roughness increases very fast. At lower temperatures this process is strongly delayed. At 110 °C different time scales of 2, 10, 30, 60, 100, and 160 min were applied, and a continuous decrease of the P3HT-fringes was observed only at longer annealing periods. In Figure 2 exemplarily the XRR-scan for 160 min annealing is shown, which demonstrates the strong reduction of the fringes, though they are still noticeable. When the temperature was reduced to 100 °C, the decay time of the P3HT fringes at high \( q_z \)-values became much longer. Only after applying an annealing period of 250 min, the fringes almost disappeared (Figure 2). For 90 °C, the decay of the P3HT-fringes was further suppressed, and so even after 250 min the P3HT-fringes were still clearly visible. To demonstrate the continuous decay of the P3HT-fringes with annealing time, Figure 3 shows XRR-scans and fits of bilayers in comparison, in pristine condition and after annealing at 100 °C for durations of 2, 20, and 250 min. The fits of these experimental data revealed continuous roughening of the polymer interface from almost zero to \( \sigma_{\text{RMS}} = 1 \text{ nm} \), as shown in the inset of Figure 3 (fit details are given in the Supporting Information). The observed increase of roughness at the P3HT/F8BT interface cannot be ascribed to effects of the glass transition temperature of the polymer bulk materials \( (T_{g,\text{P3HT}}) \) as the used temperatures were definitely below.

To determine if the obvious change of interfacial morphology correlates with a potential increase in crystallinity, GIXD measurements were performed on samples annealed at 100 °C for different periods between 2 and 250 min (compare the Supporting Information). Figure 4 exemplarily shows the integrated diffraction patterns for a pristine and an annealed bilayer at 100 °C for 250 min in comparison. Both diffraction patterns show three distinct features. The first two subtle elevations around \( q_w = 1.10 \text{ Å}^{-1} \) and \( q_w = 1.50 \text{ Å}^{-1} \) are assigned to F8BT’s characteristic side-by-side interchain distance (5.3 Å) and \( \pi \)-stacking (4.18 Å), respectively\textsuperscript{19}. The weak and quite broad signal indicates little degree of order in the film. The third stronger peak is correlated to the interplanar distance between the \( \pi \)-stacked thiophene rings of P3HT at \( q_w = 1.65 \text{ Å}^{-1} \)\textsuperscript{20}. It can be clearly seen that there is no significant change of the crystallinity of the bilayer. The small relative difference of both scans comes from a discrepancy of sample alignment and the use of different samples. Even after annealing close to \( T_{g,\text{Bulk}} \) at 140 °C, the P3HT-peak does not change, neither in the...
GIXD-scans nor in specular scans at $q_z = 0.38 \text{ Å}^{-1}$ (Figure 2). It can be concluded that the crystallinity of the P3HT/F8BT bilayer does not change up to a temperature of 140 °C. However, experiments at 160 °C (annealed for 90 min) revealed a strong increase of crystallinity (not shown).

Figure 5 shows atomic force microscopy height images of a F8BT single layer, a P3HT/F8BT bilayer with F8BT on top, and a P3HT single layer; each surface is shown pristine (top) and after annealing at 100 °C for 5 min (bottom). Both, sole F8BT (Figure 5, left) and the bilayer (middle) show a considerable coarsening of the lateral feature size upon annealing from a few nanometers to up to 40 nm. At the same time, single layer P3HT (Figure 5, right) shows a reverse trend. These observations are reflected in surface roughness ($\sigma_{\text{RMS}}$) determined by AFM image analysis. This is showing a heat-induced increase from 0.38 to 0.56 nm in $\sigma_{\text{RMS}}$ for F8BT single layers and 0.49 to 0.59 nm for bilayer samples. The same heat treatment on a P3HT single layer causes a decrease from 0.86 to 0.67 nm for $\sigma_{\text{RMS}}$. Despite the fact that different to XRR, AFM is a scanning-probe based technique with limited lateral resolution, the surface roughnesses of the pristine samples are in the same range as the corresponding fit parameters of the XRR data, with values of 0.39, 0.72, and 0.93 nm for F8BT, the bilayer, and P3HT, respectively.

Photophysics. The photoluminescence (PL) quenching can be a measure for the efficiency of exciton dissociation at the donor/acceptor interface, which is one essential step toward photocurrent generation. Exciton dissociation is strongly related to the D/A interface characteristics. Figure 6 shows emission spectra of P3HT/F8BT bilayer films, pristine and after annealing (100 °C, 5 min) upon excitation at 515 nm. This wavelength excites exclusively the donor P3HT phase, as visible from the absorbance spectrum of P3HT. The same heat treatment on a P3HT single layer causes a decrease from 0.86 to 0.67 nm for $\sigma_{\text{RMS}}$. Despite the fact that different to XRR, AFM is a scanning-probe based technique with limited lateral resolution, the surface roughnesses of the pristine samples are in the same range as the corresponding fit parameters of the XRR data, with values of 0.39, 0.72, and 0.93 nm for F8BT, the bilayer, and P3HT, respectively.

