Organic Photodetectors and their Application in Large Area and Flexible Image Sensors: The Role of Dark Current

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1. Introduction

1.1. Organic Photodetectors for Large Area Image Sensors

Organic photodetectors (OPDs) are increasingly attractive for light sensing applications as they combine a wide absorption spectrum and high photogeneration yield with low fabrication costs, lightweight, and flexibility.[1,2] Compared to amorphous silicon detectors, OPDs promise many important advantages, most notably that their compatibility with the thermomechanical properties of plastic enables large area image sensors to be solution-processed on a variety of flexible substrates.[3] By using industrially scalable coating techniques such as slot-die coating, OPD arrays can be processed from solution at a lower temperature than amorphous silicon[4] (typically less than 150 °C), thus paving the way towards lower production costs. Furthermore, to integrate OPDs in large-area image sensors, no diode patterning or alignment is required. Compared to imaging systems based on glass substrates, where refraction of light and the associated dispersion results in increased optical cross-talk between single pixels,[5] top absorbing OPD arrays with semitransparent top electrodes enable higher resolution.

In OPDs, light is detected upon absorption of incident photons with an energy equal to or larger than the optical bandgap of the photoactive material. Due to the higher absorption coefficients of organic materials (>10^5 cm⁻¹) when compared to silicon (50–100 cm⁻¹), thin active layers of only 100 nm are sufficient to absorb up to 60% of the incident light.[6] However, due to the lower relative permittivity (εᵣ ≈ 3–4) of organic relative to inorganic semiconductors, light absorption leads to photogenerated excitons with a comparatively high binding energy of ≈0.35–0.5 eV[7] rather than free electrons and holes. Thus, like many organic photovoltaic (OPV) devices, the active layer of OPDs is often based on a bulk heterojunction (BHJ) architecture[8] that comprises finely intermixed percolating networks of electron donor and acceptor phases, in which their large interface area facilitates exciton dissociation while the bicontinuous networks enable charge transport to the relevant electrodes. Figure 1a shows a typical OPD architecture...
comprising a BHJ layer sandwiched between electron and hole extraction layers (EEL and HEL, respectively).

1.2. Photosensor Arrays

In contrast to OPV devices that usually operate under positive bias at the maximum power point, a negative bias voltage is typically applied to OPDs (Figure 1b). For a sufficiently large reverse bias, all the photogenerated charges are efficiently extracted at the contacts due to the effective electric field, making the photocurrent independent of the applied voltage (Figure 1c).

Small OPD arrays can be read out using a so-called passive matrix.\cite{Ng2014} In this scheme, $N$ rows and $M$ columns define a $N \times M$ array of photodiodes. Current sneak paths however should be considered carefully. In the passive matrix scheme, the leakage currents of all pixels at the readout line add to the signal of one pixel that is addressed. For high-resolution and large-area imaging applications, so-called active-matrix addressing is typically used. Switching elements are typically introduced in the form of thin-film transistors (TFTs), although the use of blocking diodes has also been recently proposed.\cite{Janssen2019}

In the case of a TFT active matrix, the hole extracting contacts of all photodiodes are connected to a common electrode, which is connected to an external bias voltage source. The diodes are operated with a reverse bias voltage, of typically $-2$ to $-5$ V. The electron extracting contacts of all photodiodes of each column are connected to a common readout line via a switching TFT. Each readout line is connected to the input of its assigned readout amplifier. The gates of all TFTs in each row are connected to a common gate line, which is driven by a dedicated row driver output. The flat panel sensor is scanned one row at a time, in a similar way to active-matrix displays. During one frame time all the rows are sequentially selected by applying a voltage that changes the TFTs from the non-conducting to the conducting state. In this line selection time, the readout TFT transfers the charge from the photodiode capacitance to the data line and resets the voltage across the photodiode capacitance to its original value. During this time, the charge must be transferred from the photodiode to the integrating amplifier, and the output of the amplifier must be scanned. So, all pixels of an entire line are read out simultaneously.

The negative bias voltage applied to OPD arrays when integrated with a TFT matrix\cite{Janssen2019, Gelinck2013a, Gelinck2013b} ensures that the diode remains sufficiently charged, and effectively implies that the readout charge is linearly proportional to the amount of collected photocarriers. The reverse bias also increases the response speed of the diode, and often the collection efficiency, i.e., the extraction of photogenerated charge carriers at the contacts. On the other hand, dark current density will generally increase with increasing reverse bias. A reverse bias of $-2$ to $-5$ V is typically used as a good compromise.

The flexibility of OPD arrays makes them particularly suitable for applications where conformity is desirable. An important example is X-ray detectors, well-established technologies for digital radiography in the field of medical imaging. In these indirect-conversion detectors, a scintillator layer converts X-ray photons into UV or optical photons, which are then detected by an amorphous-silicon photodetector (a-Si PD) array mounted on a rigid glass substrate (Figure 1d).\cite{Ng2014} Replacing the a-Si PDs with solution-processed OPDs simplifies the manufacturing process considerably, thus reducing fabrication costs.

Flexible X-ray image sensors remain active under deformation (Figure 1g) and easily adapt to complex shapes, thereby potentially enabling more accurate imaging of the human body than inorganic X-ray detectors based on rigid glass substrates. Ng et al.\cite{Ng2014} first demonstrated a flexible photosensor array using solution-processed OPDs on a flexible a-Si matrix backplane. In 2014, Zhao et al.\cite{Zhao2014} proposed OPD-based X-ray imagers with amorphous indium gallium zinc oxide TFTs for next generation digital breast tomography systems. A proof-of-concept X-ray detector on thin plastic foil with $120 \times 160$ pixel format and 126 µm pixel size was demonstrated by Gelinck et al.\cite{Gelinck2013a} This

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combined a CsI:Tl scintillator with an OPD array and an oxide TFT backplane. The OPD layer was slot-die coated directly on the active matrix backplane on thin-plastic substrate.

OPDs have also been used in prototypes of biometric fingerprint and palmprint scanners,[14,15] where the OPD array detects visible light reflected by the finger or hand, respectively, as well as for gesture recognition. Fingerprint imaging is achieved by utilizing a difference in reflectance between the finger ridges and valleys. This technology offers a low-cost alternative to bulky inorganic detectors and can be easily integrated in mobile phones or door handles. Fingerprint scanners require higher spatial resolution compared to medical X-ray applications. Typically, a resolution of 200 pixels per inch (ppi) is high enough to unlock mobile phones. However, for high-quality fingerprint authentication a higher resolution is needed, allowing for more accurate sensing and thus a greater margin of safety. For gesture recognition (Figure 1e,h), shadowing by the hand is detected using lower resolution arrays albeit of larger size. A linear photoresponse of the OPD is important for both applications, as reliable operation under a range of different light conditions is required. This can be achieved by modulating the OPD photoresponse with an additional pulsed light source and discriminating the temporal component from the time-invariant background contribution.

Another promising application for OPDs is as the fundamental light sensing element in pulse oximeters, i.e., non-invasive medical sensors that can monitor heartbeat and measure blood oxygenation.[16,17] By using near-infrared (NIR) light, the light slightly penetrates the skin and is then partly absorbed or reflected. When used in reflective mode (Figure 1f,i), organic pulse oximeters provide a versatile alternative to rigid conventional designs that typically restrict the sensing location to finger tips or ear lobes. To estimate blood oxygenation levels, a good photoresponse at ca. 850 nm is desirable.[17] Typically,
light of at least two wavelengths is sequentially emitted. By detecting the reflected light of each color, the level of arterial hemoglobin oxygenation can be assessed.

