

# n-Type Doping of Organic Semiconductors: Immobilization via Covalent Anchoring

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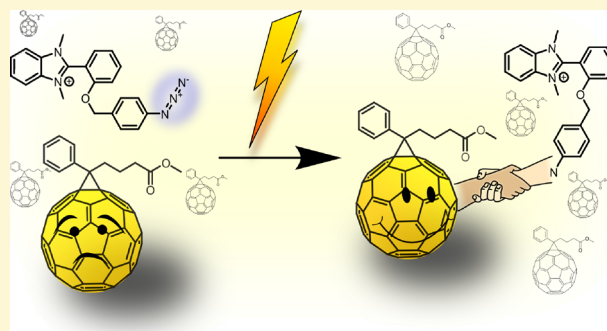
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## Supporting Information

**ABSTRACT:** Electrical doping is an important tool in the design of organic devices to modify charge carrier concentration in and Fermi level position of organic layers. The undesired diffusion of dopant molecules within common transport materials adversely affects both lifetime and device performance. To overcome this drawback, we developed a strategy to achieve immobilization of dopants through their covalent attachment to the semiconductor host molecules. Derivatization of the commonly employed n-type dopant 2-(2-methoxyphenyl)-1,3-dimethyl-2,3-dihydro-1H-benzimidazole (*o*-MeO-DMBI) with a phenylazide enables the resulting *o*-AzBnO-DMBI to photochemically generate a reactive nitrene, which subsequently binds covalently to the host material, 6,6-phenyl-C<sub>61</sub>-butyric acid methyl ester (PCBM). Both the activation and addition reactions are monitored by mass spectrometry as well as optical and photoelectron spectroscopy. A suppression of desorption and a decrease in volatility of the DMBI derivative in ultrahigh vacuum were observed after activation of a bilayer structure of PCBM and *o*-AzBnO-DMBI. Electrical measurements demonstrate that the immobilized *o*-AzBnO-DMBI can (i) dope the PCBM at conductivities comparable to values reported for *o*-MeO-DMBI in the literature and (ii) yield improved electrical stability measured in a lateral two terminal device geometry. Our immobilization strategy is not limited to the specific system presented herein but should also be applicable to other organic semiconductor–dopant combinations.



## INTRODUCTION

The applications and benefits of electrical or electrochemical doping in organic semiconductors are manifold.<sup>1,2</sup> Suitable dopants increase the conductivity of organic layers by several orders of magnitude, for example, in charge transport layers of organic light-emitting diodes (OLEDs).<sup>3</sup> Promising thermoelectric devices can be realized with doped conjugated polymers due to their low thermal conductivity.<sup>4–6</sup> Most importantly, doped layers enable quasi-ohmic contacts to metal electrodes, which greatly improves charge carrier injection and mitigates contact resistances due to offsets in energy level alignment.<sup>7–9</sup> A prerequisite for applying doped layers in dedicated device stacks is stability since diffusion of dopants into adjacent active layers has been identified as a degradation mechanism (e.g., exciton quenching), detrimental to photoactive devices.<sup>10–14</sup> Moreover, diffusion of dopants can be

problematic in organic hole transport layers for planar perovskite solar cells.<sup>15</sup> Consequently, stable and efficient doping of organic materials is of high interest in present-day device fabrication. Replacing atomic dopants such as halides or alkali metals by small molecular p- and n-type dopants reduces volatility significantly.<sup>16</sup> However, some host–dopant systems still exhibit diffusivity of organic electron donors/acceptors.<sup>17,18</sup> This effect is more pronounced within conjugated polymers compared to molecular host materials.<sup>19–21</sup> Even though the dopant lacks diffusion in the host matrix, this might change when an electric field is applied. Recently, the migration of molecular dopants in an electric field has been

