Efficient Organic Photovoltaics Utilizing Nanoscale Heterojunctions in Sequentially Deposited Polymer/fullerene Bilayer

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A highly efficient sequentially deposited bilayer (SD-bilayer) of polymer/fullerene organic photovoltaic (OPV) device is developed via the solution process. Herein, we resolve two essential problems regarding the construction of an efficient SD-bilayer OPV. First, the solution process fabrication of the SD-bilayer is resolved by incorporating an ordering agent (OA) to the polymer solution, which improves the ordering of the polymer chain and prevents the bottom-layer from dissolving into the top-layer solution. Second, a non-planar heterojunction with a large surface area is formed by the incorporation of a heterojunction agent (HA) to the top-layer solution. Poly[[9-(1-octylnonyl)-9H-carbazole-2,7-diyl]-2,5-thiophenediyl-2,1,3-benzothiadiazole-4,7-diyl-2,5-thiophenediyl] (PCDTBT) is used for the bottom-layer and phenyl-C71-butyric-acid-methyl ester (PC70BM) is used for the top-layer. The SD-bilayer OPV produced utilizing both an OA and HA exhibits a power conversion efficiency (PCE) of 7.12% with a high internal quantum efficiency (IQE). We believe our bilayer system affords a new way of forming OPVs distinct from bulk heterojunction (BHJ) systems and offers a chance to reconsider the polymers that have thus far shown unsatisfactory performance in BHJ systems.

Organic photovoltaic (OPV) devices have received intense attention due to their low-cost manufacturing¹, ease of processing² and mechanical flexibility³. These OPVs utilize a bulk heterojunction (BHJ) type photoactive layer where the electron donating polymer and electron accepting fullerene are blended into one layer to form a heterojunction in the bulk film. In order to achieve a large heterojunction area for efficient exciton dissociation, nanoscale phase separation between the polymer and fullerene is desired. Therefore, the BHJ system requires a delicate control of the processing conditions including the blending ratio between the polymer and fullerene, the amount of processing additives, the kinds of solvents used and the thermal annealing temperature.

Analogous to its use in inorganic solar cells, the bilayer structure of a p-type polymer and an n-type fullerene can be utilized for OPVs. A bilayer structure can be prepared by the sequential deposition of polymer and fullerene solutions. Usually, there is interdiffusion of polymer and fullerene at the polymer/fullerene interface in the sequentially deposited bilayer (SD-bilayer) of polymer/fullerene. Nevertheless, compared to the blending system, finding the optimum conditions for a SD-bilayer is more straightforward. However, two major requirements first need to be addressed for the satisfactory application of the SD-bilayer to OPVs. First, the top-layer solvent should not destroy the bottom-layer. Since considering the thickness of active layer in OPV devices is usually less than 100 nm, the bottom-layer should have little solubility in the top-layer solvent. Second, the interfacial area at the top-layer/bottom-layer heterojunction should be large enough for efficient exciton dissociation.

Recently, several interesting results regarding a poly(3-hexylthiophene) (P3HT)/phenyl-C61-butyric acid methyl ester (PC60BM) SD-bilayer OPV have been reported⁴⁻⁹. Dichloromethane (DCM) was used as the solvent for the PC60BM top-layer because DCM does not destroy the P3HT bottom-layer. It was found that partial
both an OA and top-layer solutions, respectively. The SD-bilayer OPV utilizing PCDTBT bottom-layer (Film 1), SD-bilayer of PCDTBT/PC70BM (Film 2) and PCDTBT bottom-layer after removing the PC70BM top-layer from the solar cell efficiency and stability when it forms a BHJ with polymers in DCM. For example, PCDTBT is known to exhibit high solubility for the PC70BM top-layer into the P3HT bottom-layer normally annealed P3HT/PC60BM SD-bilayer showed comparable efficiency to that of a BHJ OPV device. Recently, an organic solar cell based on a PTB7/PC70BM SD-bilayer was reported\(^\text{16}\). However, the power conversion efficiency (PCE) of the solar cell was found to be around 3%, which is significantly lower than the PCE of a solar cell utilizing a PTB7:PC70BM BHJ.

The proper solvent for SD-bilayer OPV fabrication must be considered as DCM is not always appropriate to use in every SD-bilayer device of polymer/fullerene due to the various solubilities of different polymers in DCM. For example, PCDTBT is known to exhibit high solar cell efficiency\(^\text{11}\) and stability\(^\text{12,13}\) when it forms a BHJ with PC70BM. However, even though the solubility of PCDTBT in DCM is low (0.54 mg/mL), the solvent almost completely removes the thin PCDTBT film (50 nm), and consequently, constructing a SD-bilayer of PCDTBT/PC70BM is impossible when using a DCM solution.