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Figure 6. P3HT emission from pristine (red squares) and annealed (orange circles) bilayer films upon excitation at 515 nm. Absorption spectrum of pure P3HT (pink triangles) is shown for completeness. (Note: Symbols on the graphs are for indication).
reach a D/A interface to dissociate (within their small diffusion radius) and instead recombine radiatively. Photovoltaic Characteristics. Figure 7 shows photocurrent characteristics under simulated solar illumination (AM1.5G) of three representative P3HT/F8BT bilayer solar cells before and after thermal treatment at 90, 100, or 110 °C for different annealing times. Symbols on the curves assign the corresponding plots.

Figure 7. Photocurrent characteristics of P3HT/F8BT bilayer photovoltaic devices, as-deposited and after thermal treatment at 90, 100, or 110 °C for different annealing times. Symbols on the curves assign the corresponding plots.

The power conversion efficiency (PCE) is a product of fill factor, $J_{SC}$ and $V_{OC}$ divided by incoming power. Figure 8 shows

*Figure 8. Evolution of power conversion efficiency (PCE) of P3HT/F8BT bilayer solar cells as a function of annealing temperature and annealing time. Shown is the PCE normalized to the initial value of the not-annealed device (0 min). The error bars give the variance from all measurements at particular annealing conditions.*
evolution of the device PCE upon annealing of the bilayer for the three temperatures 90, 100, and 110 °C with duration of the heat treatment. For better comparability the PCEs have been normalized to the initial efficiency value of a pristine bilayer device. For each of the annealing temperatures, again a similar trend is seen. The PCE starts from the value of the pristine bilayer and increases with duration of thermal treatment, up to a certain maximum point. The time after which this maximum is reached occurs after shorter durations the higher the temperature, which is after 40 min annealing at 90 °C, 20 min for 100 °C, and after only 8 min at 110 °C. Continued annealing beyond this duration causes in any case a reduction of PCE. Thereby at the highest temperature (110 °C) devices show the steepest drop in PCE, even below the value of the initial pristine bilayer device, while at 90 °C almost plateau-like behavior is observed, only decreasing over a long time. The observed trend in PCE is clearly a result of the superposition of the behavior of $J_{SC}$ and $V_{OC}$.