Finally, the detection of NIR radiation is attractive for a wide variety of non-medical applications as well, such as optical communication,[18] night vision, and 3D object recognition.[1]

Having outlined the advantages of OPDs over inorganic photodetectors, we turn to the key performance metrics that quantify suitability for specific applications. Although they have been summarized in literature,[1,2] here we provide a comprehensive definition of the most important figures of merit to enable a clear understanding of reported OPD performances. Since reported metrics are often measured using different experimental methods and conditions, direct comparison of OPD performance metrics can often generate confusion. For the most relevant metrics, we highlight the variety of experimental methods, along with their challenges or limitations, and the dependence on the main experimental parameters (e.g., bias voltage, light intensity, etc.), to increase awareness and thus promote a more critical view when comparing reported values.

2. Performance Metrics of OPDs

2.1. EQE and Spectral Responsivity

The spectral responsivity $R$ in units of A W$^{-1}$ describes how much current is generated by the OPD per incoming photon of a given energy. It can be calculated via

$$R = \frac{J_{ph}}{P_{in}}$$

(1)

where $J_{ph}$ is the photocurrent density and $P_{in}$ is the incident light intensity.[1] The external quantum efficiency (EQE), i.e., the ratio between the number of incoming photons and the number of photogenerated free electrons, can be expressed as

$$\text{EQE} = \frac{R}{h} \frac{k\nu}{q}$$

(2)

where $h$ is the Planck constant, $\nu$ the frequency of the incident photon, and $q$ is the elementary charge. In general, high EQE (and hence $R$) is desirable to ensure efficient photon flux detection. While for broad-band OPDs the EQE should ideally be spectrally invariant and as high as possible across the entire operational wavelength range, narrow-band OPDs (defined by a quasi-Gaussian spectral distribution with a full-width-at-half-maximum (FWHM) $\leq 100$ nm) that require color discrimination should display high EQE only within the desired spectral window.[1] In addition to input filtering, the EQE spectrum can be manipulated by accurately tuning the optical bandgap, and hence absorption properties of the donor and acceptor phase. An alternative approach employs a thick ($\approx 2 \mu$m) BHJ layer to induce charge collection narrowing, in which only weakly absorbed photons produce electrons sufficiently close to the EEL to be collected.[21] Further increase of the EQE in specific wavelength ranges can be obtained by exploiting microcavity effects.[22,23] Since heterojunction OPDs behave as thin film optical cavities, interference phenomena derived from multiple reflections between two metallic semitransparent electrodes can enhance absorption within the photoactive layer, leading to broadening or narrowing of certain spectral features. These effects can be manipulated by carefully tuning the photoactive layer thickness and the properties of the optical spacer layers, and have recently been employed to enhance the EQE in the infrared region by increasing the charge-transfer (CT) state absorption.[24] EQE can also be enhanced by photomultiplication, i.e., the collection of multiple charge carriers per incident photon. In OPDs, this is generally achieved when trapped charge carriers cause the energy bands to bend, resulting in enhanced charge injection under illumination.[25]

Finally, EQE (and hence $R$) under reverse bias generally increases with greater applied voltage (i.e., electric field strength) due to enhanced charge extraction efficiency, but should eventually reach the saturation limit. In contrast, the dark current density ($J_d$) will keep increasing with increasing electric field (see Section 5.1), implying a trade-off between high EQE and low $J_d$ under reverse bias.

2.2. Speed of Response

Once electrons and holes are generated upon photon absorption, they drift towards the relevant electrodes due to the applied electric field. The speed of response, defined as the time required to collect photogenerated charge carriers at their respective contacts, ultimately determines the OPD dynamic range and cut-off frequency.[26] The response time can be measured by various techniques, including time-of-flight photocurrent transients[27] or small perturbation methods such as impedance spectroscopy.[28] Although they probe similar phenomena, these techniques may lead to different experimental results at low charge carrier densities ($<10^{16}$ cm$^{-3}$) due to surface recombination at the electrodes or inhomogeneous charge distribution across the active layer.[29] For polymer:fullerene BHJ OPDs, it is widely accepted that the transit time is limited by the mobility of the slowest carrier.[30] usually attributed to holes in the polymer phase.[31] Transit times for typical OPDs with thicknesses of 100–300 nm are of the order of $\mu$s, still less than the shortest interframe time (33 ms) for imaging applications operating at 30 frames per seconds (fps).[30]

2.3. Noise equivalent power and specific detectivity ($D^*$)

The minimum incident light power that can be detected by the OPD is referred to as noise equivalent power (NEP). The NEP is also defined as the signal optical power yielding a signal-to-noise ratio (SNR) equal to 1.[3] Making use of the definition for the spectral responsivity given by Equation (1), the NEP in units of W Hz$^{-1/2}$ can be expressed as

$$\text{NEP} = \frac{I_{\text{noise}}}{\sqrt{B}} \frac{1}{R}$$

(3)

where $I_{\text{noise}}$ is the noise current and $B$ the detection bandwidth.[31] The specific detectivity $D^*$ in units of cm Hz$^{1/2}$ W$^{-1}$
(i.e., Jones) is the reciprocal of the NEP normalized to the square root of the device area \(A\):

\[
D^* = \frac{R \sqrt{AB}}{i_{\text{noise}}}
\]  

(4)

Although accurate determination of \(i_{\text{noise}}\) is essential to estimate \(D^*\), experimental measurements of \(i_{\text{noise}}\) are challenging and not always performed. For sake of simplicity, the shot noise \(i_{\text{shot}}\) from the dark current \(i_d\) is often assumed to be the dominant contribution to \(i_{\text{noise}}\), leading to

\[
D^* \approx \frac{R \sqrt{AB}}{i_{\text{shot}}} = \frac{R \sqrt{A}}{2q_i} = \frac{R}{2q_i J_d}
\]  

(5)

Under this assumption, \(J_d\) is taken to be a constant value that is subtracted from the photoinduced signal. The use of Equation (5) implies that the dark current is dominated by shot noise that limits \(D^*\), while 1/f noise and thermal noise\(^{[12]}\) are ignored. It has been pointed out that this assumption can overestimate the specific detectivity.\(^{[1,13]}\) The general applicability of Equation (5) can therefore be questioned. However, Equation (5) is widely used and thus enables a fair comparison between reported OPD performances. Nevertheless, confusion may still arise when comparing reported \(D^*\) values calculated with Equation (3), for two reasons: a) \(J_d\) depends on the applied electric field (see Section 5.1) and b) \(R\) depends on the wavelength and the applied light intensity according to Equation (1).

Because it is not always possible to compare \(D^*\) values measured under the same experimental conditions, here we will consider the maximum \(D^*\) value reported in each work.

### 2.4. Dark Current Density (\(J_d\))

Dark current is defined as any current generated under an applied reverse bias voltage in the absence of light. The detrimental influence of a high dark current density on OPD performance can be quantified by its negative impact on several key metrics. First, high \(J_d\) results in a lower SNR and decreases sensitivity to low light intensities. Furthermore, it constrains the linear dynamic range (LDR), i.e., the operational light intensity range of the image sensor.\(^{[1]}\) by increasing the minimum detectable photocurrent. Finally, the specific detectivity \(D^*\) decreases for increasing dark current densities according to Equation (5).