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reported in poly(3-hexylthiophene) (P3HT) and is considered as a drift effect.<sup>22</sup> To suppress dopant diffusion and drift, certain strategies have been proposed and demonstrated, for example, introduction of a dense dopant blocking interlayer or filling of dopant pathways using a second redox-inactive transport material.<sup>23–25</sup> However, processing becomes more difficult, and possible complications concerning charge transport or energy level alignment arise from the additional dopant blocking materials.<sup>26</sup> Furthermore, polar side chains were introduced to a polymer to increase its dopant affinity, yielding an improved thermal stability of molecular dopants and a higher melting temperature.<sup>25</sup> Larger molecular dopants offer a better morphological stability as demonstrated by thermally activated diffusion of dopants in molecular host systems.<sup>27</sup> Besides physical modifications such as size or morphology, increased intermolecular interaction also reduces the mobility of dopants. These include dispersion force, dipole–dipole interactions,  $\pi$ – $\pi$  interactions, or even hybridization.<sup>28</sup> Charged dopants are much less diffusive than neutral dopants due to electrostatic interactions.<sup>21,29</sup> A covalent bond between the dopant and the host molecule would allow much stronger “fixation” ideally for the immobilization of both neutral and charged dopants. A premodification of the host with a functional group that introduces self-doping was already used to localize dopants in assembled thin films.<sup>30,31</sup> For the covalent bond formation after blending, the host and dopant have to have complementary functional groups.<sup>32–34</sup> This concept requires modification of each transport material to fit the corresponding dopant, which is not practical given the plethora of materials in use. In order to be generally compatible with a wide scope of host materials, an anchor group for dopants needs an unspecific binding method, for example, by attachment to aliphatic C–H or olefinic C=C bonds present in almost any organic semiconductor without introduction of severe degradation or traps. A practical solution for this issue can be derived from photochemical cross-linking as the desolubilization protocol.<sup>35</sup> The utilization of various bis-azides for cross-linking, thus preventing polymer strands from redissolution in solution-processed heterostructures, is not limited to one specific polymer.<sup>36,37</sup> The azide is activated by deep-ultraviolet light (UV, 254 nm), inducing photolysis and generating reactive singlet nitrenes. When properly substituted, these predominantly insert into aliphatic CH bonds, and side reactions<sup>38</sup> involving triplet pathways, aromatic insertion, attack upon other functional groups, or ketenimine ring expansion are suppressed.<sup>36,39,40</sup> The latter is found to be dominating only in solution for non-fluorinated aryl azides,<sup>39,41</sup> whereas in thin films or matrices, CH insertion prevails.<sup>42–45</sup>

1,3-Dimethyl-2-phenyl-2,3-dihydro-1*H*-benzimidazole (DMBI) derivatives are effective, air-stable, and thus easy-to-handle, solution-processable n-type dopants. DMBIs are often used to dope n-channel semiconductors, preferably PCBM<sup>46–51</sup> or n-type polymers.<sup>47,52</sup> However, they seem to be more effective, in terms of achieved maximum conductivity, in combination with PCBM than with polymers like P(NDI2OD-T2) due to the poor miscibility of the dopant and polymer.<sup>6,47,53</sup> Although the doping mechanism of DMBI derivatives in, for example, PCBM has been investigated, it is still under debate.<sup>48,51,54</sup> The initial step either involves a hydride transfer to the host or a hydrogen radical transfer with a subsequent electron transfer—both mechanisms lead to the

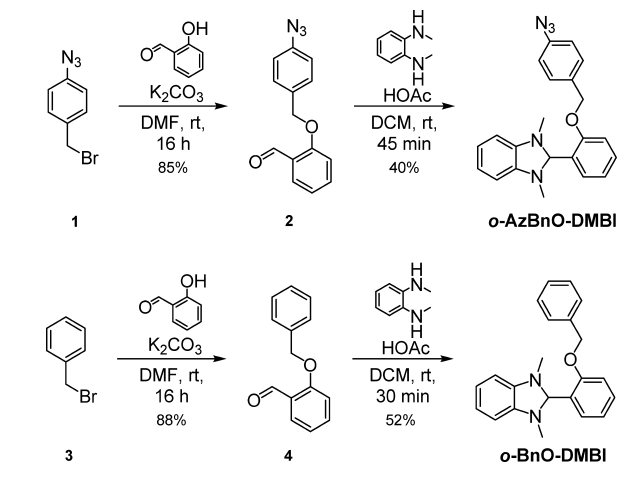
generation of cationic DMBIs and PCBM radical anions responsible for the doping effect.

In this work, we report the successful anchoring of an azide-functionalized DMBI-based dopant in PCBM, employing a nonspecific immobilization strategy through covalent attachment to the host while retaining its dopant strength compared to a nonfunctionalized counterpart. We detail the synthesis of *o*-AzBnO-DMBI and *o*-BnO-DMBI, in which the *ortho*-methoxy group of *o*-MeO-DMBI<sup>55,56</sup> is substituted by a (4-azidobenzyl)oxy and a benzyloxy functional group, respectively, and investigate their doping and anchoring properties in blends with PCBM. Infrared and photoelectron spectroscopies are employed to analyze the CH insertion or cycloaddition reaction, and conductivity measurement was employed to verify doping with and without a UV trigger. High-resolution mass spectrometry verifies a covalent linkage of the dopant to PCBM. Our immobilization strategy via nitrene generation is generally applicable and not limited to a specific host–dopant system. Furthermore, it is destined to improve the stability, lifetime, and performance of doped organic devices by ideally suppressing both detrimental diffusion and drift effects.