In this work, we report an efficient SD-bilayer solar cell based on a PCDTBT/PC70BM. We have resolved two essential problems: the construction of the SD-bilayer by a solution process and the formation of a heterojunction with a large surface area by introducing an ordering agent (OA) and a heterojunction agent (HA) to the bottom- and top-layer solutions, respectively. The SD-bilayer OPV utilizing both an OA and a HA shows PCE of 7.12% with the internal quantum efficiency (IQE) over 90%.

**Results and discussions**

We investigated whether DCM, the solvent for the PC70BM top-layer, dissolves the PCDTBT bottom-layer by spin-coating DCM onto the PCDTBT layer. The PCDTBT layer was nearly completely removed after spin-coating with DCM. This is ascribed to the amorphous (or less ordered) feature of the PCDTBT chains. In the P3HT/PC60BM SD-bilayer OPV, domains of highly ordered P3HT chains prevent the dissolution of P3HT by blocking DCM penetration. This implies that polymer chains should form ordered domains to successfully construct a polymer/PC60BM SD-bilayer using the sequential solution process. Unfortunately, many of the recently developed low band gap polymers, including PCDTBT, are amorphous due to the bulky alkyl side chains required for the solubility enhancement of the polymers\(^\text{14-16}\).

In this respect, two approaches for enhancing the ordering of the polymer chains were executed. First, thermal annealing of the polymer at a proper temperature\(^\text{17,18}\) can enhance ordering. Recently, X. Lu et al.\(^\text{19}\) reported that ordered PCDTBT domains were created after thermal annealing at 200°C. Second, utilizing an OA assists in the formation of ordered polymer domains. Proper OA selection depends upon the OA having a lower vapor pressure and little solubility for the polymer when compared with the host solvent. These properties allow the OA, when added to the host solvent, to give the polymer chains more time to achieve an ordered state by reducing the evaporation rate of the host solvent. In addition, when the host solvent has evaporated, the remaining OA (poor solvent for polymer) aids in the precipitation of the polymer leading to the formation of ordered polymer domains. We have found several suitable OAs for PCDTBT. 1,8-Diodooctane (DIO) was chosen as one OA for PCDTBT due to its negligible vapor pressure and the fact that PCDTBT is insoluble. The PCDTBT was dissolved in 99 vol% of chlorobenzene (CB) and 1 vol% of DIO. For clarity, the PCDTBT layers prepared with this solvent system (1 vol% of DIO) and those prepared without DIO will be denoted as PCDTBT(OA) and PCDTBT(N), respectively, and the thermally annealed PCDTBT will be denoted as PCDTBT(N-T).

Figure 1 shows the absorption intensity change for the PCDTBT films with different preparation methods (Film 1 of Figure 1a). The black lines in Figure 1b, c and d are absorption spectra of as-prepared PCDTBT(N), PCDTBT(OA) and PCDTBT(N-T) films, respectively. All the films show similar absorption intensity regardless of the preparation method. This absorption intensity shows that the thicknesses of the PCDTBT films were not affected by the addition of an OA or by thermal annealing. Subsequently, the PC70BM DCM

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**Figure 1** | Preparation procedure of films and the UV-visible absorption spectra of PCDTBT films. (a) Graphical representation of the as-prepared PCDTBT bottom-layer (Film 1), SD-bilayer of PCDTBT/PC70BM (Film 2) and PCDTBT bottom-layer after removing the PC70BM top-layer from the SD-bilayer (Film 3). (b) UV-visible absorption spectra of PCDTBT(N), (c) UV-visible absorption spectra of PCDTBT(N-T), and (d) UV-visible absorption spectra of PCDTBT(OA). Black lines represent the absorption of Film 1. Red lines represent the absorption of Film 3.
solution was spin-coated on the PCDTBT bottom-layer to form the PC_{70}BM top-layer (Film 2 of the Figure 1a). To verify the thickness of the PCDTBT bottom-layer after spin-coating with PC_{70}BM solution, the PC_{70}BM top-layer was selectively removed using diiodomethane (DIM) which is a good solvent for PC_{70}BM and a non-solvent for PCDTBT (Film 3 of the Figure 1a). Red lines in Figure 1 show the absorption spectrum of the corresponding PCDTBT bottom-layer after removing the PC_{70}BM top-layer. The absorption spectra and the photographs in Figure 1b indicate that the PCDTBT(N) was almost completely removed by the PC_{70}BM top-layer solution. This is in contrast to the PCDTBT(N–T), which was insoluble in the PC_{70}BM solution (Figure 1c). Figure 1d indicates that a considerable amount (65.8%) of the PCDTBT(OA) bottom-layer remained even after spin-coating with the PC_{70}BM solution.

