Determining Ultralow Absorption Coefficients of Organic Semiconductors from the Sub-Bandgap Photovoltaic External Quantum Efficiency

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Energy states below the bandgap of a semiconductor, such as trap states or charge transfer states in organic donor–acceptor blends, can contribute to light absorption. Due to their low number density or ultrasmall absorption cross-section, the absorption coefficient of these states is challenging to measure using conventional transmission/reflection spectrophotometry. As an alternative, the external quantum efficiency (EQE) of photovoltaic devices is often used as a representative of the absorption coefficient, where the spectral line shape of the EQE is considered to follow the absorption coefficient of the active layer material. Here, it is shown that the sub-bandgap EQE is subject to thickness dependent low finesse cavity interference effects within the device—making this assumption questionable. A better estimate for the absorption coefficient is obtained when EQE spectra corresponding to different active layer thicknesses are fitted simultaneously for one attenuation coefficient using an iterative transfer matrix method. The principle is demonstrated for two model acceptor–donor systems (PCE12-ITIC and PBTTT-PC71BM) and accurate subgap absorption coefficients are determined. This approach has particular relevance for both understanding subgap states and their utilization in organic optoelectronic devices.

1. Introduction

Electro-optical modeling has been extensively used to design and optimize the performance of optoelectronic devices such as solar cells,[1,2] photodetectors,[3,4] and thin-film light-emitting diodes (LEDs).[5] For example, an accurate electro-optical model can help to increase the photocurrent produced by a solar cell,[6] tune the spectral response of a photodetector,[7] and increase the outcoupling efficiency of thin-film LEDs.[8] To perform these optimizations, a full knowledge of the complex refractive indices of all active and buffer layers of the device structure is required. For photon energies $h\nu > E_g$ the complex refractive index is often determined through transmission and reflection spectrophotometry or spectroscopic ellipsometry. The same experimental techniques become very challenging when applied to the sub-bandgap optical constants, particularly because for $h\nu < E_g$ the absorbance is typically orders of magnitude weaker than for $h\nu > E_g$. Throughout this work, the absorbance correctly refers to the ratio of absorbed to incident light power, whereas absorption is the physical process of photoexcitation through light–matter interaction. Precise sub-bandgap optical constants are, however, crucial for modeling a variety of semiconductor devices operating at energies below the bandgap such as near-infrared (NIR) photodetectors[9,10] or light emitting exciplex LEDs.[11] In addition, the spectral line shape of the absorption coefficient for $h\nu < E_g$ is indicative of a variety of phenomena such as disorder in amorphous semiconductors and charge transfer absorption in organic semiconductors. Disorder-induced and charge transfer state absorptions are related to loss mechanisms in organic solar cells, and have been studied extensively to improve solar cell efficiencies.[12–14] Given the considerable interest in the sub-bandgap absorption coefficients, sensitive methods of measuring it have been developed. This includes photothermal deflection (PDS) spectroscopy as a powerful technique for measuring absorption coefficients down to 0.001 cm$^{-1}$ in the best case, but more commonly down to 1 cm$^{-1}$ in condensed matter phases.[15–18] Similar to ellipsometry, PDS requires the preparation of single layer samples and a rather complicated data analysis subject to model fitting. Another technique is Fourier transform photocurrent spectroscopy (FTPS) that is based on recording the photocurrent action spectrum using a Fourier Transform Infrared spectrometer. FTPS is often used to determine low energy trap states in inorganic semiconductors.[19,20] One should note that photocurrent-based methods may be more sensitive than PDS but can only probe the absorbance of the photocurrent-generating species. In amorphous silicon, it has been shown that FTPS systematically underestimates the optical absorbance for $h\nu < 1.4$ eV when compared with PDS.[21,22] Similar to FTPS, the external quantum efficiency (EQE) is another photocurrent-based method commonly used to characterize the photocurrent response of solar...
cells and photodetectors over the visible (VIS) and NIR spectral range. The EQE is the product of absorptance \( A(\lambda) \) of the active layer as a function of wavelength and the internal quantum efficiency \( \text{IQE} \), such that

\[
\text{EQE}(\lambda) = A(\lambda) \times \text{IQE}
\]