## RESULTS AND DISCUSSION

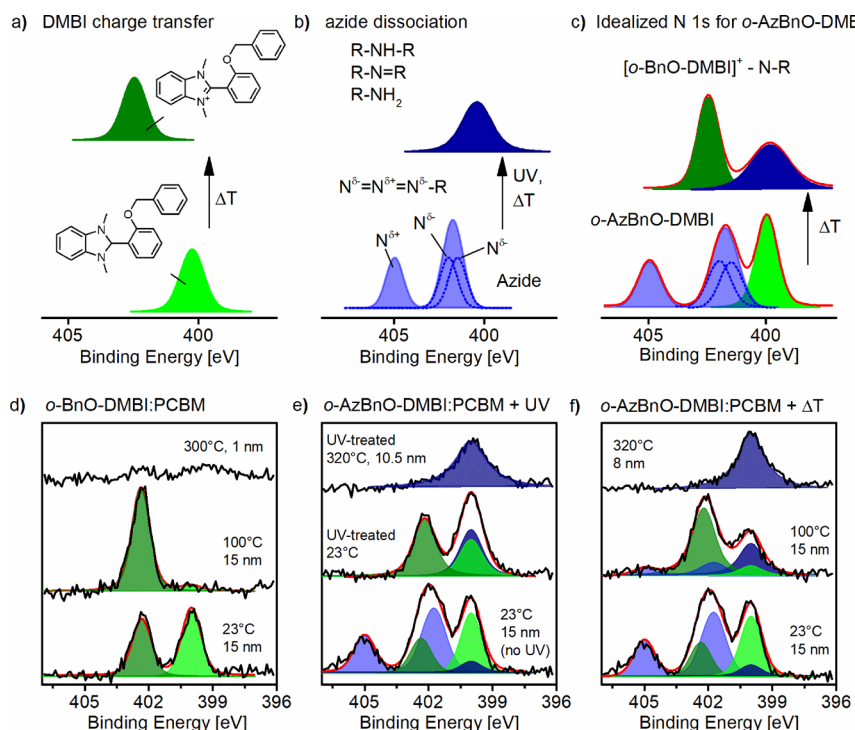
**Synthesis and Immobilization Reactions.** DMBI dopants are easily accessible via simple condensation reactions according to Scheme 1. Introduction of a phenylazide is

**Scheme 1.** Synthesis of *o*-AzBnO-DMBI and *o*-BnO-DMBI



achieved after etherification of salicylaldehyde employing 1-azido-4-(bromomethyl)benzene (1) in DMF at room temperature furnishing compound 2 in 85% yield, followed by condensation with *N,N*-dimethylbenzene-1,2-diamine to afford *o*-AzBnO-DMBI in 40% yield. The characteristic IR absorption band of the azide stretching vibration at  $2109\text{ cm}^{-1}$  (see Figure 4b and Figure S11) confirms the azide functionalization.<sup>57</sup>

To elucidate the influence of the azide group on immobilization, we synthesized the non-azido-functionalized analogue and thus nonimmobilizable *o*-BnO-DMBI employing benzyl bromide (3) for the Williamson ether synthesis. Both new dopants were isolated after crystallization from petroleum ether/methanol or DCM/hexane, respectively. The obtained colorless single crystals were suitable for X-ray crystal analyses, unambiguously proving their structure (see the Supporting Information). In order to optimize the processing parameters,



**Figure 1.** XPS measurements of the N 1s core-level spectra of thermally annealed PCBM films doped with *o*-BnO-DMBI and *o*-AzBnO-DMBI in panels (d–f). (a–c) Schematic cartoon of N 1s peak components and their transformation. (d) Recorded spectra of PCBM doped with *o*-BnO-DMBI for different annealing temperatures. (e) N 1s signal of PCBM doped with *o*-AzBnO-DMBI together with the spectrum measured after UV activation and after annealing of the UV-activated film to 320 °C. (f) Direct annealing of the *o*-AzBnO-DMBI/PCBM film without UV treatment. The assignment of peak composition in panels (d–f) is inferred from contributions of DMBI in green and from the azide in blue. The expected peak components of DMBI upon charge transfer and decomposition of pure azide are sketched in panels (a) and (b), respectively. For *o*-AzBnO-DMBI, both reactions of panels (a) and (b) can occur as illustrated in panel (c). The peak shapes of panels (a) and (b) are fitted to the data in panels (e–f) using a constrained model (see the Supporting Information for details). In the component fit, there is already a considerable fraction of charged dopants in panels (d–f) and possibly dissociated azide in panels (e) and (f) at room temperature. The initial film thickness at room temperature is 15 nm and changes in panels (d–f) for higher temperatures.

we studied the stability of the newly synthesized dopants and the literature known *o*-MeO-DMBI in solution (see the Supporting Information, Figure S13). NMR spectroscopy revealed that the dopants are only fairly stable in solution: after 1 day at ambient conditions (sunlight and air) in chloroform, oxidized species can be seen. However, the dopant appears to be stable for processing within several hours. In the solid state at  $-10$  °C, however, no degradation was observed over months.