To investigate the ordering and orientation of the polymer chains in the PCDTBT layer, grazing incidence X-ray diffraction (GIXD) experiments were performed. Figure 2a–c shows 2D GIXD images of the PCDTBT films prepared from the different processing conditions. Two broad peaks were observed for the PCDTBT(N) and PCDTBT(OA). According to X. Lu et al., the peak at around 1.5 Å⁻¹ (4.2 Å) corresponds to the π–π stacking distance and the peak in the range of 0.355–0.406 Å⁻¹ (17.7–15.5 Å) corresponds to the spacing between the polymer chains. Additional peaks between 0.5 Å⁻¹ and 1.0 Å⁻¹ along the q_z direction were observed from PCDTBT(N–T). These peaks are related to the second- and third-order reflections showing long range ordered structures parallel to the substrate. Figure 2d compares out-of-plane scattering profiles extracted from the 2D GIXD patterns of corresponding films. The peaks marked with an asterisk are caused by indium tin oxide (ITO) diffuse scattering. Their crystallographic information is summarized in Table 1. The calculated coherence length along the q_z axis (surface normal) increased in the order of PCDTBT(N) < PCDTBT(OA) < PCDTBT(N–T). The coherence lengths of PCDTBT in PCDTBT(N–T) and PCDTBT(OA) are 380% and 48.3% larger than that of PCDTBT in PCDTBT(N), respectively. The out-of-plane scattering analysis of the PCDTBT layers revealed that the PCDTBT(N) did not show significant PCDTBT chain ordering and the PCDTBT(OA) showed enhanced chain ordering along the q_z direction. Additionally, the PCDTBT(N–T) showed significant enhancement in the PCDTBT chain ordering along the q_z direction. These results suggest that including the OA in the PCDTBT solution enhances the chain ordering of PCDTBT implying that DIO plays a vital role in the chain ordering of PCDTBT. Consequently, the ordered PCDTBT domains established by the OA will be less soluble in the PC_{70}BM solvent, which corresponds well with the UV-visible absorption results in Figure 1. This clearly shows that the incorporation of an OA to the PCDTBT solution or the thermal annealing of the PCDTBT film enables the SD-bilayer of PCDTBT/PC_{70}BM to be fabricated by the solution process.

The SD-bilayer of PCDTBT/PC_{70}BM was successfully prepared by sequential spin-coating with PCDTBT and the PC_{70}BM solution. The constructed SD-bilayer films were utilized as the active layer in an OPV with the device structure ITO/PEDOT: PSS/PCDTBT/PC_{70}BM/TiO₂/Al. Table 2 compares the performance of the SD-bilayer OPV. For clarity, PC_{70}BM layers prepared with DCM will be denoted as PC_{70}BM(N). Three different SD-bilayers of PCDTBT(N)/PC_{70}BM(N), PCDTBT(N–T)/PC_{70}BM(N) and PCDTBT(OA)/PC_{70}BM(N) were used as the active layer of a SD-bilayer OPV, and the solar cell utilizing PCDTBT(OA)/PC_{70}BM(N)
showed the highest PCE (1.82%) among the three types of devices. The PCE difference was mainly ascribed to the short circuit current density ($J_{SC}$). However, all SD-bilayer solar cells exhibited similar PCEs, and the values are significantly lower than that of BHJ solar cells, which range from 6.4 to 7.1%20–22.

There are two possible reasons for these results. First, the low PCEs could be ascribed to the low solubility (4.47 mg/mL) measured for PC$_{70}$BM in DCM. The solubility is too low for the conformal formation of the PC$_{70}$BM top-layer on the PCDTBT bottom-layer with a proper thickness. The second reason could be due to the volatility of DCM. The highly volatile DCM will evaporate quickly leaving large domains of PC$_{70}$BM on the PCDTBT bottom-layer. To investigate this hypothesis, the surface of SD-bilayer of PCDTBT/PC$_{70}$BM was scanned using atomic force microscopy (AFM). The scanning electron microscopy (SEM) image of the SD-bilayer surface clearly showed large aggregations of PC$_{70}$BM on the PCDTBT bottom layer with various diameters (Figure 3a). The features of PC$_{70}$BM domains are exactly matched with the surface morphologies of the PCDTBT bottom-layer after removal of the PC$_{70}$BM top-layer from the SD-bilayer films of PCDTBT/PC$_{70}$BM (Figure 3b–d). The large holes with a diameter of around 200–800 nm were observed in all three SD-bilayer films, suggesting that the large domains of PC$_{70}$BM were formed on the PCDTBT bottom-layer during coating with the PC$_{70}$BM top-layer and left indentations when the PC$_{70}$BM domains were removed with DIM. The domain size was significantly larger than the exciton diffusion length, which would reduce the exciton dissociation efficiency. This could be a major reason for the poor performance of the PCDTBT/PC$_{70}$BM SD-bilayer OPVs without concerning the various PCDTBT bottom-layers. In addition, the low fill factor (FF) of the solar cell reflects a significant charge recombination due to an imbalance between electrons and hole transport.