As a consequence of Equation (1), the spectral line shape of the EQE must be directly transferrable to \( A \) assuming that the IQE does not feature any substantial energy dependence. The latter assumption has been proven in technologically relevant high efficiency systems such as efficient bulk heterojunctions and perovskite solar cells. Nevertheless, examples for donor–acceptor systems with an excitation energy dependent IQE have been shown recently. If the spectral line shape of the EQE changes with applied bias, the IQE is not spectrally flat and \( A \) can be more precisely inferred from the EQE measured at the largest possible bias to ensure photocurrent saturation.\(^\text{[28,30]}\)

The active layer absorptance spectrum \( A(\lambda) \) depends on the absorption coefficient \( \alpha(\lambda) \), but also the wave-optics of the multilayer stack device that is the solar cell or the photodetector. In the case of an optically thin layer with \( \alpha \ll 1 \) (layer thickness \( t \)) and in the limit of negligible cavity effects (such as a semiconductor film deposited on a transparent substrate), it is often assumed that \( A \approx \alpha t \) for \( h\nu < E_g \).\(^\text{[24,31–33]}\) Therefore the spectral line shape of the EQE should follow \( \alpha \) via \( A \). However, the assumption of \( A \approx \alpha t \) is invalid in the presence of electrodes as the device forms a low-finesse cavity with an associated thickness dependence. The well-known cavity effects are typically observed as pronounced interference fringes in the above-gap EQE (where \( \alpha t \approx 1 \)) but can have a similar effect on the subgap EQE. The prominence of these cavity effects depends primarily on the relative difference between the individual layer refractive indices. A higher mismatch of the refractive index between the organic and transparent conducting electrode interface causes substantial Fresnel reflection making the respective electrode partially reflective. This can be quite a subtle effect, but important in regions of low attenuation coefficients and/or high dispersion in the refractive index—both circumstances encountered below the bandgap again making \( A \approx \alpha t \) a questionable approximation. A better estimate for \( A \) is obtained by modeling the optical field distribution within a device, which requires knowledge of the optical constants (that is the real and imaginary components of the refractive index) for all materials constituting the layers of the device. In the so-called transfer matrix method, the electric field is described as a position- and wavelength-dependent matrix considering the absorption and reflection of every layer and interface.\(^\text{[34]}\) \( A(\lambda) \) of the active layer is then calculated as the sum of the position \((x)\)-dependent modulus squared of the electric field \( E(\lambda, x) \) over the layer thickness \( t \) multiplied by the absorption coefficient \( \alpha \) and the refractive index \( n \)

\[
A(\lambda) = \alpha(\lambda) n(\lambda) \sum_{x=0}^{t} |E(\lambda, x)|^2
\]

It is a relatively straightforward task to model a thin film solar cell when all optical constants are known but challenging when the sub-bandgap optical constants are unknown. In this work, we determined absorption coefficients as low as \( 10^{-2} \text{ cm}^{-1} \) in the subgap spectral range of two exemplary acceptor–donor blends often used in high efficiency organic solar cells. The full molecular structures of these model systems, PBTTT-PC\(_{71}\)BM and PCE12-ITIC are provided in the Supporting Information. Solar cells with active layer thicknesses between 60 and 375 nm were fabricated to study the effect of thickness on the shape of the EQE spectra in the subgap spectral range. The absorption coefficients were obtained by numerically fitting for multiple EQE spectra and including thickness-dependent optical field simulations via an iterative transfer matrix method. We validated the modelled coefficients by simulating EQE spectra of devices with different thicknesses where we find a good agreement between the simulated and experimental results. It becomes clear that the EQE line shape is strongly dependent on the active layer thickness and hence assuming that \( A \approx \alpha t \) can only be true for a small range of thicknesses. In the PCE12-ITIC system, the interference-induced thickness dependence of EQE in the subgap regime is more pronounced due to the large refractive index of this material system. These results indicate that one cannot directly relate the subgap EQE spectra to the subgap absorption coefficient without performing appropriate optical simulations. This is particularly important in systems with larger refractive index such as the recently introduced nonfullerene donor/acceptor blends.