In acetonitrile, the UV–vis spectrum of *o*-BnO-DMBI exhibits absorption features at  $\lambda = 221$ , 274, and 313 nm, whereas for *o*-AzBnO-DMBI, an additional absorption band at 253 nm is detected and attributed to the azide functional group (Figure S14).<sup>42,58</sup> Irradiation at 254 nm excites the azide and induces nitrene generation.<sup>42,45</sup> Throughout this work, this wavelength is used to initiate the immobilization process. Thermogravimetric analysis (see Figure S15) of *o*-AzBnO-DMBI shows irreversible mass loss at 117 °C corresponding to the extrusion of nitrogen. During additional heating/cooling cycles below the decomposition temperature of 240 °C, no further mass loss is observed. Similar to the azide-functionalized dopant, *o*-BnO-DMBI exhibits a rather low melting point at 108 °C (86 °C for *o*-AzBnO-DMBI) and decomposes at 200 °C. To achieve maximum doping effects for the host/DMBI blend, annealing at 75 °C is required.<sup>48</sup> Thus, *o*-AzBnO-DMBI is stable enough to test the doping of the PCBM host upon thermal treatment.

In the following, we provide spectroscopic data to corroborate successful anchoring of the *o*-AzBnO-DMBI dopant. From X-ray photoelectron spectroscopy (XPS) core-level spectra, we gain insight into the activation mechanism of both the dopant and the anchor group as well as possible decomposition reactions. In order to additionally test the thermal stability of the DMBI derivative in the blend, we annealed thin films of high doping concentration (50 mol %) in ultrahigh vacuum (UHV,  $10^{-9}$  mbar) and measured XPS at each temperature step (for details, see Figure S20). The N 1s emission of *o*-AzBnO-DMBI contains a contribution of the functional azide group and the DMBI core, illustrated in Figure 1c in blue (azide) and in green (DMBI).

To assign the N 1s peak components of *o*-AzBnO-DMBI correctly, it is instructive to analyze spectral changes expected from DMBI and the azide group, sketched in Figures 1a and 1b, respectively. The spectra of *o*-BnO-DMBI in Figure 1d only exhibit peak components of the DMBI core located at 402.5 and 400 eV. We identify the nitrogen emission at 402.5 eV as charged dopants having formed a radical and transferred an electron to PCBM. This assignment agrees with the N 1s emission of the ionic DMBI salt at about 402.5 eV reported in the literature.<sup>59</sup> With rising temperature, the peak component at 400 eV transforms into charged DMBI species at 402.5 eV, which correlates with an increase in conductivity observed for annealed PCBM films doped with DMBI. The N 1s signal reported for pure azides sketched in Figure 1b has three

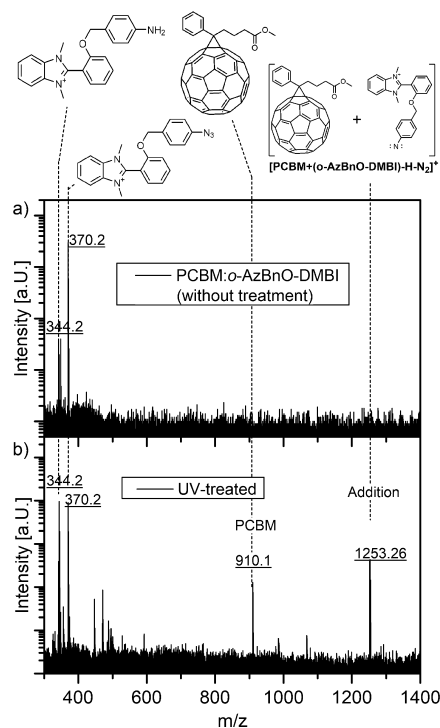


components, whereas two components coincide between 400 and 402 eV leaving a single peak component isolated at high binding energies.<sup>60,61</sup> From density functional theory (DFT) calculations, it is known that the central nitrogen in  $R-N=N=N^{\delta+}=N^{\delta-}$  is electron-deficient and has a N 1s level at high binding energies.<sup>62</sup> Azide decomposition triggered by UV light or thermal energy leads to nitrogen release and ideally to the generation of singlet nitrene, which can then form new carbon–nitrogen bonds. Regarding the azide in *o*-AzBnO-DMBI, the peak component at 405 eV in Figure 1e was therefore assigned to the central nitrogen, and the contributions of the lateral nitrogen atoms were between 400 and 402 eV. Upon UV activation at room temperature in Figure 1e, the former peak at 405 eV vanishes completely, reducing the peak area around 402 eV and increasing at 400 eV. During annealing of the doped PCBM film in Figure 1f, the N 1s emission at 405 eV disappears above 100 °C suggesting that the azide group becomes thermally activated and produces a nitrene. As a result, carbon–nitrogen bonds are formed causing an additional N 1s emission at 400 eV, which persists at temperatures higher than 300 °C. At the same time, the DMBI part of *o*-AzBnO-DMBI donates an electron, similar to *o*-BnO-DMBI, thus transferring the peak component at 400 eV into the charged DMBI species at 402.5 eV. When reaching higher temperature in UHV, pure and *o*-BnO-DMBI-doped PCBM completely evaporate from the substrate at 300 and 200 °C, respectively. Interestingly, the film doped with *o*-AzBnO-DMBI is stable even beyond 300 °C, and only a partial decrease in coverage is observed. However, from the N 1s emission, we infer that the N 1s peak component at 402.5 eV, attributed to charged dopants, disappears almost completely above 210 °C leaving only carbon–nitrogen bonds at 400 eV when approaching 320 °C in Figure 1 (for details, see Figure S20).