| Bottom-layer (Donor layer) | Top-layer (Acceptor layer) | $V_{OC}$ (V) | $J_{SC}$ (mA/cm$^2$) | FF | PCE (%) |
|---------------------------|---------------------------|-------------|------------------|----|---------|
| PCDTBT(N)                 | PC$_{70}$BM(N)            | 0.84        | 2.68             | 0.48 | 1.09    |
| PCDTBT(N–T)               | PC$_{70}$BM(N)            | 0.87        | 3.02             | 0.42 | 1.10    |
| PCDTBT(OA)                | PC$_{70}$BM(N)            | 0.94        | 4.27             | 0.45 | 1.82    |
| PCDTBT(N)                 | PC$_{70}$BM(HA)           | 0.88        | 5.18             | 0.63 | 2.88    |
| PCDTBT(N–T)               | PC$_{70}$BM(HA)           | 0.91        | 7.99             | 0.51 | 3.70    |
| PCDTBT(OA)                | PC$_{70}$BM(HA)           | 0.90        | 12.02            | 0.66 | 7.12    |

Figure 3 | Surface morphologies of the SD-bilayer of PCDTBT(OA)/PC$_{70}$BM obtained by SEM (a) and PCDTBT bottom-layers (b–d) obtained by AFM. (a) PCDTBT(OA)/PC$_{70}$BM SD-bilayer, (b) PCDTBT(N) bottom-layer after removal of the PC$_{70}$BM top-layer from the PCDTBT(N)/PC$_{70}$BM film, (c) PCDTBT(N–T) bottom-layer after removal of the PC$_{70}$BM top-layer from the PCDTBT(N–T)/PC$_{70}$BM and (d) PCDTBT(OA) bottom-layer after removal of the PC$_{70}$BM top-layer from the PCDTBT(OA)/PC$_{70}$BM. All the images were obtained by AFM in tapping mode (2.5 μm × 2.5 μm).
In order to address these shortcomings, conformal formation of the PC70BM top-layer on the PCDTBT bottom-layer without significant destruction of the PCDTBT bottom-layer is critical. To satisfy this requirement, we introduced a HA that helps conformal formation of the PC 70BM top-layer without destroying the PCDTBT bottom-layer. The following properties are desirable when considering a suitable HA: good solubility for PC 70BM with low vapor pressure and little solubility for PCDTBT. In this context, we identified several HAs for the PC 70BM top-layer. Among the HAs, DIM was found to be appropriate. PC 70BM is highly soluble in DIM whereas PCDTBT is not soluble. Additionally, the vapor pressure of DIM is low enough for the conformal formation of the PC 70BM top-layer on the PCDTBT bottom-layer. According to J. S. Moon et al., DIO and DIM are known to inhibit excessive crystallization of the fullerene, which will reduce the roughness of the PC 70BM film. The solubility of PC 70BM was greatly improved to above 15 mg/mL by adding 5 vol% of DIM to the PC 70BM DCM solution. The DIM containing solutions exhibited good film forming properties compared to DCM only solutions. The surface roughness of the PCDTBT/PC 70BM SD-bilayer film was remarkably reduced by adding DIM to the PC 70BM solution. Hereafter, the PC 70BM top-layer prepared from the DCM solvent with 5 vol% of DIM will be denoted as PC70BM(HA). The root mean square roughness ($R_q$) of PCDTBT(OA)/PC70BM(N) (27.5 nm) in Figure 3a. This clearly shows the effect of HA on SD-bilayer film roughness.

In addition to reducing the surface roughness of the PCDTBT/PC 70BM SD-bilayer film, maximizing the interfacial area of the PCDTBT/PC 70BM heterojunction is desirable when constructing an efficient SD-bilayer OPV. To investigate the heterojunction morphologies of PCDTBT/PC 70BM SD-bilayers, cross-section transmission electron microscopy (TEM) was conducted on the PCDTBT(N–T)/PC70BM(HA) and PCDTBT(OA)/PC70BM(HA) films. Clear interface formation between PCDTBT and PC 70BM was not observed in the TEM image of both SD-bilayers when the image was taken at focusing conditions (Figure 5a and 5e). The PCDTBT/PC70BM interfaces became distinguishable after underfocusing the SD-bilayer samples to $-15 \mu m$. The prepared PCDTBT(N–T)/PC70BM(HA) SD-bilayer showed a clear polymer/PC 70BM interface suggesting the formation of a planar heterojunction (Figure 5b). However, a polymer/PC 70BM interface was not clearly observed for the PCDTBT(OA)/PC70BM(HA) SD-bilayer (Figure 5f). The cross-section TEM sample was mapped for sulfur and carbon by energy dispersive X-ray spectroscopy (EDS) (Figure 5c, 5d, 5g and 5f). Since only PCDTBT contains sulfur, the PCDTBT polymer is expected to lie where sulfur is observed. Sulfur was found only in the PCDTBT layer in PCDTBT(N–T)/PC70BM(HA), which is coincident with the cross-section TEM result. In the PCDTBT(OA)/PC70BM(HA) SD-bilayer film, how-
ever, sulfur was found everywhere, and there was a distinct sulfur distribution gradient. The sulfur intensity was most intensive in the PCDTBT bottom-layer, and the intensity decreased in the PC70BM upper-layer. Considering that the spatial resolution of the EDS is larger than the polymer domain size and the AFM image taken of the PC70BM removed PCDTBT layer, it is thought that the PCDTBT and PC70BM domains are interdigitated to form a non-planar heterojunction with a large surface area.