Unlike \(R\), \(J_d\) can span multiple orders of magnitude depending on material properties and device architecture.\(^{[34]}\) meaning that reducing \(J_d\) is of utmost importance for improving SNR, LDR, and \(D^*\) in OPDs. Exploration of the origins of dark current, so essential to minimize when developing high-performing OPDs, and specific strategies to reduce \(J_d\) form the bulk of this progress report.

Over recent years, great effort has been devoted to lowering dark current density in OPDs. State-of-the-art OPDs exhibit values in the range of \(10^{-6}–10^{-7}\) mA cm\(^{-2}\).\(^{[34,35]}\) However, direct comparison of the reported values is sometimes hampered by the fact that \(J_d\) can strongly depend on the applied bias, and no specific measurement protocol is agreed. To avoid confusion, in this work we report \(J_d\) values at a given effective electric field \(F = 6 \times 10^6\) V m\(^{-1}\), unless otherwise stated.

Corresponds to an applied voltage of \(V = -2\) V over an active layer with a thickness \(L = 300\) nm (assuming \(F = V/L\)), and thus falls into a reasonable range for OPD applications.

### 3. Dark Current in OPDs: Origin and Solutions

#### 3.1. Dark Current Mechanisms

Effective suppression of OPD dark current requires a deep understanding of its fundamental origin. When the OPD is operated under an applied reverse bias voltage, intrinsic \(J_d\) (i.e., in the absence of pinholes or other extrinsic leakage phenomena) is mainly attributed to the charge carrier injection rate from the contacts into the semiconductor, or the rate of thermal generation of charge carriers within the bulk of the active layer, followed by drift towards their respective electrodes under the applied electric field. In Figure 2a, the energy levels of donor and acceptor are represented by the orange full lines and the black dotted lines, respectively. The density of state (DOS) distributions of the donor highest occupied molecular orbital (HOMO) and the acceptor lowest unoccupied molecular orbital (LUMO) are assumed to be Gaussian.

In the first scenario, electrons are injected from the high work function electrode into energy states in the tail of the Gaussian DOS of the acceptor LUMO, whereas holes are injected from the low work function electrode into tail states of the donor HOMO. The charge injection rate is expected to be strongly dependent on the injection barrier \(\Phi_b\) defined as the energy difference between the acceptor LUMO and Fermi level of the high work function electrode in the case of electrons. A higher \(\Phi_b\) results in a lower dark current for a given applied bias. Assuming perfect Ohmic contacts and alignment between the low (high) work function electrode Fermi level and the acceptor LUMO (donor HOMO), the injection-limited \(J_d\) is expected to be proportional to the effective bandgap \(E_g\) defined as the energy difference between the acceptor LUMO and the donor HOMO, as donor and acceptor materials are in contact with both electrodes.

In the second scenario, charge carriers are thermally generated and collected at the contacts. It is often stated that bulk thermal generation within the polymeric semiconductor can be neglected due to the large bandgap \(E_g\) of organic materials (i.e., \(E_g > 1\) eV),\(^{[2]}\) resulting in negligibly small concentration of thermally generated carriers \(n_i\), as \(n_i \propto \exp(–E_g/k_B T)\) with \(k_B\) the Boltzmann constant and \(T\) the absolute temperature. However, thermally generated carriers close to the donor–acceptor (D-A) interface might be easily dissociated within the effective bandgap \(E_{ip}\) leading to \(n_i \propto \exp(–E_g/k_B T)\). Although thermal generation at the D-A interface via ground-state CT interaction can typically be neglected in the case of UV–visible OPDs\(^{[2,36]}\) due to the deep HOMO energy of the donor material resulting in large \(E_g\), its contribution may be relevant for NIR-sensitive OPDs.

#### 3.2. Dark Current Reduction Strategies

Figure 2b summarizes the main experimental approaches to reduce the dark current density in OPDs. First, \(J_d\) can be reduced by astute selection of the energy levels within the
photoactive layer. Choosing a donor material with a deep HOMO energy would minimize hole injection from the low work function electrode (see Section 4.1). Equally, an acceptor material with a shallow LUMO would be expected to minimize electron injection, but this route is less explored as most OPD are developed using fullerene derivatives with similar LUMO values. Furthermore, donor and acceptor materials with smaller energetic disorder, i.e., a narrower Gaussian DOS, are expected to increase the effective injection barrier, thereby reducing $J_d$.

In addition, the photoactive layer stratification can be adjusted specifically with the aim of reducing the dark current density (see Section 4.2). Planar heterojunction (PHJ) OPDs made via sequential deposition of individual donor and acceptor layers are widely investigated. Alternatively, sequentially solution-processed (SSP) OPDs can lead to a partial donor-acceptor interpenetration, resulting in a well-mixed interdiffusion phase in the middle of the bulk. Both approaches can lead to the formation of pure phases at the electrodes, which can effectively block unfavorable charge carrier injection under reverse bias. The injection barrier $\Phi_b$, and thus the charge injection rate is expected to scale with the energy difference between the donor LUMO and Fermi level of the high work function electrode in the case of electrons.

Another common strategy to reduce $J_d$ consists of improving charge selectivity at the contacts (see Section 4.3). This is obtained by raising energetic barriers to charge injection under reverse bias, while maintaining an energy cascade between the active layer and the corresponding electrodes to facilitate extraction of photogenerated carriers. To achieve this, electron-blocking layers (EBLs) and hole-blocking layers (HBLs) are often employed. These comprise an additional interlayer between the electrode and the photoactive layer, such that $E_{\text{LUMO,EBL}} > E_{\text{LUMO,acceptor}}$ for electrons and $E_{\text{HOMO,HBL}} < E_{\text{HOMO,donor}}$ for holes, resulting in an increased energetic barrier for charge injection. Ideally, $E_{\text{HOMO,EBL}} = E_{\text{HOMO,donor}}$ and $E_{\text{LUMO,HBL}} = E_{\text{LUMO,acceptor}}$, so that photogenerated carrier collection is not impeded.

Table 1 contains a summary of recently published OPD performances. In each work, $J-V$ characteristics in dark conditions and/or $D^*$ spectra are presented, the main $J_d$ mechanisms are mentioned and/or a description of the experimental approach for $J_d$ reduction is provided, along with the experimental results. $J_d$ values are chosen at electric field $F = 6 \times 10^6$ V m$^{-1}$. $D^*(\lambda_{max})$ is the specific detectivity at the wavelength corresponding to the maximum of the first allowed optical absorption band of the semiconductor. Reported $D^*$ values were calculated using Equation (5), unless otherwise stated. Interestingly, fullerene derivatives are used as electron acceptors in all work, except for...
Table 1. Overview of the most common J<sub>d</sub> mechanisms and experimental approaches for J<sub>d</sub> reduction. Dark current density (J<sub>d</sub>) and specific detectivity (D<sup>∞</sup>) at the wavelength corresponding to the maximum of the first allowed optical absorption band of the semiconductor (λ<sub>max</sub>).