**DISCUSSION**

Multiple studies reported interface roughening between two polymer layers by thermal annealing at glass transition temperature of the bulk material ($T_g$) in the range of several nanometers. Here, although a clear change of polymer morphology would not be expected by annealing below the polymers $T_g$, X-ray reflectivity clearly revealed increasing interface roughness of the polymer—polymer interface between P3HT and F8BT. Its origin is either molecular interdiffusion generating a nanoscopic mixed phase or caused by microscopic interpenetration of the two phase-separated polymers. However, X-ray reflectivity is not able to distinguish between these two cases. The reason for such an effect occurring below $T_g$ is not completely understood. A reduction of $T_g$ in the order of 10 °C–30 °C was observed near free surfaces of polystyrene (PS) reaching tens of nanometers deep into the material. For other polymers this effect was also observed but weakened by a factor of 3. Different studies reported enhanced segmental mobility of polymers at free surfaces. In the present case, reorganization at the film surface was confirmed with AFM imaging for single P3HT films (refinement) and single F8BT films (coarsening) after 100 °C annealing (see Figure 5). However, at the bilayer interface, being no free surface, such strong changes are not expected; but the X-ray reflectivity results in Figure 2 and Figure 3 show a clear increase of the interface roughness, if annealed below $T_g$. Thereby the interface roughens faster (i.e., after shorter annealing time), the higher the temperature. Photophysical studies show clear quenching behavior of the P3HT photoluminescence by the presence of the F8BT acceptor layer in the pristine samples. However, thermal annealing of the bilayer causes increase in photoluminescence. As the single P3HT films’ PL do not change with annealing, this must originate from diminished PL quenching at the D/A interface. Further, there are the device characteristics of the solar cells. The power conversion efficiency shows a particular trend, characterized by an initial increase with annealing time, followed by a drop. The maximum of the PCE thereby shifts to shorter annealing durations with increasing temperature (40 min @ 90 °C, 20 min @ 100 °C, and 8 min @ 110 °C). Thereby the PCE trend clearly reflects the superposition of photocurrent and open circuit voltage evolution. For the possible origin of the changes in $J_{SC}$ and $V_{OC}$ with thermal annealing below $T_g$, the following effects could be responsible: (1) degradation of the active layer, (2) changes at the electrode interface (affected by contact resistance, interfacial area between active layer at the electrode), (3) decrease in charge mobility by molecular reorganization, and (4) change of the D/A interfacial area. There was no degradation of the active layers with ongoing annealing, as this would have shown as diminished PL also for sole P3HT films. Variations at the contact (e.g., by dewetting of the polymer at the inorganic interface) can be excluded since the observed fill factor is thermally invariant, which is a direct measure for the interfacial charge-transfer at the contact. The charge carrier mobility of the individual polymer layers changes only modestly upon thermal annealing below $T_g$ because substantial molecular reorganization is still suppressed. Different studies have shown that charge mobility of F8BT does not change by annealing at 100 °C and even begins to decrease at $T_g$. For P3HT, annealing temperatures below $T_g$ cause, as in this case, the charge carrier mobility by less than 1 order of magnitude. Only if P3HT is annealed at/above $T_g$, its hole-mobility increases considerably. This is also supported by XRD and XRR, as neither of them showed significant modifications in P3HT or F8BT at these temperatures. In conclusion, the evolution of the interface roughness must be directly and exclusively responsible for the observed development in $J_{SC}$ and $V_{OC}$ and PCE in consequence. It is suggested that at the onset of interface roughening of the bilayer there is indeed an increase of interfacial area between donor and acceptor, which has a direct beneficial impact on the amount of photocurrent generated, visible as increase of $J_{SC}$. The $V_{OC}$ on the other hand, is only at maximum for the most sharp pristine interface, as the probability for flow of recombination currents (loss mechanism) is at minimum. The ongoing increase of interface roughness beyond a certain point (max. $J_{SC}$) seems to create a morphology at the D/A interface, which no longer enhances photocurrent generation. Lowered $J_{SC}$ and $V_{OC}$ together with the decreased PL quenching indicates an increased rate of recombination. This usually occurs when the distance between a photogenerated exciton and the D/A interface (required for dissociation and generation of charge pairs) exceeds its diffusion length (in the range 10 nm). With regard to the bilayer, this would imply that with ongoing interface roughening beyond the point of maximum device performance, too large inversions or even migrated separated droplets of one material within the other might have formed. However, as the highest observed interface roughness by XRR is 1 nm, this remains unclear.

**CONCLUSIONS**

The polymer bilayer of F8BT and P3HT was thermally annealed at temperatures below the glass transition temperature ($T_g$ ∼ 140 °C) of both polymers used, namely at 90, 100, 110, and 140 °C. No change of the crystallinity was observed in this temperature region. However, X-ray reflectivity showed a decrease of Kiessig fringes, which corresponds directly to the morphology at the bilayer interface. A continuous decrease of those fringes was observed for longer annealing periods (from 2 to 250 min) indicating a continuous increase of the interface roughness from almost zero to 1 nm. This effect of interface roughness appears at smaller annealing times when higher temperatures were applied. Also the photocurrent characteristics of the organic solar cells showed a serious change in the power conversion efficiency (PCE). In a first step an increase of up to 80% in PCE was measured, the reached maximum in PCE
shifted from 40 min annealing at 90 °C to 20 min annealing at 100 °C, to only 8 min annealing at 110 °C. After the observed maximum, an extension of the annealing time leads to a decrease of PCE, which is more pronounced at higher temperatures. Interestingly, the defined alteration of the device characteristics occurs at temperatures below the bulk-glass transition and therefore cannot be explained by changes of the charge carrier mobility but merely by the detected increase of the interface roughness.

**ASSOCIATED CONTENT**

Supporting Information

X-ray reflectivity investigations together with fit of the experimental data are given for the single layers of P3HT and PBTT as well as for the bilayer in the pristine state and annealed at 100 °C for 2, 20, and 120 min. All parameters of the fits are given. Grazing incidence X-ray diffraction of a pristine bilayer and of bilayers annealed at 100 °C between 2 and 250 min. Atomic force microscopy phase images of the single layers of P3HT and PBTT and of a bilayer in the pristine state and annealed at 100 °C for 5 min. Statistics in the device performance for devices annealed at 90, 100, and 110 °C for annealing times between 2 min up to 250 °C. The Supporting Information is available free of charge on the ACS Publications website at DOI: 10.1021/acsami.5b04972.

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**Notes**

The authors declare no competing financial interest.

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