2. Results
2.1. Optical Constants
PBTTT-PC\(_{71}\)BM (1:4) and PCE12-ITIC (1:1) were chosen as model material systems for investigations of subgap ultralow attenuation coefficients. PBTTT-PC\(_{71}\)BM is known for its extended subgap EQE shoulder due to charge transfer state (CT) absorption,\(^\text{[35]}\) while PCE12-ITIC shows a narrow CT state-related-EQE contribution that is barely distinguishable from the above-gap EQE caused by singlet excitations.\(^\text{[33]}\) First, we obtained the optical constants from single layer samples of spin-coated material with thicknesses ranging between 100 and 400 nm on both glass and silicon substrates. Ellipsometry is considered as an indirect method for obtaining the optical constant, as it measures polarization parameters \( \Psi(\lambda) \) and \( \cos \Delta(\lambda) \), which are then globally fitted for the film thickness \( t \) and refractive index \( n \) in the transparent region between 1200 and 1600 nm (\( \chi = 0 \), i.e., the Cauchy region). Thereafter, \( n \) and \( \chi \) are fitted stepwise over the entire spectrum with fixed \( t \). Figure 1 shows the optical constants obtained for PCE12-ITIC and PBTTT-PC\(_{71}\)BM. We find that the ellipsometric data could be reliably determined in good agreement between the experimental data and the mathematical model (mean-squared error \(<1\%\)) as well as high reproducibility between samples in the VIS spectral range. The dashed lines in Figure 1 indicate the sensitivity limit for determining \( \chi \), which roughly corresponds to the bandgap energy beyond which this method is not applicable to \( \chi \) (as seen from the divergence of \( \chi \) for different \( t \)). At the same time, the relative change in \( n \) with \( t \) in the Cauchy regime compared to \( \chi \) is insignificant, as \( n \) lacks spectral features and converges to a constant value around...
2. In summary, we used $n$ as obtained from ellipsometry for transfer matrix simulations in the entire spectral range but dismissed $\kappa$ for wavelengths beyond the dashed lines illustrated in Figure 1. We further determined the optical constants of indium tin oxide (ITO), MoO$_3$, and ZnO experimentally, while relying on reported tabulated data for Ag$^{[36]}$. As a next step, electrically inverted photovoltaic cells with the structure ITO (105 nm)/ZnO (37 nm)/active layer/MoO$_3$ (7 nm)/Ag (130 nm) were fabricated with different active layer thicknesses (PCE12-ITIC: 60, 132, 215, 375 nm and PBTTT-PC$_{71}$BM: 60, 81, 110, 140, 190 nm).