At room temperature, pure and untreated *o*-AzBnO-DMBI is volatile under UHV conditions as observed by XPS. However, when *o*-AzBnO-DMBI is sequentially processed on top of PCBM in monolayer-range thickness, a desorption experiment in UHV at room temperature reveals that the anchoring of *o*-AzBnO-DMBI also appears to be quantitative, rendering *o*-AzBnO-DMBI completely resistant to desorption after UV activation (see the Supporting Information for details). A similar trend is observed for thicker bilayer structures, which means that in pure dopant films *o*-AzBnO-DMBI may also react with neighboring dopants (see Figure S19).

Concerning the anchoring mechanism, it is known that azides react with fullerenes in a 1,3-dipolar [3 + 2] cycloadditions affording triazolines,<sup>63,64</sup> which, upon thermal treatment, undergo loss of nitrogen,<sup>65,66</sup> or [2 + 1] cycloaddition with nitrenes yielding aziridinofullerenes or azafulleroids.<sup>67,68</sup> Additionally, CH insertion into the alkyl chains of PCBM, though much less probable, may occur. To demonstrate covalent attachment of the dopant to the host, blends of *o*-AzBnO-DMBI and PCBM were analyzed by electrospray ionization (ESI) and matrix-assisted laser desorption ionization (MALDI) high-resolution mass spectrometry. We prepared blends of PCBM and *o*-AzBnO-DMBI at 25 mol % in a solution of chlorobenzene, which were subsequently dried in vacuo. Azide activation was achieved by either annealing at 135 °C or UV activation of spin-cast thin films, which were redissolved, and the material was dried afterward. Whereas no [3 + 2] cycloaddition products are

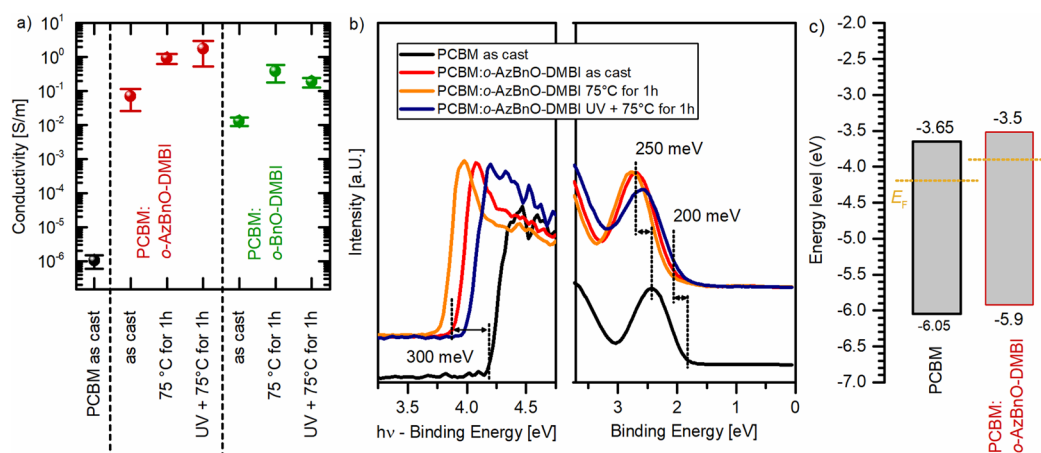
found after simple blending, additional peaks of covalent PCBM-nitrene adducts at 1253.2 *m/z* (attributed to  $[PCBM + (o\text{-AzBnO-DMBI})\text{-H-N}_2]^+$ ) are detected after heating to 135 °C (thermal azide generation) or photochemical activation (Figure 2 and Figures S24 and S26). Photochemical activation



