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The devices based on VC90:PC$_{71}$BM processed from CF and DIO (3%v)/CF exhibits power conversion efficiency of 4.58 % and 6.07 %, respectively.
Solution processed organic solar cells based on A-D-D’-D -A Small Molecule with Benzo[1,2-b:4,5-b’]dithiophene donor (D’) unit, cyclopentadithiophene donor (D) and ethylrhodanine acceptor unit having 6 % light to energy conversion efficiency

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Abstract

Solution processed small molecule A-π-D-π-A denoted as BDT(CDTRH)_2 consists of 2-ethylhexoxy substituted BDT (donor D’unit) as the central building block and 3-ethylrhodanine (RH) as end capped terminal (acceptor group) unit with π-linkage of cyclopentadithiophene (CDT) (donor D) have been designed and synthesized. We have investigated the optical and electrochemical properties of this small molecule and found that its energy levels are compatible with the energy levels of fullerene derivatives for efficient exciton dissociation. This small molecule has been used as electron donor along with PC_{71}BM as electron acceptor for the fabrication of solution processed small molecule bulk heterojunction (BHJ) solar cells. The BHJ solar cell processed with BDT(CDTRH)_2 :PC_{71}BM (1:1 wt ratio) showed power conversion efficiency (PCE) of 4.58 % with J_{sc}= 8.66 mA/cm^2, V_{oc}= 0.98 V and FF= 0.54. The high V_{oc} value of the device has been attributed to the deeper HOMO energy level of BDT(CDTRH)_2. The overall PCE of the device has been increased up to 6.02 % (J_{sc} =10.42 mA/cm^2, V_{oc} = 0.94 V  FF = 0.62) when the blend was processed with 3% v/v CN/CF solvent. This increase is mainly due to an increase in J_{sc} and FF, which was linked to the increase in crystallinity and favorable nanomorphology of the active layer improving exciton dissociation and achieving a more balanced charge transport in the device.

Key words: Donor-acceptor, cyclopentadithiophene, bulk heterojunction solar cells, solvent additives
Introduction

Organic solar cells based on solution processed active layers consisting of small molecule (SM):fullerene blends have attracted a lot of attention on account of their advantages of large area, flexible, light weight devices by roll to roll printing techniques, high purity and discrete molecular weight [1]. They can be a viable alternative to the polymer: fullerene systems as demonstrated by the impressive power conversion efficiencies in the range of 8 - 10 % [2]. At present the research on solution processed small molecule (SM)-based organics solar cell is targeted to achieve higher PCEs by designing and synthesizing new organic small molecules with D-A-D or A-D-A structures [3]. The most promising organic small molecule donor materials for photovoltaic applications are generally built by connecting various electron donating (donor) and electron accepting (acceptor) units through the \( \pi \)-conjugated spacer [4]. Among the various donor units used in the D-A copolymers, benzof[1,2-b:4,5-b']dithiophene (BDT) is one of the most promising electron-donor units for applications in high-performance polymer semiconductors. Its large planar \( \pi \)-conjugated structure promotes enhanced \( \pi-\pi \) stacking and improves hole mobility [5]. Compared to one dimensional (1D) BDT based conjugated polymers, the two dimensional (2D) BDT polymers based solar cells show high \( V_{OC} \) due to its deeper HOMO energy levels. Recently a D-A copolymer with benzodithiophene (D) unit showed a PCE of 8.3 % processed with 1-Chloronaphthalene (CN) as additive [6]. The efficient photovoltaic response of these 2D BDT copolymers makes them a promising donor materials for solution processed small molecule organic solar cells (SMOSCs) [7]. Therefore, small molecules with high efficiency could be designed by tailoring the HOMO and LUMO energy levels of the 2D BDT unit by introducing appropriate electron accepting units and \( \pi \)-bridge linkers.

A series of dye building blocks including 3-ethylrhodanine (RH), 1,3 indandione and 1,3 dimethylbarbituric acid have been introduced as electron accepting units along with electron rich oligothiophenes and/or BDT providing low bandgap small molecule donor materials [8]. A PCE up to 8.12 % has been achieved with a small molecule using thienyl benzodithiophene (BDT) as core and 3-ethylrhodanine as terminal group [8b]. Bazan et al. reported DTS(PTTh\(_2\))\(_2\) [9] and DTS(FBTTh\(_2\))\(_2\) [10] using dithienosilole (DTS) as a donor core with [1,2,5]-thiadiazolo-[3,4-c]-pyridine or fluorobenzothiadiazole as pendant moieties, both of which displayed promising photovoltaic performance with a PCE of up to 9.02% [10b]. Li et al. and Chen et al. synthesized two 2D small molecules, D2 and DR3TBDDTT
with an appropriate A-π-D-π-A molecular structure, which showed excellent photovoltaic performance with PCE up to 6.75 % and 8.12 %, respectively [11, 12].