2.2. Numerical Determination of $\kappa$

The EQE spectra are used in the procedure depicted in the process flowchart of Figure 2 to obtain ultralow attenuation coefficients. The full protocol was implemented in MATLAB and is provided in the Supporting Information. The numerical basis for this process is the Nelder–Mead simplex algorithm that minimizes a nonlinear and constrained objective function.$^{[37]}$ Importantly, the presented method does not depend on the algorithm used for the minimization. Other derivative-free algorithms like the Brent’s method or Powell’s conjugate direction method can also be employed.$^{[38]}$ To numerically determine the attenuation coefficient $\kappa$, two steps were followed as shown in Figure 2a,c. (1) Estimation of the precise active layer thickness for the single device (defined as the “pixel”) and (2) calculation of the attenuation coefficient $\kappa$ under consideration of the pre-optimized thickness. First, the absorption $A$ of each device is modeled by the transfer matrix method. The experimental input data required for the modeling is the measured active layer thickness $t_0$ and the optical constants in the VIS spectral range that is from 400 to 600 nm for PBTTT-PC$_{71}$BM and 400–700 nm for PCE12-ITIC. Thereafter, IQE$_{VIS}$ and the mean (IQE$_{mean}$) are obtained using Equation (1). From the available ellipsometric constants, values were chosen that resulted in the minimum deviations of the IQE$_{VIS}$ from its mean. The IQE of one exemplar PCE12-ITIC device (375 nm active layer layer) is depicted in Figure 1b showing a deviation of the IQE of no more than 13 % from its mean. Such deviations can be caused by thickness variations over the device area which are common for solution-processed layers and could even be seen with the naked eye for the 210 nm PBTTT-PC$_{71}$BM device. We therefore preoptimized the active layer thickness $t$ within ±10 nm boundaries around the experimental thickness $t_0$. To do so, the absorbance $A'$ is calculated from the IQE$_{mean}$ as $A' = EQE \times IQE^{-1}_{mean}$. Figure 2b shows that the absolute difference between $A'$ and $A$ is largest where the IQE deviates most from its mean. To match $A'$ to $A$, $t$ is numerically varied to minimize the function $(A' - A)^2 / A$. In the case of $A = A'$, IQE$_{VIS}$ would be spectrally flat, which is impossible to achieve given the experimental errors and uncertainties related to optical constants and thicknesses of other device layers. Nevertheless, the IQE spectra remains approximately flat within these unavoidable errors. IQE spectra of the devices under investigation are shown in Figures S1 and S2 of the Supporting Information. The experimental and the numerically optimized thicknesses of all the investigated devices are shown in Table S1 of the Supporting Information. Once the optimized thicknesses and IQE$_{mean}$ for all devices are obtained, the absorption spectra below the bandgap can be estimated by $A'' = EQE \times IQE^{-1}_{mean}$ using the experimental EQE over the full spectral range. Importantly, $A''$ is now a matrix containing the estimated absorbance of all devices. Figure 2c shows the process of numerically obtaining $\kappa$. The fixed input comprises the absorption matrix $A''$, the experimental attenuation coefficient at the last wavelength within the VIS spectral range $\kappa(\lambda_{m-1})$, the attenuation coefficient $\kappa$ over the full spectral range, and the numerically obtained thicknesses $t$. The optimization of $\kappa$ proceeds wavelength by wavelength starting from the $\lambda_m$ where $\lambda_m$ is the last wavelength within the VIS spectral range. At $\lambda_m$, the transfer matrix calculates $A''$ using $\kappa(\lambda_{m-1})$ as the initial guess of the attenuation coefficient. To further match $A''$ to $A''(\lambda_{m})$, the $\kappa$ is varied between 0 and $\kappa(\lambda_{m-1}) + \kappa(\lambda_{m-1})/10$ with the goal to minimize the function $(A''(\lambda_{m}) - A''(\lambda_{m}))^2 / A''(\lambda_{m})$. Once a predefined number of iterations has been exceeded, $\kappa(\lambda_{m})$ is fixed and the algorithm proceeds to the next wavelength $\lambda_{m+1}$, choosing $\kappa(\lambda_{m})$ as the initial estimate for fitting $\kappa(\lambda_{m+1})$.

2.3. EQE Simulations

A computer code for this method is provided in the Supporting Information with exemplar data. The presented method is applicable to a single or more devices of a different active layer thickness. However, inaccuracies in the determined thickness and small deviations from the flat IQE assumption will have a large impact on the numerically obtained attenuation coefficient if only one device is considered. Therefore, it will be more
accurate to numerically fit devices with different thicknesses. From the globally obtained attenuation coefficient, the active layer absorptance can be simulated and then scaled by IQEmean to obtain the EQE. It is important to consider that electronic effects, such as the charge collection imbalance of thick junctions, are not considered in the optical model.