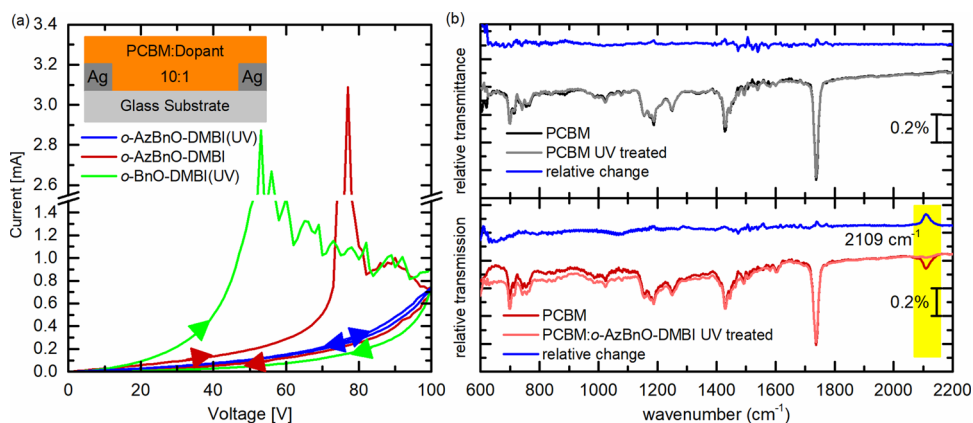
**Figure 2.** MALDI high-resolution mass spectrometry data are plotted for (a) a blend of PCBM and *o*-AzBnO-DMBI and (b) after UV treatment on a log scale. In principle, the MALDI laser used for desorption can cause azide decomposition, but we find a pronounced main peak at 370.2 *m/z* assigned to intact *o*-AzBnO-DMBI in panel (a). Comparable ESI spectra can be found in Figures S24–S26. Only trace amounts of unfavorable amine products are detected at 344.2 *m/z* in both panels (a) and (b) relative to *o*-AzBnO-DMBI at 370.2 *m/z*.

is the more benign method as shown in the ESI spectra (Figure S25); fragmentation of the ether bond within the thermal activated blends is detected. We find comparably small mass peaks for amines (these may also be a fragment of mass spectroscopy created during ionization)<sup>69,70</sup> and unreacted azide from the UV-activated blends (Figure 2). We attribute the latter to the slightly modified sample preparation for mass spectroscopy because XPS and IR experiments (Figures 3 and 4, respectively) demonstrate quantitative azide photolysis.

**Doping and Functional Characterization.** The conductivity measurements were performed under nitrogen on pure and doped PCBM layers (host-to-dopant ratio, 10:1), spin-cast from chlorobenzene onto prestructured silver contacts on glass substrates (Figure 3a and Figure S17). Whereas neat PCBM thin films exhibit conductivities in the range of 10<sup>-6</sup> S/m, this characteristic value rises up to 0.1 and 0.01 S/m for as-cast films doped with *o*-AzBnO-DMBI and *o*-BnO-DMBI, respectively. The final annealing step for doped films at 75 °C increases the conductivity further to 1 S/m. After UV treatment (254 nm) of as-cast films, the conductivities of the doped PCBM layers decrease by roughly half an order of magnitude in comparison to as-cast blends. The subsequent thermal treatment (1 h at 75 °C) increases the



**Figure 3.** (a) Conductivities of pure and doped PCBM layers with a molar doping ratio of 10:1 as-cast, after UV treatment, and after subsequent annealing. (b) UPS spectra of pure and doped PCBM layers with a molar doping ratio of 10:1 as-cast, after UV treatment, and after subsequent annealing. The secondary electron cutoff with respect to the vacuum level and the HOMO onset with respect to the Fermi level are given. (c) Calculated energy levels of pure and doped PCBM layers with a band gap of 2.4 eV.



**Figure 4.** (a)  $I$ – $V$  curves of annealed layers as-cast and after UV treatment. (b) Relative IR transmission spectra of 30 nm thick pure and doped PCBM layers with a molar doping ratio of 10:1. The spectra were measured on as-cast layers and after UV treatment as given in the legend. Dividing the spectra after the UV treatment by the as-cast spectra reveal the relative spectral change caused by the UV treatment. The vibrational mode of the azide group at 2109  $\text{cm}^{-1}$  is marked in panel (b) and strongly reduced by the UV treatment.

conductivities for both dopants by more than 2 orders of magnitude to 1 S/m. In total, this is a gain of 6 orders of magnitude compared to neat PCBM layers (see Figure 3a).

UV photoelectron spectroscopic (UPS) measurements of corresponding PCBM layers on silicon are presented in Figure 3b. The secondary electron cutoffs of *o*-AzBnO-DMBI-doped PCBM reveal strong shifts, which translate to a reduction of the work function of about 300 meV compared to the neat PCBM layer. Additionally, in doped layers, the highest occupied molecular orbital (HOMO) broadens, and the HOMO maximum shifts by 250 meV to higher binding energies compared to undoped PCBM and is accompanied by a shift of the core-level emission lines (see Figure S19). For comparison, Naab et al. observed a Fermi level shift of 500 meV in a PCBM blend with 1 wt % *o*-MeO-DMBI on ITO.<sup>51</sup>

The Fermi level shift goes along with an increase of free charge carrier concentration and thus leads to an increased conductivity revealing a strong n-type doping effect of both DMBI derivatives in PCBM. Annealing of the thin film results in an additional shift of the Fermi level toward the LUMO of about 50–100 meV for *o*-AzBnO-DMBI arriving at a total 350–400 meV shift. Furthermore, we find a broadening of the emission of the HOMO level, yielding an increase of tail states

for doped PCBM films. The results of the UPS measurements were used to calculate the energy level diagrams of pure and doped PCBM layers that are depicted in Figure 3c, assuming a constant band gap of about 2–2.5 eV.<sup>71–73</sup>