For further investigation into the heterojunction morphology of the PCDTBT/PC70BM SD-bilayers, the PC70BM top-layer was selectively removed from the SD-bilayer films using DIM (refer the Film 3 of Figure 1a). And the surface morphology of the PC70BM-removed PCDTBT bottom-layer were scanned using AFM and shown in b–d of Figure 4.

Considering both the morphology of the PC70BM(N) top-layer deposited on the PCDTBT(OA) bottom-layer (Figure 3a) and the surface morphology of the PCDTBT(OA) bottom-layer after removal of the PC70BM(N) top-layer (Figure 3d), it was expected that the selective removal of PC70BM by DIM would not significantly alter the morphology of the PCDTBT bottom-layer because DIM does not dissolve PCDTBT but only selectively dissolves PC70BM. We have investigated the effect of the PC70BM removal process on the morphology of the PCDTBT bottom-layer in detail. UV-visible absorption measurement of the DIM solution revealed no absorption from PCDTBT but only absorption due to PC70BM (Figure S1 of Supplementary Information). However, it is difficult to completely remove the PC70BM from the SD-bilayer because the PC70BM is supposed to be inter-diffused into the PCDTBT bottom-layer during the PC70BM top-layer deposition process. To investigate the amount of residual PC70BM left on the PCDTBT bottom-layer, photoluminescence (PL) measurement of the DIM solution after excluding the PC70BM removed PCDTBT films were performed. As shown in Figure S2a and b of the Supplementary Information, PL quenching of ~24% was observed for both the PCDTBT(N–T) and PCDTBT(OA) films after PC70BM removal. Based on the PL quenching experiment for the PCDTBT:PC70BM films with various blending ratios (Figure S2c and d of Supplementary Information), the 24% PL quenching indicates only 0.1 wt% of PC70BM (PC70BM = 999 : 1) remained in the PCDTBT(N–T) and PCDTBT(OA) bottom-layers after PC70BM removal with DIM. However, the amount of residual PC70BM is thought to be negligible compared to the amount of PCDTBT. Therefore, it is expected that the obtained surface morphology of PCDTBT bottom-layer without any residual PC70BM.

However, there is another factor that can influence PCDTBT film morphology. It is possible that DIM can cause the PCDTBT to swell changing its morphology. To investigate this, we have examined the surface of the PCDTBT film after several DIM treatments. The PCDTBT film was immersed in the DIM solution for 10 seconds, and the surface morphology of the PCDTBT film was examined by AFM. As expected, no noticeable change in the film morphology was observed even after repeated DIM treatments (Figure S3 of Supplementary Information).

Based on the above experiments, it was concluded that there is no significant change in the morphology of the PCDTBT bottom-layer during the selective PC70BM removal process, and that the surface morphology of Figure 4b–d represents the morphology of the heterojunction formed at the PCDTBT/PC70BM interface.

There was no significant difference in the surface roughness for the PC70BM(HA) top-layer removed PCDTBT(N), PCDTBT(T) and PCDTBT(OA) bottom-layers (3.21 nm, 1.74 nm and 3.37 nm, respectively). However, a significant difference in the surface area was found among the PCDTBT bottom-layers. The surface areas of the films were calculated from the AFM images after excluding the geometric area (1 μm²) from the total surface area. The surface areas of PCDTBT(N) and PCDTBT(N–T) are similar, whereas the surface area of PCDTBT(OA) is 6 times larger than the PCDTBN bottom-layer and 5 times larger than the PCDTBN bottom-layer. This represents a 5–6 fold increase in the heterojunction area of the PCDTBT(OA)/PC70BM(HA) SD-bilayer compared to the PCDTBN/PC70BM(HA) and PCDTBN/PC70BM(HA) SD-bilayers. In addition, the surface of the PCDTBT(OA) bottom-layer showed a well developed nanoscale morphology compared to the PCDTBN and PCDTBN–T bottom-layers. The width and height of the PCDTBN domains are around 50 nm and 15 nm, respectively. Considering the outcome of our morphological investigation and surface area calculations at the PCDTBN/PC70BM(HA) interface, the PCDTBN(OA)/PC70BM(HA) is expected to form the most effective non-planar heterojunction for exciton dissociation among the SD-bilayer films.