| Year | Ref. | J<sub>d</sub> mechanism | Experimental approach for J<sub>d</sub> reduction | Results |
|------|------|-------------------------|-----------------------------------------------|---------|
|      |      |                         | Active layer materials<sup>a</sup> | Active layer morphology | Blocking layers | J<sub>d</sub> [mA cm<sup>−2</sup>] | D<sup>∞</sup>(λ<sub>max</sub>) [I0] | λ<sub>max</sub> [nm] |
| 2007 | [44] | ✓ | − | P3HT:PC<sub>61</sub>BM | BHJ | PFB | − | 2.0 × 10<sup>−4</sup> | − | 500 |
| 2008 | [45] | ✓ | − | P3HT:PC<sub>61</sub>BM | BHJ | − | − | 5.0 × 10<sup>−6</sup> | 7.0 × 10<sup>12</sup> | 468 |
| 2011 | [46] | ✓ | − | Squarine:PC<sub>61</sub>BM | BHJ | MEH-PPV | − | 3.5 × 10<sup>−5</sup> | 3.4 × 10<sup>12</sup> | 700 |
| 2012 | [47] | ✓ | − | P3HT:PC<sub>61</sub>BM | BHJ | − | TAZ | 4.5 × 10<sup>−5</sup> | 3.0 × 10<sup>12</sup> | 500 |
| 2013 | [48] | ✓ | − | C<sub>60</sub> | Single layer | C-TPD | − | 7.0 × 10<sup>−6</sup> | 3.6 × 10<sup>11</sup> | 370 |
| 2013 | [49] | ✓ | − | P3HT:PC<sub>61</sub>BM | BHJ | P3HT | − | 3.0 × 10<sup>−6</sup> | − | 600 |
| 2013 | [50] | ✓ | − | PBDTT-T:PC<sub>61</sub>BM | BHJ | − | PEIE | 2.0 × 10<sup>−5</sup> | 8.5 × 10<sup>12</sup> | 680 |
| 2015 | [51] | ✓ | − | P3HT:Indigo | BHJ | − | − | 2.9 × 10<sup>−5</sup> | 1.0 × 10<sup>12</sup> | 630 |
| 2015 | [52] | ✓ | − | PDPDTT:PC<sub>71</sub>BM | BHJ | poly-TPD | − | 1.0 × 10<sup>−5</sup> | 1.0 × 10<sup>13</sup> | 870 |
| 2015 | [53] | ✓ | − | PCDTB:PC<sub>71</sub>BM | BHJ | − | PEIE | 1.4 × 10<sup>−7</sup> | 3.4 × 10<sup>13</sup> | 532 |
| 2016 | [54] | ✓ | − | PCDTB:PC<sub>71</sub>BM | BHJ | − | − | 7.0 × 10<sup>−6</sup> | 3.0 × 10<sup>13</sup> | 575 |
| 2016 | [55] | ✓ | − | P3HT:PC<sub>61</sub>BM | BHJ | − | YbF<sub>3</sub> | − | 1.7 × 10<sup>12</sup> | 520 |
| 2016 | [56] | ✓ | − | P3HT:PC<sub>61</sub>BM | BHJ | − | C<sub>60</sub> | 1.3 × 10<sup>−5</sup> | 2.4 × 10<sup>21</sup> | 580 |
| 2016 | [57] | ✓ | − | ZnO:F8T2 | PHJ | − | − | 6.0 × 10<sup>−7</sup> | 1.2 × 10<sup>21</sup> | 450 |
| 2017 | [42] | ✓ | − | PBDDTTF:PC<sub>61</sub>BM | BHJ | − | − | 1.5 × 10<sup>−5</sup> | 3.3 × 10<sup>21</sup> | 690 |
| 2017 | [58] | ✓ | ✓ | TAPC-C<sub>60</sub> | BHJ | − | − | 1.0 × 10<sup>−5</sup> | − | − |
| 2017 | [59] | ✓ | − | PFB:T2OBT:PC<sub>61</sub>BM | BHJ | − | − | 6.5 × 10<sup>−7</sup> | − | − |
| 2017 | [60] | ✓ | − | PCDTB:TPD:PC<sub>61</sub>BM | BHJ | − | TIPS-P | 1.0 × 10<sup>−7</sup> | 1.4 × 10<sup>13</sup> | 610 |
| 2018 | [61] | ✓ | − | C<sub>60</sub> | Single layer | TAPC | − | 2.0 × 10<sup>−4</sup> | 1.2 × 10<sup>12</sup> | 530 |
| 2019 | [62] | ✓ | − | PV:PC<sub>61</sub>BM | BHJ | − | − | 3.5 × 10<sup>−7</sup> | 6.4 × 10<sup>12</sup> | 360 |
| 2019 | [53] | ✓ | − | PV:PC<sub>61</sub>BM | BHJ | − | − | 3.4 × 10<sup>−5</sup> | 2.2 × 10<sup>13</sup> | 660 |
| 2019 | [43] | ✓ | − | PV:O-IDTBR:PC<sub>61</sub>BM | BHJ | − | − | 2.0 × 10<sup>−5</sup> | 4.6 × 10<sup>12</sup> | 755 |

<sup>a</sup>Abbreviations are defined in the text; PFB = poly[9,9’-dioctylfluorene-co-bis-N,N’-(4, butylyphenyl)-bis-N,N’-phenyl-1,4-phenylene-diamine]; MEH-PPV = poly[2-methoxy-5-(2’-ethyl-hexyl)-oxy]-1,4-phenylene-vinylene]; TAZ = 3-phenyl-4(1’-naphthyl)-5-phenyl-1,2,4-triazole; C-TPD = cross-linked 4,4’-bis[(p-trichlorosilyl)propyloxy]phenylamine]-bisphenyl; PVK = poly(9-vinylcarbazole) and F8T2 = poly[9,9-dioclyfluorene-al bithiophene]; poly[4,4,9,9-tetrakis(4-hexylphenyl)-4,9-dihydro-s-indaceno][2,3-b:5,6-’b’][dithiophene-2,7-diyl-alt-S(2-ethylhexyl)-4H-thieno[3,4-d]pyrrole-4,6(5H)-dione]-1,3-diy]; TIPS-P = 6,13-bis(trisopropylsilylhexyl)pentacene; O-IDTBR = rhodanine-benzothiadiazole-coupled indacenodithiophene; <sup>b</sup>λ<sub>max</sub> at reverse bias electric field F = 6 × 10<sup>6</sup> V m<sup>−1</sup>; <sup>c</sup>D<sup>∞</sup> values based on experimental measurements of the noise current I<sub>n</sub>noise.

some recent exceptions.\textsuperscript{[18,43]} This implies that the difference in optical and electrical characteristics reported here are largely due to the different polymers used as donor materials. Furthermore, dark current densities below 10<sup>−2</sup> mA cm<sup>−2</sup> are rarely achieved within a relevant electric field range for applications. Nevertheless, very low J<sub>d</sub> in the order of 10<sup>−7</sup> mA cm<sup>−2</sup> can be obtained using various methods, which suggests that there is currently no unique strategy to reduce the dark current density in OPDs. This indicates that J<sub>d</sub> depends in a complex way on several material and design parameters, and that its underlying mechanisms are still largely unknown.