Figure 3a,b illustrate the absorption coefficients, the experimental EQE (EQEexp) and simulated EQE (EQEsim) spectra of PCE12-ITIC and PBTTT-PC71BM. The agreement between simulation and experiment for different active layer thicknesses was quantified by calculating the relative percentage difference. EQEsim is typically within 40% deviation around the absolute EQEexp despite the simulation covering seven orders of magnitude. Moreover, the difference in EQEexp with active layer thickness in the sub-bandgap region is mostly larger than the deviation of the simulation around EQEexp. Consequently, EQEsim spectra follow the overall trend of the thickness dependent EQEexp spectra. To exemplify this, the maximum difference in EQE of 60 and 215 nm PCE12-ITIC devices in the sub-bandgap region is 20-fold, whereas the fit-error for one thickness is at most two times the experimental EQE, i.e., a maximum relative percentage error of 100%. The strongest deviations of EQEsim from EQEexp (65% and 75%) occur in the spectral range of the bandgap (210 nm PBTTT-PC71BM and 60 nm PCE12-ITIC) and at the sensitivity limit of the EQE measurement (e.g., 60 nm PCE12-ITIC and 60 nm PBTTT-PC71BM above 1300 nm). From a Gaussian fit to the absorption coefficient of PBTTT-PC71BM, a charge transfer energy (E_{CT}) of 1.16 eV and a reorganization energy (\lambda) of the donor–acceptor complex of 163 meV were determined as shown in Figure S3 of the Supporting Information.

E_{CT} of PBTTT-PC71BM was previously reported to be 1.15 eV in a thin solar cell, where the active layer absorptance A in the device was approximated by A \approx \alpha t.\textsuperscript{[32]} As both E_{CT} values are in excellent agreement, we performed the Gaussian fit on EQEexp spectra for different active layer thicknesses under the assumption of A \approx \alpha t.\textsuperscript{[32]} The 190 nm PBTTT-PC71BM, could not be reliably fitted, while other devices with active layer thicknesses of 60, 81, 110, and 140 nm resulted in E_{CT} and \lambda values ranging from 1.15 to 1.18 eV and 143 to 186 meV, respectively (see Table S2, Supporting Information). Given the

Figure 2. Process flow for obtaining ultralow attenuation coefficients in the weakly absorbing spectral range using numerical optimization. a,b) The active layer thickness t is optimized such that the difference between A and A’ in the visible spectral range is minimal resulting in a spectrally flat IQEvis. c) From the mean of the IQE and sensitively measured sub-bandgap EQE, an absorptance matrix A” for several device thicknesses is calculated. Using the preoptimized t, \kappa is numerically obtained by recursive simulation of the active layer absorptance A”(\lambda_m) for all device thicknesses at one wavelength with the goal to fit A’.
dependence of the fit parameters on the chosen fitting range, an uncertainty of ±10 meV for \(E_{CT}\) and ±20 meV for \(\lambda\) is estimated. To conclude this phase of the analysis, the assumption of \(A = \alpha t\) leads to an insignificant error in determining \(E_{CT}\) and reorganization energy for the material system PBTTT-PC\(_{71}\)BM for layer thicknesses below 190 nm. For PCE12-ITIC devices, the thickness dependent EQE features above 1000 nm were identified as resonances of the device acting as a low-finesse cavity. The resonance modes are highly dependent on the active layer thickness as expected. To eliminate the possible effect of higher order harmonics derived from the illumination system (see the Experimental Section), the EQE of the 190 nm device was remeasured with long-pass filters placed in front of the sample. No significant changes in the EQE were observed as depicted in Figure S5 of the Supporting Information. We should also note that the recorded EQE spectra are all above the electrical noise level of the apparatus as shown in Figure S5 of the Supporting Information. In Figure 3a,b, the numerically obtained \(\alpha\) is scaled to compare its spectral line shape to the EQE spectra of devices with different active layer thicknesses \(t\). The assumption \(A = \alpha t\), according to which the spectral line shape of \(\alpha\) follows \(A\) and therefore the EQE, is not applicable to any PCE12-ITIC device without greatly over- or underestimating \(\alpha\). To find the thickness at which this assumption is valid, we simulated the active layer absorptance \(A = \alpha t\) can give a better estimate for the numerical \(\alpha\) over a wide range of thicknesses. At the same time, the change in thickness between 60 and 150 nm also affects the spectral line shape of the simulated absorptance. From these two examples, it becomes clear that active layer thickness plays a significant role in the observed spectral line shape of the EQE and care must be taken to not mistake material properties with device properties.