Figure 4a shows  $I$ – $V$  curves of doped *o*-AzBnO-DMBI and *o*-BnO-DMBI PCBM layers as-cast and after UV treatment. At high electric field strength, the untreated films including *o*-AzBnO-DMBI feature a pronounced rise and subsequent falloff in current. As a result, the  $I$ – $V$  curve displays a distinct hysteresis, which can be also observed when repeating the cycle (slightly shifted to higher voltages). This can be explained either by contact modification and/or by ionic migration. A drift of charged dopant molecules away from a contact causes a similar hysteresis in  $I$ – $V$  curves, which was found for films of poly(3-hexylthiophen-2,5-diyl) (P3HT) doped with the p-dopants 2,3,5,6-tetrafluoro-7,7,8,8-tetracyanoquinodimethane (F4TCNQ) and molybdenum tris[1-(methoxycarbonyl)-2-(trifluoromethyl)-ethane-1,2-dithiolen] ( $\text{Mo}(\text{tfd-CO}_2\text{Me})_3$ ).<sup>22</sup> On the other hand, the  $I$ – $V$  curve of an *o*-AzBnO-DMBI doped layer that was UV-treated (followed by a final annealing step) displays no or reduced hysteresis on average. This observation is a first indication for a successful dopant immobilization after nitrene generation.

Moreover, hysteresis persists in *o*-BnO-DMBI-doped and UV-treated PCBM, suggesting that the observed characteristic is not due to changes in morphology during UV irradiation but due to the reactions of *o*-AzBnO-DMBI.

The HOMO level of PCBM broadens after UV exposure, which is attributed to an oxidation of the thin film surface most likely from residual oxygen in the glovebox triggered by UV light (measured by XPS in Figure S18). IR transmission measurements of PCBM layers before and after UV exposure, given in Figure 4b, do not show an oxidation of the PCBM bulk material, meaning the observed oxidation takes place only at the surface. Unfortunately, this hinders the further evaluation of the UV-treated doped PCBM layers by UPS since the observed surface degradation strongly influences the measured Fermi level position. IR transmission measurements of PCBM layers doped with *o*-AzBnO-DMBI before and after UV treatment are presented in Figure 4b (see the Supporting Information for additional spectra). Due to the doping concentration of 10 mol %, most of the observed vibrational modes stem from PCBM and fail to reveal a change caused by the UV treatment. Only the characteristic vibrational mode of the dopant's azide group at 2109 cm<sup>-1</sup> vanishes completely, revealing a quantitative activation and dissociation of the azide by UV light.

## CONCLUSIONS

In this work, we detail the synthesis of *o*-AzBnO-DMBI, a derivative of the literature known *o*-MeO-DMBI. An azidophenyl group is introduced to DMBI by etherification of salicylaldehyde employing 1-azido-4-(bromomethyl)benzene and a subsequent condensation with *N*<sup>1</sup>,*N*<sup>2</sup>-dimethylbenzene-1,2-diamine. The modification of DMBI with a reactive azide aims at providing a universally applicable anchor-like functional group, which should be able to covalently bind to a wide scope of common organic semiconductors. By creating a covalent bond with the host material, the dopant is expected to be immobilized or at least rendered significantly less mobile, which ideally reduces problems in devices arising from diffusion or drift of dopants. In an exemplary study, *o*-AzBnO-DMBI was blended with PCBM and activated by UV light in order to generate highly reactive nitrenes to ultimately form covalent carbon–nitrogen bonds between PCBM and the dopant. The azide activation was monitored by infrared and photoelectron spectroscopy. High-resolution mass spectrometry reveals addition products, firmly demonstrating the covalent attachment of the dopant to PCBM molecules. The thermal stability of *o*-AzBnO-DMBI in the blend was tested by annealing thin films of high doping concentration in UHV and by measuring temperature-dependent XPS. We find that the activated dopant renders the doped film more resistant to desorption when approaching the desorption temperature of PCBM in UHV. At room temperature, activation of *o*-AzBnO-DMBI in a bilayer structure was shown to prevent desorption from PCBM under UHV. *I*–*V* measurements further prove that UV-activated *o*-AzBnO-DMBI maintains its doping properties giving rise to conductivities typical for blends of DMBI and PCBM. At high electric field strength, the UV-activated doped PCBM films show no or neglectable hysteresis, but a pronounced hysteresis is observed if *o*-AzBnO-DMBI is not exposed to UV light. The increased electrical stability can be explained by a suppression of dopant drift due to anchoring of the dopant to PCBM after activation.