Figure 3 shows a graphical overview of OPDs reported in the recent years. In Figure 3a, the dark current density at reverse bias electric field F = 6 × 10<sup>6</sup> V m<sup>−1</sup> is plotted as function of the wavelength corresponding to the maximum of the first allowed optical absorption band of the polymeric semiconductor. Here, OPDs are classified in four categories according to the wavelength of operation, namely UV, blue, green, and NIR, with the clear majority lying in the region between 500 and 700 nm. Although the lowest dark currents are generally reached for wide bandgap materials, there is no evident correlation between J<sub>d</sub> and λ<sub>max</sub>. Adding blocking layers...
(full symbols) clearly helps reducing $J_d$ while maintaining the same OPD spectral response. In the case of NIR OPDs, BLs seem to be essential to achieve low dark current densities in the order of $10^{-6}$ mA cm$^{-2}$. However, green-sensitive OPDs with blocking layers still do not outperform the best BHJ OPDs without BLs at the same operational wavelengths. In the attempt to plot $J_d$ in a meaningful way for OPDs with and without BLs, Figure 3b shows $J_d$ as function of the effective injection barrier $\Phi_e$. For OPDs with Ohmic contacts, $\Phi_e = E_0 = |LUMO_A - HOMO_D|$ assuming pinning of the BHJ energy levels to the work function of the injecting contacts under reverse bias, as explained in Section 3.1. For OPDs with electron or hole BLs, $\Phi_e = |LUMO_{BLl} - W_e|$ or $\Phi_e = |W_e - HOMO_{Dl}|$, respectively, with $W_e$ the work function of the electron or hole injecting contact under reverse bias. Finally, for OPDs with PHJ active layers, $\Phi_e = |LUMO_b - W_b|$ or $\Phi_e = |W_b - HOMO_b|$ for the case of electron or hole injection under reverse bias, respectively. The dashed line indicates the intrinsic limit of the injection-limited $J_d$ as calculated from Equation (7) for a reference OPD, as explained in Section 5.1. Here, the absolute values of LUMO$_b$ and HOMO$_D$ as well as the energy levels of the BLs and the work functions of the injecting contacts under reverse bias, were taken from each work; uncertainty regarding their absolute values that arises from different measuring techniques must be considered, as will be discussed in Section 6. Nevertheless, it becomes clear that low dark current densities can rarely be achieved when the effective injection barrier is reduced, due to the enhanced probability of charge carrier injection and/or bulk thermal generation. This also sets an intrinsic limit to the maximum $D^*\text{ as function of }\Phi_e$ (Figure 3c).

Here, the intrinsic limit of $D^*$ was assessed with Equation (5) using the calculated $J_d$ and assuming a reasonable value for the responsivity $R$, i.e., $R = 0.2$. While the use of BLs in many works decreases $J_d$ (increases $D^*$), most results are scattered and still lie far from the intrinsic limits for a given value of $\Phi_e$.

This suggest that BLs can effectively lower the dark current but do not reduce $J_d$ to the expected intrinsic limit. In addition, this indicates that extrinsic factors may affect the experimental values of $J_d$ and $D^*$ in many cases. Possible explanations for this are the object of further discussion in Section 6. Notably, the graphical overview in Figure 3b enables to identify the best reported OPDs (data points approaching the intrinsic limits), some of which are object of a more detailed study in Section 4.

### 4. Effect of OPD Parameters on Dark Current

In this section, we provide a more detailed analysis to the different experimental approaches employed to reduce the OPD dark current. For each approach, the best examples of OPDs in literature are cited and their $J_d$ values are reported, with the aim to select the most successful strategies for $J_d$ reduction.

#### 4.1. Active Layer Energetics

Having a large offset between the Fermi level of the electron extracting contact and the donor polymer HOMO energy has been reported to be a prerequisite to reduce hole injection and thus minimize $J_d$. Indeed, the lowest dark current densities reported in literature for OPDs are achieved using polymers with a deep HOMO energy. It is no coincidence that the record values of $J_d$ are achieved for poly[N9′-heptadecanyl-2,7-carbazole-alt-5,5-(4′,7′-di-2-thienyl-2′,1′,3′-benzothiadiazole)] (PCDITBT),\textsuperscript{[34,52]} which has one of the deepest HOMO energies amongst photovoltaic polymers ($≈ 5.5$ eV).\textsuperscript{[34]} This implies that careful selection of the photoactive layer materials is a key element to reducing $J_d$ without additional interlayers, resulting in a faster and cheaper fabrication process. However, a trade-off between $J_d$ and the maximum absorption wavelength of the photoactive layer is

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**Figure 3.** Overview of recently published OPD parameters (for details see Figures S1–S3, Supporting Information). Empty symbols refer to BHJ OPDs without BLs, full symbols refer to BHJ OPDs with BLs or PHJ OPDs. a) Dark current density at a reverse bias electric field $F = 6 \times 10^6$ V m$^{-1}$ as function of wavelength corresponding to the maximum of the first allowed optical absorption band of the semiconductor. Symbols linked by full lines belong to the same publication. b) Dark current density at reverse bias electric field $F = 6 \times 10^6$ V m$^{-1}$ and c) maximum specific detectivity $D^*$ as function of the effective bandgap, i.e., the difference between the acceptor LUMO and the donor HOMO. The dashed line in b) indicates the intrinsic limit of the injection-limited $J_d$ as calculated from Equation (7) for a reference OPD, as explained in Section 5.1. The dashed line in c) was determined using Equation (5) with the calculated $J_d$ and assuming $R = 0.2$. © 2019 The Authors. Published by WILEY-VCH Verlag GmbH & Co. KGaA, Weinheim.
reached, given the need for a suitably shallow donor polymer LUMO that facilitates exciton separation under illumination and ensures a high open circuit voltage \( (V_{oc}) \). If the LUMO of the donor is too deep, this would cause inefficient exciton separation and thus low responsivity.\(^{[62]}\) Optimal energetic offset in the order of \( \approx 0.2-0.3 \) eV is necessary to achieve CT from donor to acceptor.\(^{[63]}\) It must be emphasized that this rule is empirical and that exceptions may occur where efficient free charge carriers are photogenerated even at lower energy offsets. Nevertheless, this requirement sets an intrinsic limitation to the deepest HOMO energy for NIR polymers, where a narrow optical bandgap must be maintained to enable the detection of the NIR radiation. As an example, assuming light absorption mainly in the donor phase and \( 0.25 \) eV LUMO offset relative to the acceptor, an effective bandgap \( E_g \) of \( \approx 1.1 \) eV is necessary to detect light up to \( 850 \) nm for applications in pulse oximeters (see Section 1.2). The relatively low \( E_g \) of NIR OPDs enhances charge injection from the metal electrodes—assuming Ohmic contacts between the electron (hole) extracting layer and the LUMO of the acceptor (HOMO of the donor)—or thermal generation within the photoactive layer. This makes it challenging to achieve NIR-sensitive OPDs with low \( J_D \) and high \( D^* \) by only optimizing the photoactive layer energetics.

4.2. Optimization of the Photoactive Layer: Stratification and Thickness

The PHJ architecture provides a valid approach to reduce the injection-limited as well as the thermally generated \( J_D \). Recently, Yoon et al.\(^{[38]}\) achieved \( J_D = 6 \times 10^{-7} \) mA cm\(^{-2} \) at \( F = 6 \times 10^{8} \) V m\(^{-1} \) for a color-selective inverted-polarity PHJ OPD using a blue-sensitive polymer donor in combination with non-absorbing ZnO acceptor. Shekhar et al.\(^{[39]}\) reached \( J_D \) in the order of \( 10^{-6} \) mA cm\(^{-2} \) in combination with EQE well above 10% in the spectral range between 400 and 800 nm. This promising result was obtained using very thin donor and acceptor layers [20 nm of boron subphthalocyanine chloride (SubPc) and 40 nm of C\(_{60}\), respectively] to ensure fast migration of the photogenerated exciton to the D-A interface and efficient charge separation.