Cross-section TEM was performed on the PC70BM removed PCDTBT bottom-layers (Figure 6). Aluminum of 10 nm thickness...
was thermally evaporated onto the PC$_\text{70BM(HA)}$ top-layer removed PCDTBT(N), PCDTBT(N–T) and PCDTBT(OA) bottom-layers to discern PCDTBT from the background. The TEM images were contracted to the scale of the AFM images (The original TEM images of the films can be found in Figure S4 of Supporting Information), and the TEM results corresponded perfectly with the AFM results. TEM images of the PC$_\text{70BM(HA)}$ top-layer removed PCDTBT bottom-layers showed relatively flat profiles for the PCDTBT(N) and PCDTBT(T) bottom-layers, whereas a rough profile was observed for the PCDTBT(OA) bottom-layer. Thus, TEM results support the formation of a non-planar heterojunction with nanoscale morphology in the PCDTBT(OA)/PC$_\text{70BM(HA)}$ SD-bilayer film, which is in agreement with the AFM height profile analysis.

Based on the TEM analysis of the SD-bilayers, PCDTBT(N–T)/PC$_\text{70BM(HA)}$ forms a SD-bilayer with little inter-diffusion, whereas the PC$_\text{70BM}$ in the PCDTBT(OA)/PC$_\text{70BM(HA)}$ SD-bilayer can penetrate into the amorphous PCDTBT domain and form an inter-diffused bilayer. Consequently, the PCDTBT(N–T)/PC$_\text{70BM(HA)}$ SD-bilayer forms a planar heterojunction, and the PCDTBT(OA)/PC$_\text{70BM(HA)}$ SD-bilayer forms a non-planar heterojunction with a large surface area.

Figure 7 compares the performance of solar cells having a PCDTBT/PC$_\text{70BM}$ SD-bilayer processed with an OA and HA. Their photovoltaic parameters are listed in Table 2. The solar cell performance was significantly enhanced by the incorporation of the HA to the PC$_\text{70BM}$ solution. As expected from the AFM and TEM results, the OPV having a PCDTBT(OA)/PC$_\text{70BM}$ SD-bilayer showed the highest PCE of 7.12% with $J_{\text{SC}}$ of 12.02 mA/cm$^2$, $V_{\text{OC}}$ of 9.80 V, and FF of 0.66. The average efficiency of 6.73 ± 0.14% was obtained from 30 devices. This clearly shows the effect of the HA as the OPV fabricated without any HA, PCDTBT(OA)/PC$_\text{70BM}$, only exhibited an efficiency of 1.82%. The difference in PCEs of the SD-bilayer OPVs was mainly attributed to the difference in $J_{\text{SC}}$.

Among the SD-bilayer OPVs having a HA processed top-layer, the PCDTBT(N)/PC$_\text{70BM(HA)}$ SD-bilayer showed the lowest $J_{\text{SC}}$, and the PCDTBT(OA)/PC$_\text{70BM(HA)}$ SD-bilayer OPV showed the highest $J_{\text{SC}}$.

The $J_{\text{SC}}$ of a solar cell is affected by the overall carrier dynamics, such as the light absorption efficiency ($\eta_{\text{LD}}$), the exciton dissociation efficiency ($\eta_{\text{ED}}$), and the charge collection efficiency ($\eta_{\text{CC}}$). Comparing $\eta_{\text{A}}$ of SD-bilayers, the $\eta_{\text{A}}$ of the PCDTBT(N)/PC$_\text{70BM}$ film will be the lowest among the three different bilayer films. As it was confirmed by the absorption spectra (Figure 1) and TEM (Figure 6a), significant amounts of PCDTBT was removed during the formation of the PC$_\text{70BM(HA)}$ top-layer due to an insufficient ordering of the PCDTBT chains in the PCDTBT(N) bottom-layer.