### 2.4. Trap States

Figure 3 further indicates a broad feature in the absorption coefficient spectrum of PCE12-ITIC between 1000 and 1400 nm with an ultralow absorption coefficient of 0.1–0.01 cm\(^{-1}\). According to the empirically established relation \(E_{CT} \approx qV_{OC} + 0.6\) eV, the CT state energy of PCE12-ITIC is expected to be around 1.5 eV. This has been experimentally shown\(^{[39,40]}\) and can be observed in Figure 3 as a shoulder in the absorption coefficient spectrum at 800 nm. The corresponding Gaussian-distributed absorptance of CT states occurs at much higher energies than the observed tail states. Importantly, the ability to determine absorption coefficients between 0.1 and 0.01 cm\(^{-1}\), allows one to observe ultraweak optical transitions that are typically associated with defect absorption in this spectral range.\(^{[21,22,41]}\) Moreover, the tail of the absorption coefficient observed here can be easily fitted by an exponential function. Similar to FTPS measurements, the presented method determines exclusively photocurrent-generating trap states. Therefore, it possibly underestimates the optical absorption coefficient in the spectral range of defect absorption. However, the thus obtained “effective absorption coefficient” is advantageous for modeling solely photocurrent-based devices like solar cells and photodetectors, and to perform detailed balance analysis to determine the open circuit voltage of the solar cells and thermodynamic limit of the detectivity of the photodetectors.
2.5. Minimal EQE

Using the presented method, the sensitivity to the absorption coefficient depends solely on the ability to measure small photocurrents in the weakly absorbing spectral region. Figure S5a of the Supporting Information shows the QE of PCE12-ITIC cells measured up to 1700 nm. Depending on the device noise current and measurement integration time, the noise floor is reached at different wavelengths, but always above 1400 nm. In Figure S5b of the Supporting Information, the current response of a 105 nm thick PCE12-ITIC device in the dark is depicted as the noise-equivalent QE of the measurement system utilized in this work. We believe it is therefore possible to obtain $\alpha$ values down to $10^{-3}$ cm$^{-1}$ if the device noise is decreased via (for example) increasing the shunt resistances (i.e., lowering the thermal noise). Importantly, this $\alpha$ is only valid for charge-generating states. Another limitation of this method concerns finding the accurate optical constants via spectroscopic ellipsometry needed to determine the IQE. Anisotropy or thickness dependent morphology as well as birefringent optical properties can be very difficult to model. Moreover, it has been reported that some material systems show an excitation energy dependent IQE.$^{[21,28]}$ In that case the sub-bandgap IQE cannot be accurately predicted when modeling the device using the as-obtained absorption coefficients. While the charge transfer energy of the PBT TT:PC$_{71}$BM was determined as previously reported, sub-bandgap states were observed for the PCE12-ITIC blend, which are most likely linked to photocurrent generating states. The method presented here refines how to obtain the absorption coefficient from EQE spectra while offering a novel way to study sub-bandgap absorption with relatively low experimental overhead.

3. Conclusion

In summary, we have presented a numerical and experimental approach to determine ultrasmall absorption coefficients of subgap states from external quantum efficiency spectra and applied it to organic semiconductors-based solar cells. Due to non-negligible cavity effects, the spectral shape of the sub-bandgap EQE is influenced by interference fringes that are greater when the refractive index is more dispersive as exemplified with the PCE12-ITIC system. The thickness-dependent EQE spectra of PCE12-ITIC and PBT TT:PC$_{71}$BM devices were accurately predicted when modeling the device using the as-obtained absorption coefficients. While the charge transfer energy of the PBT TT:PC$_{71}$BM was determined as previously reported, sub-bandgap states were observed for the PCE12-ITIC blend, which are most likely linked to photocurrent generating states. The method presented here refines how to obtain the absorption coefficient from EQE spectra while offering a novel way to study sub-bandgap absorption with relatively low experimental overhead.