Agostinelli et al.\(^{[40]}\) investigated the effect of D-A vertical segregation on the dark current density of a SSP OPD based on poly(3-hexylthiophene) (P3HT) and phenyl-C\(_{61}\)-butyric acid methyl ester (PC\(_{61}\)BM). Changes in the vertical composition profile induced by annealing led to better contact selectivity that reduced \( J_D \) under reverse bias. However, the OPD with the lowest \( J_D = 10^{-8} \) mA cm\(^{-2} \) also showed a relatively low EQE of 0.024 at 500 nm, mainly attributed to the high series resistance of the thick PC\(_{61}\)BM layer and the inefficient separation of photogenerated excitons due to insufficient interdiffusion at the P3HT-PC\(_{61}\)BM interface. In the work of Wang et al.\(^{[41]}\) optimal interdiffusion of at the D-A interface led to high EQE of 0.69, but \( J_D \) did not decrease below \( 10^{-4} \) mA cm\(^{-2} \) at \( F = 6 \times 10^{4} \) V m\(^{-1} \). Thus, the trade-off between EQE and dark current density depending on the degree of D-A vertical segregation is the main limitation of SSP OPDs.

Finally, \( J_D \) is found to decrease with increasing photoactive layer thickness \( L \) for two main reasons, namely 1) the decrease of extrinsic defects\(^{[64]}\) (e.g., film imperfections, pinholes, and other leakage paths) resulting in higher manufacturing yields and lower probability of soft breakdown above a certain electric field, and 2) the reduction of image charge effect,\(^{[65]}\) resulting in a higher effective barrier for charge injection. Biele et al.\(^{[5]}\) optimized the spray coating process anticipated by Tedde et al.\(^{[66]}\) by using high boiling point 1,8-diiodooctane (DIO) as processing additive, which enabled the single droplets to coalesce, resulting in a thick active layer (800 nm) with reduced root-mean-square roughness. This led to an impressively low dark current density of \( J_D = 3.4 \times 10^{-8} \) mA cm\(^{-2} \) and EQE = 0.82 at 650 nm under \(-5 \) V (Figure 4). Pierre et al.\(^{[52]}\) reported high manufacturing yields for OPDs with the maximum thickness limited by the solubility of donor and acceptor phases in solution (\( \approx 570 \) nm). On the other hand, too thick active layers (\( \approx 1000 \) nm) cause a decrease in responsivity as charge carriers are not efficiently extracted and thus recombine within the photoactive layer.\(^{[5]}\) This implies a trade-off between \( J_D \) and photocurrent as \( L \) is varied.\(^{[67]}\)

4.3. Improving the Charge Selectivity at the Contacts

4.3.1. Electron-Blocking Layers

In the recent years, many groups have employed EBLs to effectively reduce the dark current in OPDs. Zhou et al.\(^{[15]}\)

![Figure 4](https://www.afm-journal.de)
demonstrated a 25 nm thick poly[N,N′-bis(4-butylphenyl)-N,N′-bisphenylbenzidine] (poly-TPD) EBL to suppress the reverse-bias dark current in an NIR OPD based on poly[2,5-bis(2-hexyldecyl)-2,3,5,6-tetrahydro-3,6-dioxopyrrolo[3,4-c]-pyrrole-1,4-diyl]-alt-[2,2′:5′,2″-terthiophene]-5,5″-diyl] (PDPP3T) mixed with phenyl-C71-butyric acid methyl ester (PC71BM). Compared to the standard OPD with a PEDOT:PSS interlayer, introducing the poly-TPD EBL reduced the dark current density by more than two orders of magnitude due to the superior electron-blocking ability, leading to $J_d \approx 10^{-6}$ mA cm$^{-2}$ under a reverse electric field of $6 \times 10^6$ V m$^{-1}$ and thus a specific detectivity $D^*$ greater than $1.0 \times 10^{13}$ Jones at wavelengths from 350 to 870 nm. Although no loss in the EQE was reported under applied reverse bias, the small energy barrier of $\approx 0.2$ eV at the interface between poly-TPD and the photoactive layer (Figure 5a) lowered the extraction efficiency near the $V_{oc}$, resulting in s-shaped $J$-$V$ characteristics under illumination (Figure 5b). Nevertheless, poly-TPD was demonstrated to be a promising EBL material being a wide-bandgap hole transporting polymer with shallow LUMO level ($-2.3$ eV) and HOMO energy close to that of the donor polymer (PDPP3T).

4.3.2. Hole-Blocking Layers

In addition to EBLs, the use of polyethyleneimine-ethoxylated (PEIE) as an effective HBL has been widely demonstrated. When a PEIE layer is solution-processed on the indium tin oxide (ITO) bottom electrode, simple aliphatic amine groups in this polymer are physisorbed onto the ITO and the CT nature of their interaction with the surface is found to reduce the ITO work function. Therefore, the minimization of $J_d$ in OPDs with PEIE interlayers is often attributed to its hole-blocking properties under reverse bias. Saracco et al. first reported $J_d \approx 2 \times 10^{-6}$ mA cm$^{-2}$ at $F = 6 \times 10^6$ V m$^{-1}$ in OPDs consisting of a blend of poly[(4,8-bis-(2-ethylhexyloxy)-benzo(1,2-b:4,5-b′)dithiophene)-2,6-diyl-alt-(4-(2-ethylhexanoyl)-thieno[3,4-b] thiophene-)2,6-diyl)] (PBDTTT-C) with PC71BM, using PEIE interlayer. Pierre et al. achieved an impressively low dark current density of $1.4 \times 10^{-7}$ mA cm$^{-2}$ and $D^* = 3.45 \times 10^{13}$ Jones using PCDTBT as donor polymer (Figure 6). Similarly, Kielar et al. reached $J_d = 3 \times 10^{-7}$ mA cm$^{-2}$ and $D^* = 3.21 \times 10^{13}$ Jones at $\approx 530$ nm, which is amongst the highest reported specific detectivities at this wavelength.
5. Exploring the Intrinsic Limits of the Dark Current in OPDs

In this section, we provide a quantitative analysis of two main dark current mechanisms, that is charge injection and bulk thermal generation. In the former case, we use an analytical model to show the dependence of the injection-limited \( J_{\text{d}} \) on the main material and device parameters, namely active layer thickness \( L \), effective bandgap \( E_{\text{g}} \), energetic disorder \( \sigma \), and charge carrier mobility \( \mu \), all within a reasonable range for OPD applications. For the latter, we calculate the contribution from thermally generated carriers by treating the device as a blackbody, demonstrating that charge injection can be safely considered as the dominant contribution to \( J_{\text{d}} \). Finally, based on this theoretical insight, we suggest guidelines to improve OPD performance.

5.1. Injection-Limited Dark Current

A major contributor to \( J_{\text{d}} \) charge carrier injection under reverse bias proceeds via a thermally activated jump that promotes the injection of an electron (hole) from the Fermi level of the hole (electron) extracting electrode to a tail state in the acceptor LUMO (donor HOMO). We invoke the formalism proposed by Arkhipov et al.,\(^\text{[6]}\) which captures the fundamental two-step process that contributes to the injection-limited dark current, namely a first carrier jump into the semiconductor followed by a certain probability to escape to the opposite electrode. This injection model considers 1) the image charge effect at the electrode, 2) the energetic site disorder inherent to organic semiconductors, and 3) the hopping charge transport mechanism. Charge injection occurs from the electrode Fermi level into tail states of the DOS of the active layer materials, as previously described by Gartstein and Conwell.\(^\text{[69]}\) The charge carrier can escape recombination with its image charge under the conditions that a neighboring hopping site at equal or lower energy is encountered.\(^\text{[6]}\) As the carrier moves away from the electrode interface, it drifts in the electric field towards the opposite electrodes via a diffusive random walk in the disordered energetic landscape. It should be noted that the injection process will not be spatially homogeneous but instead filamentary,\(^\text{[70]}\) since the first injection event is dominated by the presence of spatially fixed tail states in the DOS distribution close to the interfaces.