PL measurements on the SD-bilayer films were carried out to investigate the $\eta_{\text{LD}}$ of the SD-bilayer films (Figure S5 of Supplementary Information). The PL of PCDTBT films was quenched to 88.5% and 89.3% after construction of PCDTBT(N)/PC$_\text{70BM}$ and PCDTBT(OA)/PC$_\text{70BM}$ SD-bilayer films, respectively. The PL of PCDTBT(N–T)/PC$_\text{70BM}$ SD-bilayer film, however, shows 81.6% quenching compared to the PL of PCDTBT film. Considering the fact that little amount of PC$_\text{70BM}$ in PCDTBT film significantly quenches PL intensity of PCDTBT film (Figure S3 of Supplementary Information), the difference in PL intensity between PCDTBT(OA)/PC$_\text{70BM}$ and PCDTBT(N–T)/PC$_\text{70BM}$ is not substantial. Based on the GIXD and TEM results, the thermal annealing enhances the ordering of PCDTBT and results in increased size of ordered PCDTBT domains that prevent penetration of PC$_\text{70BM}$ into PCDTBT layer. As a result, the PCDTBT(N–T)/PC$_\text{70BM}$ SD-bilayer forms a relatively flat interface compared to the PCDTBT(OA)/PC$_\text{70BM}$ SD-bilayer, which was also confirmed by surface area calculation results in Figure 4c and d. This lowers the $\eta_{\text{LD}}$ of PCDTBT(N–T)/PC$_\text{70BM}$ SD-bilayer films.
Consequently, the inferior \( \eta_{\text{sc}} \) of PCDTBT(N)/PC\(_{70}\)BM(HA) and the inefficient \( \eta_{\text{sc}} \) of PCDTBT(N-T)/PC\(_{70}\)BM(HA) is considered to be the main reason for the lower \( J_{\text{sc}} \) of corresponding SD-bilayer OPVs compared to that of PCDTBT(OA)/PC\(_{70}\)BM(HA) OPV. Although the high efficiency of PCDTBT(OA)/PC\(_{70}\)BM(HA) is thought to be ascribed to non-planar heterojunction formation with high interfacial area, other factors could also be responsible including changes in the carrier mobility of material on either side of the junction, changes in the interfacial recombination rate due to differences in the orientation of the polymer chains, or changes in the exciton diffusion length.

The maximum external quantum efficiency (EQE) of 72.8% was obtained at 540 nm (Figure 7b). The EQE and PCE values of the SD-bilayer OPV are comparable to the reported EQE and PCE values obtained from the PCDTBT/PC\(_{70}\)BM BHJ OPV\(^{1,12}\). The IQE of the PCDTBT(OA)/PC\(_{70}\)BM(HA) OPV device was calculated considering the internal absorption obtained by the reflectance measurement on the device (Figure 7c). The detailed process for the IQE of the SD-bilayer OPV device is described in Methods section. Briefly, the IQE value of the OPV device was obtained after dividing IQE values of the device by the calculated absorption of the active layer only. The calculated IQE value of the OPV device is in the range of 90% to 100%, which indicates a very efficient charge collection ability for the SD-bilayer OPV device.

Interestingly, the optimized thickness of the PCDTBT bottom-layer of the PCDTBT(OA)/PC\(_{70}\)BM(HA) SD-bilayer is only around 20 nm (Figure 6c). Considering the optimum thickness of the PCDTBT:PC\(_{70}\)BM layer in BHJ OPVs was found to be 80–100 nm, the thickness of the PCDTBT bottom-layer of our SD-bilayer OPV seems relatively thin. However, when considering that the blending ratio of PCDTBT:PC\(_{70}\)BM for a BHJ is around 1:4, the amount of PCDTBT in our SD-bilayer and the BHJ is similar. The absorption spectrum of the PCDTBT/PC\(_{70}\)BM SD-bilayer film almost coincides with that of the BHJ film (PCDTBT:PC\(_{70}\)BM = 1:4). Figure S6 of Supporting Information). Furthermore, the absorption intensities of PCDTBT films after removing the PC\(_{70}\)BM from the BHJ film and SD-bilayer film are similar. This implies the amount of PCDTBT in the SD-bilayer film is similar with that in the BHJ film. In addition, the reflectance data of the PCDTBT(OA)/PC\(_{70}\)BM(HA) SD-bilayer device obtained by absorption measurement using a reflective mode clearly shows that light in the range of 350–600 nm is sufficiently absorbed by the PCDTBT(OA)/PC\(_{70}\)BM(HA) SD-bilayer (Figure S7 in Supporting Information).

Besides DIO and DIM, we have also tested 1-chloronaphthalene, 3-methylthiophene, 3-hexylthiophene and 1,8-octanedithiol for use as the OA or HA in the production of the PCDTBT/PC\(_{70}\)BM SD-bilayer OPV, and the results are summarized in Figure S8, Figure S9, Table S1 and Table S2 of Supporting Information. Among the PCDTBT/PC\(_{70}\)BM SD-bilayer OPVs utilizing OA and HA, PCDTBT(DIO)/PC\(_{70}\)BM(DIM) showed the highest PCE.

