One-pot selective synthesis of a fullerene bisadduct for organic solar cell applications

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One-pot selective synthesis of a fullerene bisadduct for organic solar cell applications

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A single isomer of fullerene bisadduct, PC60PF, was obtained from commercially available fullerene derivative, PC60BM, in one pot over two steps. The tether-directed remote functionalization approach provided a very simple and fast method to produce single isomer of fullerene bisadducts with good yield and easy purification. Bulk heterojunction organic solar cells containing the bisadduct was fabricated and tested.

Over the past decade, organic solar cells (OSCs) have been a hot topic of research as an emerging renewable energy technology. In comparison with conventional devices, such as silicon solar cells, OSC technology promises lightweight flexible devices that can be produced at low cost through roll-to-roll printing methods. One specific type of OSC is the bulk-heterojunction (BHJ) solar cell which contains a blend of electron donor and acceptor materials in its active layer. Among electron acceptor materials, fullerenes are one of the most attractive compounds, due to their reversible reduction behavior, good electron transport properties and easily functionalized structures. Numerous fullerene derivatives have been reported and used as effective electron acceptor materials in OSC devices, such as [6,6]-phenyl-C61-butyric acid methyl ester (PC60BM) and indene C60 bisadduct (IC60BA).

The power conversion efficiency (PCE) of OSCs can be optimized by adjusting the device parameters of open-circuit voltage ($V_{oc}$), short-circuit current density ($J_{sc}$) and fill factor (FF). In particular, the $V_{oc}$ is relatively easy to control by tuning the highest occupied molecular orbital (HOMO) energy level of donor material and the lowest unoccupied molecular orbital (LUMO) energy level of acceptor material. A common strategy to increase the voltage is to elevate the LUMO energy level of the fullerene acceptor and this is normally achieved by attaching additional substituents to the fullerene core. There has been a few examples of fullerene bisadducts employed in OSC devices but these bisadduct materials are typically regioisomer mixtures. There has been a handful of studies looking at the effect of regioisomers on device performance thus far. In particular, our group isolated a crystalline single isomer of the indene C60 bisadduct (IC60BA) and examined its device performance. While the results were promising, the single isomer was obtained by careful chromatographic separation which meant that the production of material was both time consuming and difficult to scale up.

In order to target pure fullerene bisadduct materials in a more efficient manner, we decided to investigate a regioselective synthesis method, namely tether-directed remote functionalization. In general, this idea aims at increasing the regioselectivity of reactions on the fullerene surface by using a linker between the reactive head groups. This method has been successfully demonstrated on both C60 and C70 substrates and perfect regioselectivity as well as stereoselectivity has been achieved. In this work, the tether-directed functionalization approach was used to obtain selectively a C60 bisadduct, N-methyl-phenyl-C61-propyl-2-fulleropyrrolidine (PC60PF). A range of characterization methods were performed to show that the final product was a single regioisomer. To compare the properties of the single isomer with an isomeric mixture, a non-tethered PC60BM-based fulleropyrrolidine (PC60BM-FP) was also prepared (Figure 1). The single isomer of PC60PF as well as PC60BM-FP and PC60BM were used in BHJ OSC devices to examine their photovoltaic performance.

Figure 1. Chemical structures of single isomer, PC60PF, bisadduct isomeric mixture, PC60BM-FP, and widely used PC60BM.

The fullerene bisadduct PC60PF was synthesized from PC60BM in one pot over two steps in 53% yield. (Figure 2a) Firstly,
disobutylaluminium hydride (DIBAL) was added dropwise to a stirred solution of PC\textsubscript{60}BM at -78 °C in toluene under nitrogen atmosphere. After stirring for one hour, the reaction was quenched with methanol to give the crude aldehyde intermediate. Solvents were removed under vacuum from the reaction vessel and the aldehyde was used in the next step without isolation. The aldehyde was heated with N-methylglycine in 1,2-dichlorobenzene (o-DCB) at 180 °C with to form PC\textsubscript{60}PF. The 1,3-dipole cycloaddition step was monitored by thin layer chromatography (SiO\textsubscript{2}, tolune) to optimize the formation of the fulleropyrrolidine. The PC\textsubscript{60}PF product was purified by flash chromatography (SiO\textsubscript{2}, tolune). The purity of the product was analyzed using high performance liquid chromatography with a Cosmosil Buckyprep-D column which indicated the sample contained one major product (see Supporting Information). The isomeric mixture PC\textsubscript{60}BM-FP was synthesized from PC\textsubscript{60}BM and N-methylglycine in 29% yield (Figure 2).