Applying the assumptions of the Arkhipov et al. formalism, we define the injection-limited \( J_{\text{d}} \) as the product between the charge carrier density at the metal–semiconductor interface \( n_{\text{int}} \) and the carrier mobility \( \mu_{\text{d}} \) in the bulk of the semiconductor:

\[
J_{\text{d}} \propto q n_{\text{int}} \mu_{\text{d}} F
\]

Assuming a Gaussian DOS, the injection barrier \( \Phi_{\text{inj}} \) is equal to \( \Phi_{\text{bd}} \), minus a term proportional to the thermal equilibrium energy \( \frac{\sigma_{\text{i}}}{2k_{\text{B}}T} \) with \( \sigma_{\text{i}} \) the energetic disorder at the metal–semiconductor interface. The effective injection barrier \( \Phi_{\text{inj}} \) is further reduced by the image potential barrier lowering \( \Phi_{\text{i}} = \frac{\sigma_{\text{i}}}{2k_{\text{B}}T} \), being \( \varepsilon = \varepsilon_{\text{bd}} \) the permittivity. Under these conditions, \( n_{\text{int}} \approx N_{\text{i}} \exp \left( \frac{\Phi_{\text{inj}}}{k_{\text{B}}T} \right) \) with \( \Phi_{\text{inj}} = \Phi_{\text{bd}} - \frac{\sigma_{\text{i}}}{2k_{\text{B}}T} \) (Figure 7a) and with \( N_{\text{i}} \) the volume density of molecular sites between which the hopping takes place. The mobility in the zero-field limit and at carrier density in the independent-particle (Boltzmann) regime \( \mu_{\text{d}} \) is equal to the product of the mobility in the infinite temperature limit, \( \mu_{\text{d}}^{*} \), and a temperature-dependent factor \( \exp \left( -\frac{\sigma_{\text{i}}}{k_{\text{B}}T} \right) \) with \( \sigma_{\text{i}} \) the energetic disorder in the bulk of the semiconductor and \( C_{\text{i}} = 0.4 \) within the extended Gaussian density model.\(^\text{[71]}\) Thus, the current density is given by:

\[
J = A q N_{\text{i}} \exp \left( \frac{\Phi_{\text{inj}}}{k_{\text{B}}T} \right) \mu_{\text{d}}^{*} \exp \left( -C_{\text{i}} \frac{\sigma_{\text{i}}}{k_{\text{B}}T} \right) \frac{V}{L} \tag{7}
\]

In this expression, the dimensionless prefactor \( A \) is considered equal to unity, as expected for OPDs with large injection barriers and uniform carrier density across the device thickness.\(^\text{[7]}\)

The injection of the main OPD parameters on the injection-limited dark current under reverse bias voltage of \( V = -3 \) V as estimated from Equation (7) is shown in Figure 7b. Data are normalized to a reference case with \( L = 100 \) nm, \( \mu_{\text{d}} = 10^{-7} \text{ m}^{2} \text{ V}^{-1} \text{ s}^{-1} \), \( \Phi_{\text{bd}} = 1 \) eV, and \( \sigma_{\text{i}} = 0.1 \) eV. Each parameter is varied within a reasonable range for OPDs applications while keeping the other parameters constant, thereby assuming their mutual independence. The \( V \)-dependence of \( J_{\text{d}} \) upon variations in \( L \), \( \mu_{\text{d}} \), \( \Phi_{\text{bd}} \), and \( \sigma_{\text{i}} \) is shown in Figure 7c–e, respectively. The dark current density decreases with increasing active layer thickness, as observed in Section 4.2. However, the thickness dependence is weaker for increasing \( L \), since \( J_{\text{d}} \) depends linearly on the electric field but exponentially on \( \Phi_{\text{bd}} \). Overall, the intrinsic thickness dependence of \( J_{\text{d}} \) is much less significant when compared to the other material parameters. However, very thick active layers (\( \sim 800 \) nm) are beneficial in reducing \( J_{\text{d}} \) due to extrinsic effects, such as the reduction of pinholes and film imperfections,\(^\text{[7]}\) as will be discussed in Section 6. For a single carrier, the dark current density depends linearly on the charge carrier mobility according to Equation (7). However, effective \( \mu_{\text{d}} \) below \( 10^{-9} \text{ m}^{2} \text{ V}^{-1} \text{ s}^{-1} \) for a charge carrier in a BHJ active layer is unrealistic, implying that \( \mu_{\text{d}} \) has limited importance in defining \( J_{\text{d}} \). The major role in defining the injection-limited dark current density is played by the effective injection barrier, leading to a decrease by 5 orders of magnitude in \( J_{\text{d}} \) upon variations in \( \Phi_{\text{bd}} \) from 1.0 to only 1.3 eV. Remarkably, \( J_{\text{d}} \) increases by a factor 10 when the interface disorder \( \sigma_{\text{i}} \) is increased from 0.1 eV to only 0.12 eV, which underlines the importance of choosing active layer materials with low energetic disorder (such as PCBM).

5.2. Thermally Generated Dark Current

To determine whether bulk thermal generation makes a significant contribution to the dark current, we adapt an approach commonly applied to solar cells under illumination.\(^\text{[7,72]}\) For solar cells, the short-circuit current \( J_{\text{sc}} \) is determined by the overlap integral between the incident photon flux, \( \phi(E) \) and the external quantum efficiency spectrum, EQE(\( E \)) via

\[
J_{\text{sc}} = q \int \text{EQE}(E) \phi(E) dE \tag{8}
\]
The thermally generated dark current can be calculated by replacing the typical AM1.5G solar spectrum with the blackbody spectrum. To determine the thermally generated photon flux, we begin with the well-known photon energy distribution inside a blackbody in the interval from $E$ to $E + dE$, given by

$$e(E) dE = \frac{8\pi}{\hbar c^2} \frac{E^3}{\exp(E/k_B T) - 1} dE$$

as taken from Vandewal et al.,\cite{Vandewal2012} with $c$ is the speed of light. Dividing by the energy of each photon $E$ and a factor of $c$ gives the photon flux emitted into a hemisphere from a planar unit surface, specifically

$$\phi(E) dE = \frac{2\pi}{\hbar c^2} \frac{E^3}{\exp(E/k_B T) - 1} dE$$

Multiplying by a factor of two to account for photon flux from both sides of the planar device, and converting to units of mA cm$^{-2}$ eV$^{-1}$, gives the spectral photon flux over a range of temperatures (Figure 8a). Since for a blackbody emission and absorption are equivalent, the thermally generated $J_d$ can simply be determined from Equation (8).

Three distinct variations to EQE spectra are made, specifically the maximum magnitude EQE$_{\text{max}}$, the optical bandgap $E_0$ and the energetic disorder $\sigma$. As for the injection-limited $J_d$, we define a reference case (black solid line) with EQE$_{\text{max}} = 0.1$, $E_0 = 0.12$ eV, and $\sigma = 0.1$ eV. The value of $E_0$ determines the edge of the EQE spectrum as defined in Figure S4 (Supporting Information). In particular, choosing $E_0 = 0.12$ eV enables a direct comparison with the reference OPD in Figure 7 ($\Phi_b = 0.1$ eV, therefore $E_g = 0.1$ eV in case of OPD with Ohmic contacts and without blocking layers), assuming energetic offset of 0.2 eV to achieve efficient CT from donor to acceptor.\cite{Johansson2013} As for the injection-limited $J_d$, two parameters are held constant while the influence of the other parameter on $J_d$ is varied. Unsurprisingly, a larger EQE contribution at low energies increases the thermal dark current (Figure 8b), demonstrating the impact of sub-bandgap absorption. However, perhaps the most important point is that for a similar material parameter set, the injection-limited $J_d$ (dotted line) is about 5 orders of magnitude above the thermally generated $J_d$ (black solid line), meaning that charge injection can be regarded as the dominant contribution to the dark current density.