4. Experimental Section

Device Preparation: On a precleaned glass substrate with a prestructured layer of 105 nm ITO, a 35 nm thin layer of ZnO was deposited via a sol–gel spin-coating at 4000 rpm from a 0.5 M solution of zinc acetate dihydrate in ethanolamine and 2-methoxy ethanol (volume ratio 3:97). The as-spun layer was annealed at 150 °C for 10 min in air. The ZnO hole-transporting layer was followed by the active layer comprising either PBT TT:PC$_{71}$BM or PCE12-ITIC. For PBT TT:PC$_{71}$BM as the active layer, PBT TT (Sigma-Aldrich, $M_w > 50$ 000 g mol$^{-1}$, PDI < 3)) and [6,6]-Pheny lC$_{61}$butyric acid methyl ester (PC$_{71}$BM) was mixed in a weight ratio of 1:4 and dissolved in chloroform:1,2-dichlorobenzene (6:4 volume ratio). The solution was stirred overnight, filtered through a 0.2 μm PTFE filter in the cold and spin-coated at 65 °C for 60 s. For film thicknesses between 100 and 200 nm, the concentration of the solution was chosen to be 32 mg mL$^{-1}$ and the spin-speed varied between 1000 and 1500 rpm. For PCE12-ITIC as the active layer, the polymer PCE12 (PBDB-T) and the acceptor ITIC were mixed in a weight ratio of 1:1 and dissolved in chlorobenzene at 50 °C overnight to form a 20 mg mL$^{-1}$ solution. 0.5 vol% diiodooctane was added prior to spin-coating, while keeping the solution at 50 °C throughout the deposition process. Film thicknesses between 60 and 375 nm were obtained by varying the spin-speed between 400 and 2700 rpm and subsequent thermal annealing at 160 °C for 10 min. The active layer was followed by 7 nm of MoO$_3$ and 130 nm of silver thermally evaporated under high vacuum. All devices were defined by the geometrical overlap of the bottom and the top contact that equals 15 mm$^2$. To avoid exposure to ambient conditions, the organic part of the device was covered by a small glass substrate glued on top.

Ellipsometry: The optical constants of spin-coated films were determined experimentally by spectroscopic ellipsometry (instrument: J.A. Woollam M-2000; software: CompleteEASE 5.23). Samples comprised a single layer of material spin-coated from the same solution as utilized for device fabrication on either quartz glass or silicon substrates.

Sensitive EQE Measurements: A Spectrophotometer (LAMBDA 950, PerkinElmer) with an integrated monochromator was used as a light source spanning 400–1800 nm. The output light from the spectrometer was chopped using a Thorlabs MC2000B chopper at 273 Hz and calibrated using a photodiode. The short-circuit current derived from the device fed to a current preamplifier (Femto DHCPA-100) before being analyzed with a lock-in amplifier (SR860 Stanford Research Systems). To resolve low photocurrents, the preamplifier was set to high amplification, while the time-constant of the lock-in amplifier was chosen to be 1 s in the strongly absorbing and 30 s in the weakly absorbing spectral range. For the calibration process, a NIST-calibrated silicon photodiode (for wavelength between 400 nm and 1100 nm) and a germanium photodiode (for wavelength between 780 nm and 1800 nm) from Newport were used.

Supporting Information

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

Acknowledgements

This work was funded through the Welsh Government’s Sêr Cymru II Program ‘Sustainable Advanced Materials’ (Welsh European Funding Office – European Regional Development Fund). C.K. and S.Z. are recipients of a UKRI EPSRC Doctoral Training Program studentship. P.M. is a Sêr Cymru II Research Chair and A.A. a Rising Star Fellow also funded through the Welsh Government’s Sêr Cymru II ‘Sustainable Advanced Materials’ Program (European Regional Development Fund, Welsh European Funding Office, and Swansea University Strategic Initiative).

Conflict of Interest

The authors declare no conflict of interest.
Keywords
absorption coefficient, external quantum efficiency, organic photodetectors, organic solar cells, sub-bandgap states

Received: September 12, 2019
Revised: October 21, 2019
Published online: November 22, 2019

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