![Figure 2](image)

Figure 2. Synthesis of fullerene bisadducts (a) PC\textsubscript{60}PF and (b) PC\textsubscript{60}BM-FP with reaction conditions: i. DIBAL, -78 °C, toluene, stir 1h; ii. N-methylglycine, o-DBC, reflux 1.5 h; iii. N-methylglycine, o-DBC, reflux 3 days. The nomenclature for fullerene bisadducts illustrated in (c).

The 1,3-dipolar cycloaddition on fullerenes usually results in reaction on the double bonds between six-member rings (namely the [6,6] bonds). In the case of C\textsubscript{60} bisadducts, there are eight potential reacting position for the second substituent, which can result in at least eight isomers (Figure 2c). Due to the length of the linker on PC\textsubscript{60}PF, possible regioisomers should be limited to the cis-hemisphere of the molecule. Notably, potential products will contain stereoisomers because of the flexibility of the linker.

To predict the configuration of the final product, theoretical calculations were carried out with Gaussian 09, applying density functional theory at the B3LYP level. The basis set of 6-311G, with polarization functions (d,p), was used in the calculations. Considering both the length of the three-member carbon chain and the result of theoretical calculation, the possible positions of the pyrrolidine substituent were narrowed down to the cis-1, cis-2 and cis-3 positions of C\textsubscript{60}. Apart from regioisomers, geometric isomers are also possible at the cis-1 position, namely cis-1a, cis-1a* (the H\textsubscript{6} on cis-1a* pointing toward the phenyl group) and cis-1b (Figure 3a). The relative energy of the transition state of the 1,3-dipolar cycloaddition as well as the energy of the start material and possible products were calculated (Figure 3b and Table S1). The starting point of the calculation was the azomethine intermediate which originated from the reaction between the aldehyde and the amino acid (see Supporting Information for details). The activation barriers in o-DCB solvent environment are estimated to be 14.03 kcal mol\textsuperscript{-1} for cis-1a, 21.10 kcal mol\textsuperscript{-1} for cis-1b and 42.67 kcal mol\textsuperscript{-1} for cis-3. On the other hand, stepwise cycloaddition is suggested for cis-1a* and cis-2 via the intermediates 1a*-IM and 2-IM with the rate-limiting barriers of 13.39 and 29.76 kcal mol\textsuperscript{-1}, respectively. Based on the computational results, cis-1a and cis-1a* are the kinetic products with the lowest activation energy, while cis-1a is also the thermodynamic product with the deepest product Gibbs free energy. These theoretical calculations were used to guide the structural assignments.

![Figure 3](image)

Figure 3. (a) Five possible configurations of PC\textsubscript{60}PF and (b) theoretical calculation results showing the relative energy of start point, transition states and final products of the four possible configurations.

Further data on structure assignment was obtained with UV-vis absorption spectroscopy (Figure 4). It is known that the UV-vis absorption spectrum of fullerene derivatives contains distinct bands and features for different regioisomers. This is related to the variations of the conjugated surface of the fullerene derivatives. The UV-vis spectrum of PC\textsubscript{60}PF showed an onset of 730 nm and a peak at 440 nm (Figure 4). These features are similar to that observed for a reported C\textsubscript{60}–benzyne bisadduct with cis-1 configuration. This observation is in agreement with the assignments in the NMR experiments and in the theoretical calculations. The 1\textsuperscript{H}-NMR spectrum of the single isomer of PC\textsubscript{60}PF was well-resolved (Figure 5a). Most interestingly, the six protons on the propyl linker all have different chemical shifts in the 1 to 5 ppm region of the NMR spectrum. These separated proton resonances strongly supported the fact that the propyl linker was locked in a fused cyclic system with the protons residing in different environments above...
the fullerene surface. This was also indicative of the regioselectivity of the reaction. Further indication of selectivity was evident by comparing the $^1$H NMR spectrum of the bisadduct mixture PC$_{60}$BM-PF to that of the PC$_{60}$PF sample (Figures 5a and b). Full assignments of the proton and carbon resonances of the substituents on PC$_{60}$PF was achieved using correlation spectroscopy and heteronuclear single quantum coherence spectroscopy 2D-NMR experiments (see Supporting information). With these assignments, it was possible to use heteronuclear multiple-bond correlation spectroscopy (HMBC) experiment to aid in the structural determination. By focusing on the C$_{10}$ carbon, it was apparent that there was coupling with protons H$_5$, H$_{2a}$ and H$_{2b}$ (Figure 5c). This correlation suggested a structural configuration that matched the cis-1a isomer. In addition, the configuration of the cis-1b isomer would lead to coupling of carbon C$_{10}$ with protons H$_{2a}$ and H$_{2b}$. This was not observed in the HMBC experiment.