6. Why Does the Experimental $J_d$ Deviate from its Intrinsic Limit?

Having explored the intrinsic limits of dark current density in Section 5, we now consider factors that may cause deviation of the experimental value from this calculated intrinsic lower limit. This deviation is illustrated by considering that a reference OPD
with typical parameters of $L = 100$ nm, $E_g = 1$ eV, $\mu_0 = 10^{-7}$ m$^2$ V$^{-1}$ s$^{-1}$, and $\sigma_i = 0.1$ eV was calculated (at an effective electric field $F = 6 \times 10^6$ V m$^{-1}$) to have an intrinsic $J_d \approx 10^{-6}$ mA cm$^{-2}$, close to the empirical limit in Figure 3b assuming $E_g = 1$ eV.

While a few data points in Figure 3b lie near the empirical limit, a clear majority of literature $J_d$ values are significantly higher. One explanation for this discrepancy lies in the experimental uncertainty in determining LUMO A and HOMO D, as their absolute value can depend on the measurement technique.[75] As an example, the HOMO energy of PCDTBT was reported to be $-5.5$[52,34] or $-5.4$ eV.[13] Furthermore, reported LUMO energies of fullerene acceptors range between $-4.3$[52] and $-3.8$ eV.[13,75] While these uncertainties might affect the results in Figure 3b, they would not explain deviations from the $J_d$ empirical limits by several orders of magnitude. Instead, in many cases $J_d$ values are raised above the intrinsic limit by extrinsic factors, associated either with the OPD fabrication process or the $J_d$ measurement protocol. Pinholes in the photoactive blend can lead locally to high current densities, especially for thinner layers. In some cases, lateral leakage current paths outside the active area effectively increase $J_d$. This effect can be studied by measuring OPDs with different active area and checking if $J_d$ is independent of the active area. Similarly, local current density variation between the edge and the center of the device lead to area-dependent $J_d$ values. Furthermore, displacement currents become non-negligible for state-of-the-art OPDs with low intrinsic dark currents, particularly around $V = 0$ V; such contributions can be ruled out ensuring that $J_d$ is independent of the $J$–$V$ scan rate and scan direction.

In addition, the dark current density in OPDs can be affected by energy states within the bandgap of the active layer, often referred to as traps. Close to the band edges (within $\pm 0.2$ eV) lie tail states due to intrinsic energetic site disorder of organic materials[76] (quantified above with $\sigma$) that originate from variation in local microstructure. At similar energies lie extrinsic shallow traps, broadly attributed to chemical impurities such as oxygen, moisture and chemical impurities introduced during synthesis. Deep traps reside further into the bandgap,[77] attributed primarily to conjugation breaks, which are either due to synthetic defects or induced by impurities.[57] The precise origin of trap states is a topic of extensive ongoing discussion, with a universal trap level at around $-3.6$ eV identified in 2012 for a variety of conjugated polymers and attributed to the formation of hydrated oxygen complexes.[78] More recent

![Figure 8](image_url)  
*Figure 8. Thermally generated dark current calculations. a) Blackbody thermal photon flux $\phi(E)$ incident on a planar device, and hypothetical EQE spectra with various magnitudes, bandgaps $E_g$ and energetic disorders $\sigma$. b) The temperature-dependent dark current is determined from the overlap integral between EQE($E$) and $\phi(E)$. The injection-limited dark current at $V = -3$ V for the reference case in Figure 7 is shown for comparison.*
work\textsuperscript{[79,80]} support this attribution to moisture, suggesting that water molecules are present in voids within the active layer, even when the device is processed under notionally inert conditions, with Zuo et al.\textsuperscript{[81]} recently demonstrating that they lie \( \approx 0.3\text{--}0.4 \text{ eV} \) above and below the HOMO and LUMO levels, respectively.

Regardless of their origin, both shallow traps/tail states and deep traps can enhance \( J_d \) by both reducing the energetic jump required for charge injection and increasing the density of thermally generated carriers, since random thermal excitations cause electrons to jump to an unoccupied trap level and subsequently to the acceptor LUMO. Indeed, Fallahpour et al.\textsuperscript{[73]} demonstrated that deep trap states within the photoactive layer can influence the transport mechanism of the OPD due to trap-mediated capture and emission of electrons from/into the HOMO and LUMO levels (Figure 9a,b). Using a drift-diffusion model to simulate the dark current density of a P3HT:PC\textsubscript{61}BM OPD, excellent agreement with the experimental \( J \text{--} V \) characteristics was obtained assuming the presence of deep and shallow traps (Figure 9c). This underlines the importance of designing photoactive materials with as low trap densities as possible to achieve high sensitivity OPDs.

Shekhar et al.\textsuperscript{[39]} studied the effect of deep subgap states on the dark current density of PHJ OPDs. Measuring EQE spectra extended up to 1600 nm, the presence of subgap states and/or defect-induced states in the proximity of the D-A interface was investigated. The reduction in \( J_d \) on exchanging copper phthalocyanine (CuPc) for TAPC and then for boron SubPc chloride was attributed to the smaller tail state density, and thus lower charge generation/recombination rate at the D-A interface (Figure 10). This suggests that the dark current density can be effectively reduced by minimizing the density of deep sub-gap states at the D-A junction.

In addition to \( J_d \), extrinsic trap states were found to affect other OPD metrics such as the absolute EQE and the speed of response (see Sections 2.1 and 2.2). Cowan et al.\textsuperscript{[82]} quantified the role of traps in a PCDTBT:PC\textsubscript{65}BM BHJ solar cell by adding PC\textsubscript{65}BM as a well-defined trap state impurity. The addition of PC\textsubscript{65}BM decreased the efficiency of photon-to-electron conversion, thereby decreasing the EQE. Furthermore, trap states at the electrode–semiconductor interface were found to be the main cause of the persistent photocconductivity behavior,\textsuperscript{[83]} decreasing the speed of response. In OPDs, these interface traps can originate if the HEL or EEL leads to oxidation of adjacent species, as it was identified for PEDOT-PSS.\textsuperscript{[84]} Water and oxygen must thus be excluded from the OPD to ensure stability and durability.\textsuperscript{[85]} Therefore, commercial OPDs often demand thin-film encapsulation\textsuperscript{[86]} and suitable device architecture\textsuperscript{[87]} to achieve viable lifetimes.

7. Conclusion

In summary, OPDs offer significant advantages over conventional inorganic photodetectors, most notably flexibility and bandgap tunability, which potentially facilitate new applications. Achieving sufficiently high-performance metrics requires lowering \( J_d \) as much as possible. An overview of dark current densities reported in recent years shows a large spread in \( J_d \) values that merits further analysis. We identified charge injection from the contacts into the semiconductor as the dominant intrinsic contribution to \( J_d \) and we found that state-of-the-art OPDs are close to the expected intrinsic limit. Thus, once extrinsic contributions such as leakage are eliminated, for a given effective bandgap, the dark current density can be reduced by minimizing the active layer energetic disorder and trap state density.

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Conflict of Interest

The authors declare no conflict of interest.

Keywords

bulk heterojunction, dark current, image sensors, organic photodetectors, traps

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