In conclusion, we have developed a SD-bilayer system of PCDTBT/PC\(_{70}\)BM having a non-planar heterojunction with a high interfacial area via solution processes, and we have applied this concept to the construction of highly efficient SD-bilayer OPVs. The construction of the PCDTBT/PC\(_{70}\)BM SD-bilayer was achieved by adding the OA, for example DIO, to the polymer solution. The OA improved the ordering of the PCDTBT chains and reduced the solubility of PCDTBT in the top-layer solution. The formation of a nanoscale non-planar heterojunction was achieved by adding the HA, for example DIM, to the PC\(_{70}\)BM solution. The HA enabled the conformal deposition of the PC\(_{70}\)BM top-layer onto the PCDTBT bottom-layer and enabled the formation of the large area heterojunction. The SD-bilayer OPV utilizing an OA and HA shows a PCE of 7.12%. We expect that this method can be applied to polymers that are not adequate for use in the BHJ system due to the large surface energy difference between the polymer and fullerene as this difference is not significant in the SD-bilayer system. We believe that our SD-bilayer system establishes an alternative to the BHJ system for the fabrication of efficient OPVs, and also allows for the reassessment of those polymers which have shown insufficient performance in a BHJ system.

**Methods**

**Device fabrications.** Pre-patterned ITO-coated glass substrates (20 Ω/□) were cleaned by ultrasonic treatment for 10 min in isopropyl alcohol (IPA), acetone and IPA again then dried at 100 °C in a convection oven for 30 min. The cleaned substrates were treated with UV ozone for 20 min. The poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate) (PEDOT: PSS) (Clevios™P VP AI 4083, Germany) layer was spin-coated onto the cleaned substrates with a speed of 4000 rpm for 35 s and then dried at 100 °C in a vacuum oven for 10 min. A PCDTBT (1-Material, Canada) solution (8 mg/mL) was dissolved in CB without OA or with 1 vol% of OA. For the OA, DIO (TCI, Japan) was used. The PCDTBT layer was dried at 100 °C in a vacuum oven for 15 min after the layer was spin-coated onto the PEDOT: PSS layer with a speed of 2500 rpm for 30 s. A PC\(_{70}\)BM (Nano-C, USA) solution (6 mg/mL) was prepared in DCM without HA or with 5 vol% of HA. DIM was used for the HA. The PC\(_{70}\)BM layer was spin-coated on top of the PCDTBT layer with a speed of 4000 rpm for 30 s and subsequently dried at 100 °C in a vacuum oven for 15 min. TiO\(_2\) nanoparticles were dispersed in n-butanol with 0.4 wt% TiO\(_2\) interlayer was deposited onto the PC\(_{70}\)BM layer by spin-coating at 1500 rpm for 30 s and dried at 60 °C on a hot plate for 30 min. A 100 nm-thick Al metal electrode was thermally evaporated under a pressure of 3.0 × 10\(^{-4}\) Torr.

**Measurements.** Current density-voltage characteristics were measured by a Keithley 2400 source meter under AM 1.5 G irradiation (100 mW/cm\(^2\)) from a 150 W Xenon lamp based solar simulator (McScience, Korea). Incident photon-to-current efficiency (IPCE) was measured at different wavelengths with a monochromatic light generated from a 300 W Xenon lamp in the range of 300 nm to 800 nm using K3010 EQX (McScience, Korea). UV-visible spectrophotometer UV-2450 (SHIMADZU, Japan) is used to obtain the absorbance spectrum of the films. AFM images were recorded in non-contact mode using XE-70 (Park Systems Corp., Korea) and TEM
images were obtained using Tencal™ G2 E20 (FEI, USA) with 200 kV accelerating voltage. GIXD measurements were conducted at PLS-II 9A U-SAXS beamline of Pohang Accelerator Laboratory (Korea). The X-rays coming from the in-vacuum undulator (IVU) are monochromated (wavelength $\lambda = 1.114 \AA$) using a double crystal monochromator and focused both horizontally and vertically (450 (H) × 60 (V) $\mu$m² at FWHM at sample position) using K-B type mirrors. GIXD sample stage is equipped with a 7-axis motorized stage for the accurate alignment and the incidence angle of X-ray beam was set to 0.11° in this study. GIXD patterns were recorded with a 2D CCD detector (Rayonix SX165, USA) and X-ray irradiation time was 5–10 s dependent on the saturation level of detector. Diffraction angles were calibrated by a crystal monochromator and focused both horizontally and vertically (450 (H) × 60 (V) $\mu$m²) and the sample-to-detector distance was about 243 mm3. To estimate the absorption of active layers only and the IQE transfer-matrix formalism (TMF) was used19–22. The multilayer structure was assumed to be glass/TTO (120 nm)/PEDOT:PSS (30 nm)/PCDTBT:PCBM (1:4 w/w, 80 nm)/TiO₂ (8 nm)/Al (100 nm). The refractive indices of glass and Al were obtained from Ref. 28 and Ref. 29, respectively, and those of other layers were acquired by ellipsometry.

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