![Figure 5](image)

Figure 5. (a) 1H NMR spectrum of PC$_{60}$PF and its simplified structure, (b) 1H NMR spectrum of PC$_{60}$BM-FP mixed isomers and (c) HMBC spectrum of PC$_{60}$PF showing the long distance coupling between C$_{10}$ and H$_{2a}$, H$_{2b}$, H$_5$.

The relative configuration of the PC$_{60}$PF material was confirmed by single crystal X-ray analysis (Figure 5). Good quality single crystals were obtained by recrystallization from o-DCB and petroleum spirit 40-60 °C. Crystals of PC$_{60}$PF consisted of a racemic mixture of the cis-1a isomer with o-DCB solvent incorporated in the crystal structure. The packing diagram of crystalline PC$_{60}$PF showed alternating rows of enantiomeric pairs with o-DCB filling gaps between PC$_{60}$PF molecules. (Figure 6). The nearest distance between fullerenes is 3.38 Å.

With the molecular structure of PC$_{60}$PF solved, relevant data on its materials properties were acquired. To estimate the LUMO energy levels of the materials, cyclic voltammetry was performed (Figure S14). Data was also obtained for the PC$_{60}$BM-FP mixture and PC$_{60}$BM for comparison (Table 1). The fullerene bisadducts PC$_{60}$PF and PC$_{60}$BM-FP had relatively higher LUMO energy levels of -3.76 eV and -3.66 eV respectively compared to that of PC$_{60}$BM at -3.85 eV. This was the predicted outcome and improved open circuit voltages were expected for devices using the bisadducts as the electron acceptor material. Thermal analysis, such as differential scanning calorimetry and thermo gravimetric analysis, were also performed on the materials (Figure S9 to S13). No phase transitions were observed up to 300 °C and the decomposition temperature (5% weight loss) for fullerene compounds PC$_{60}$PF and PC$_{60}$BM was more than 400 °C (Table 1).

![Figure 6](image)

Figure 6. Structure obtained by X-ray crystallography of PC$_{60}$PF grown from o-DCB and petroleum spirit: (a) thermal ellipsoid illustration, (b) packing diagram where disordered atoms have been omitted for clarity.

<table>
<thead>
<tr>
<th>Electron Acceptor Materials</th>
<th>$E_{\text{red}}^{1/2}$ (eV)</th>
<th>$E_{\text{LUMO}}$ (eV)</th>
<th>UV-vis abs $\lambda_{\text{max}}$ (nm)</th>
<th>$T_d$ (°C)</th>
</tr>
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<tr>
<td>PC$_{60}$PF</td>
<td>-1.04</td>
<td>-3.76</td>
<td>437 (8.7)</td>
<td>428</td>
</tr>
<tr>
<td>PC$_{60}$BM-FP</td>
<td>-1.15</td>
<td>-3.65</td>
<td>463 (4.7)</td>
<td>379</td>
</tr>
<tr>
<td>PC$_{60}$BM</td>
<td>-0.95</td>
<td>-3.85</td>
<td>432 (4.1)</td>
<td>432</td>
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Table 1. Characterization results of PC$_{60}$PF, PC$_{60}$BM-FP and PC$_{60}$BM.

$^a$ Half-wave reduction potential from cyclic voltammetry; $^b$ $E_{\text{LUMO}} = - (E_{\text{red}}^{1/2} + 4.8$) eV; $^c$ Absorption coefficient ($\times 10^3$ M$^{-1}$cm$^{-1}$) in brackets; $^d$ Decomposition temperature (5% weigh loss).