The great advantage of our approach is the use of unspecific binding of the intermediately generated reactive dopant toward the host material, offering instant application to a plethora of organic semiconductors. Although chemical derivatization of dopants, either *n*- or *p*-type, may be challenging due to the instabilities resulting from their extreme frontier molecular orbital energies, azide incorporation is possible and has the potential to reduce undesired drift and diffusion in multilayer stacks of OLEDs or OPV and can be applied for *p*–*n* homojunctions based on polymers. The appropriate chemical derivatization of *p*-type dopants such as F4TCNQ and F6TCNNQ is ongoing in our laboratories.

## EXPERIMENTAL SECTION

**Synthesis.** All reactions requiring exclusion of oxygen and moisture were carried out in heat-gun-dried glassware under a dry and oxygen-free nitrogen or argon atmosphere using Schlenk and glovebox techniques. All reagents were obtained from commercial suppliers and were used without further purification if not otherwise stated. Deuterated solvents were purchased from Sigma-Aldrich Laborchemikalien GmbH (Seelze, Germany).

**Sample Preparation for Analysis.** If not noted otherwise, the dopants and PCBM were dissolved separately in chlorobenzene at a concentration of 5 g L<sup>-1</sup> for XPS and 10 g L<sup>-1</sup> for IR/*I*–*V* measurements and were mixed right before spin-coating. The dopants were kept in solution for 5–10 min, but the PCBM solution was stirred overnight at 50 °C. The films were spin-coated at 1000 rpm for 60 s in a nitrogen glovebox on a cut *n*-doped silicon wafer for XPS, intrinsic silicon for IR, and on glass for *I*–*V* measurements. Activation by UV light was performed with a GPH135TSL/4 mercury-vapor lamp from Peschl Ultraviolet with a nominal UV-C power of 1.2 W at 254 nm in the glovebox. The penlight was mounted on a stand at a distance of 10 cm above the sample. We measured the UV intensity at the sample position using a Newport power meter 1936-C equipped with a 818-UV photodiode plus attenuator and obtain a power density of 0.8 mW cm<sup>-2</sup> for a 20 min exposure time.

**IR Spectroscopy.** For IR transmission measurements, a Fourier transform IR spectrometer (Vertex 80v, Bruker) was used. To prevent absorptions from ambient air (H<sub>2</sub>O and CO<sub>2</sub>), the spectrometer was evacuated to 3 mbar. All samples were recorded near normal transmittance (7°) with an MCT detector and a resolution of 4 cm<sup>-1</sup>. For each spectrum, 200 scans were averaged.

**Electrical Measurements.** *I*–*V* measurements were performed under a nitrogen atmosphere at room temperature. For this purpose, finger-like structured silver contacts with channel lengths between 25 and 30 μm were used. Voltages were applied on these contacts using the 4155C semiconductor parameter analyzer by Agilent Technologies, which reads out corresponding currents for conductivity determination. Conductivities were then calculated from a set of at least three different samples.

**Photoelectron Spectroscopy.** Photoelectron spectroscopy (PES) measurements were carried out using a PHI5000 VersaProbe scanning photoelectron spectrometer, which was equipped with a monochromated Al Kα X-ray source at 1486.7 eV photon energy for X-ray photoelectron spectroscopy (XPS) and a differentially pumped helium discharge lamp operated to achieve He I emission at 21.2 eV photon energy for ultraviolet photoelectron spectroscopy (UPS). The focused X-ray spot was approximately 200 μm in diameter and was moved to a different position on the sample for longer acquisition in order to avoid radiation damage.

**Mass Spectrometry.** The mass spectra were recorded using the following instruments: JEOL JMS-700 magnetic sector (EI); Bruker ApexQehybrid 9.4 T FT-ICR (ESI, DART); Finnigan LCQquadropole ion trap (DART); and Bruker Autoflex Speed (MALDI).



## ■ ASSOCIATED CONTENT

### Supporting Information

The Supporting Information is available free of charge on the ACS Publications website at DOI: [10.1021/acs.chemmater.9b01150](https://doi.org/10.1021/acs.chemmater.9b01150).

Synthesis, NMR spectroscopy, crystal structure, IR spectroscopy, NMR stability test, UV–vis absorption, TGA/DSC, ESI mass spectra, and XPS analysis (PDF)

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### Notes

The authors declare no competing financial interest.

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## ■ ABBREVIATIONS

(DMBI)	1,3-dimethyl-2-phenyl-2,3-dihydro-1H-benzoimidazole
( <i>o</i> -MeO-DMBI)	2-(2-methoxyphenyl)-1,3-dimethyl-2,3-dihydro-1H-benzoimidazole
( <i>o</i> -BnO-DMBI)	2-(2-(benzyloxy)phenyl)-1,3-dimethyl-2,3-dihydro-1H-benzoimidazole
( <i>o</i> -AzBnO-DMBI)	2-(2-((4-azidobenzyl)oxy)phenyl)-1,3-dimethyl-2,3-dihydro-1H-benzoimidazole

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