The final part of the study was to compare the solar cell performance of the single isomer PC$_{60}$PF material with PC$_{60}$BM and the isomer mixture PC$_{60}$BM-FP. Bulk heterojunction devices were made in the following geometry: ITO/PEDOT:PSS/active layer/Ca/Al, in which the active layer was a blend of poly(3-ethylthiophene) P3HT and each acceptor materials with the ratio of 1:1 by weight (Figure S15). The processing solvent was o-DCB and the thicknesses of the devices was optimized by adjusting solution concentration and spin coating speed (see Supporting Information). The device performance parameters are summarized in Table 2. It is clear that the $V_{oc}$ of devices increased significantly, from 0.58 to 0.70 V, when using PC$_{60}$PF instead PC$_{60}$BM as acceptor material (Figure S15). This is in agreement with previous literature reports of using fullerene bisadduct materials to boost the $V_{oc}$.$^4$ Comparing the devices containing PC$_{60}$PF and the isomer mixture PC$_{60}$BM-FP, both $J_{sc}$ and FF were higher for the PC$_{60}$PF devices (Table 2). A reason for this observation is the pure single isomer material PC$_{60}$PF should have better charge transport properties than the isomer mixture PC$_{60}$BM-FP (*vide infra*). The external quantum efficiency (EQE) spectrum of the devices were obtained to investigate the photo-current contribution of the active layers in OPV devices. The better performing device containing PC$_{60}$PF showed significant photocurrent enhancement from 300 nm to 630 nm comparing to
the devices containing PC$_{60}$BM-FP (Figure S15). The maximum EQE of device containing PC$_{60}$PF was 63% at 500 nm which was higher than the maximum EQE of 59% in the devices containing PC$_{60}$BM-FP. The EQE data of the PC$_{60}$BM device showed slightly higher efficiency across the spectrum. This could be attributed to the higher charge mobility of the PC$_{60}$BM material (vide infra). It is interesting to note that the higher $E_{LUMO}$ measured in electrochemical experiments for PC$_{60}$BM-FP was not reflected in the $V_{oc}$ of devices. Two factors may account for this observation. The PC$_{60}$BM-FP material consisted of a mixture of isomers and there could be energy level variations between the various species present in the sample. Another way to look at these results is to consider the variation in device optimization. Assuming the electrochemical data gave a true reflection on the $E_{LUMO}$ of the PC$_{60}$BM-FP material, then the $V_{oc}$ of device should be higher. Device optimizations, such as blend ratios and processing methods, have been shown to affect $V_{oc}$. It may be possible to improve the $V_{oc}$ of the PC$_{60}$BM-FP device through extensive device optimization.

Table 2. The performance of devices based on PC$_{60}$PF, PC$_{60}$BM-FP and PC$_{60}$BM with P3HT (1:1 blend ratio by weight).

<table>
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<tr>
<th>Active layer</th>
<th>$J_{sc}$ (mA/cm$^2$)</th>
<th>$V_{oc}$ (V)</th>
<th>FF (%)</th>
<th>PCE (%)</th>
<th>Electron mobility $10^4$ (cm$^2$/Vs)</th>
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<tr>
<td>P3HT PC$_{60}$BM</td>
<td>8.5 ±0.3</td>
<td>0.58 ±0.01</td>
<td>63 ±2</td>
<td>3.1 ±0.2</td>
<td>8.50</td>
</tr>
<tr>
<td>P3HT PC$_{60}$PF</td>
<td>8.1 ±0.3</td>
<td>0.70 ±0.02</td>
<td>58 ±3</td>
<td>3.3 ±0.3</td>
<td>3.80</td>
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<tr>
<td>P3HT PC$_{60}$BM-FP</td>
<td>7.7 ±0.4</td>
<td>0.69 ±0.02</td>
<td>50 ±3</td>
<td>2.7 ±0.3</td>
<td>1.35</td>
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To examine the charge transport behavior of the acceptor materials in the bulk heterojunction, the space charge limited current (SCLC) measurements were obtained (Figure S16). The electron mobility in devices containing PC$_{60}$PF and PC$_{60}$BM-FP were calculated to be 3.80 $\times$ 10$^{-4}$ and 1.35 $\times$ 10$^{-4}$ cm$^2$/V(s) respectively. This result showed that the single isomer of fullerene bisadduct had better electron mobility than the isomer mixture PC$_{60}$BM-FP in these BHJ devices. In addition, the hole mobility of the P3HT from SCLC was previously measured at 3.30 $\times$ 10$^{-4}$ cm$^2$/V(s) in BHJ devices. This meant that more balanced charge transport occurred in the P3HT/PC$_{60}$PF blend films. The surface morphology of the films were examined using atomic force microscopy. Nanoscale features were observed in the topography and phase images and were similar in all P3HT:fullerene blend films (Figure S17). This means the difference in the solar cell device performance between the PC$_{60}$PF and PC$_{60}$BM-FP materials can be attributed to their electron mobility in the BHJ blends.

Regioselective synthesis of a single C$_{60}$ bisadduct isomer, PC$_{60}$PF, was achieved efficiently in a one-pot two step procedure from commercially available PC$_{60}$BM. The tether-directed functionalization strategy was applied successfully to control the regioselectivity of the final product in the cis-1 configuration. In BHJ solar cell devices, the single isomer PC$_{60}$PF outperformed the isomeric mixture PC$_{60}$BM-FP and was on par with PC$_{60}$BM. Further research in modifying the properties, such as solubility and crystallinity, of PC$_{60}$PF derivatives by changing the amino acid in the synthesis process is currently in progress.

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Notes and references