Many promising end group acceptors, i.e. dicyanovinyl [13, 14], alkyl cyanoacetate [15-18] and 3-ethylrodanine [19, 20] have been used for conjugated small molecules with A-D-A structure. Chen et al. reported an A-D-A small molecule comprised of a 3-ethylrodanine end group, yielding a certified PCE of 7.6 % [21]. More recently, using a 2D conjugated small molecule (SMPV1), the best PCE of OSCs reached 8 % and 10 % for single junction and double tandem OSCs [22], indicating the promising potential of BHJ organic solar cells based on these materials at low cost.

In this work, we have synthesized a novel A-D-D’-D-A small molecule denoted as BDT(CDTRH)$_2$ by employing BDT as core donor (D’) unit and 3-ethylrhodanine (RH) as electron withdrawing end groups linked through cyclopentadithiophene (CDT) (D) π spacer have investigated its photophysical and electrochemical properties. CDT has been used as donor units in CDT-benzothiazole oligomer small molecules and showed PCE of 1.61 % when blended with fullerene derivatives [23]. We have used this small molecule as electron donor along with PC$_{71}$BM as electron acceptor for the fabrication of solution processed BHJ organic solar cells. The optimized device with a blend of BDT(CDTRH)$_2$:PC$_{71}$BM (1:1) processed with chloroform (CF) achieved a PCE of 4.58 % with $J_{sc}$ = 8.66 mA/cm$^2$, $V_{oc}$ = 0.98 V and FF= 0.54. The high value of $V_{oc}$ has been attributed to the deeper HOMO level of BDT(CDTRH)$_2$. The PCE was further improved to 6.07 % when small amount of CN (3% v/v) was added to the blend solution. The enhancement of PCE upon using solvent additives may be attributed to an increase in crystallinity of the active layer film, enhancement in the absorption profile and favorable domain sizes of the donor and electron molecules, leading to improved charge transport.

**Experimental part**

**General**

Unless stated otherwise, all reagents were purchased from commercial sources and used without further purification. Dry CHCl$_3$ and toluene were obtained by passing them through an activated alumina column on a PureSolv™ solvent purification system (Innovative Technologies, Inc., MA). Flash column chromatography was carried out using Silica gel 60, 40-63 μm (Panreac Química SLU) as the stationary phase. Size exclusion chromatography was carried out in a large elution column (1000mm x 38 mm) with Biobead SX3 (Bio-Rad
Laboratories, Inc.) as the stationary phase. The eluent was passed through the column under gravity.

**Instruments**

UV-Visible absorption spectra were measured with a 1 cm path-length quartz cell using a Shimadzu model 1700 spectrophotometer. Steady state fluorescence spectra were recorded on a Spex model Fluoromax-3 spectrofluorometer using a 1 cm quartz cell. All $^1$H and $^{13}$C NMR spectra were recorded on a Brüker AV 300 and AV 400 instruments respectively, at a constant temperature of 300 K, unless stated otherwise. Chemical shifts are reported in parts per million and referenced to residual solvent. Coupling constants ($J$) are reported in hertz (Hz). Standard abbreviations indicating multiplicity were used as follows: m = multiplet, quint. = quintet, q = quartet, t = triplet, d = doublet, s = singlet, br = broad. MALDI TOF mass spectra were recorded on a Water Quattro micro (Water Inc, USA) apparatus. Cyclic voltammetric spectra were recorded with a PC-controlled CH instruments model CHI620C electrochemical analyzer.

**Synthesis and characterization**

The relevant synthetic route is outlined in Scheme 1. 6-bromo-4,4-dihexyl-4H-cyclopenta [2,1-b:3,4-b’] dithiophene-2-carbaldehyde (1) [24] and 2,6-bis (trimethyltin)-4,8-bis(2-ethylhexoxy) benzo-[1,2-b:4,5-b’]dithiophene (2) [25] were prepared according to literature procedures. BDT-CPDT-CHO (3) was synthesized Stille coupling between 1 and 2 in refluxing toluene under Argon in the presence of Pd(PPh$_3$)$_4$, as a catalyst for 48 hours. The target compound BDT(CDTRH)$_2$ was obtained using Knoevenagel condensation with 3-ethylrhodanine. To obtain the best possible performance in OSC devices BDT(CDTRH)$_2$ was purified by consecutive flash chromatography, size exclusion chromatography and recrystallization methods. This combination of techniques was found effective in removing most impurities, however one by-product remained in the final product as seen in the $^1$H NMR spectrum of BDT(CDTRH)$_2$. The corresponding NMR peaks could not be assigned to an exact structure, however they are likely to correspond to a benzodithiophene derivative originating from precursor 2. It was estimated that the amount of by product in the final product was inferior to 5% by NMR. BDT(CDTRH)$_2$ was found to be soluble in common organic solvents.

**Synthetic Procedures**

**Synthesis of BDT-CPDT-CHO (3):**
A solution of 6-bromo-4,4-dihexyl-4H-cyclopenta [2,1-b:3,4-b’] dithiophene-2-carbaldehyde (i) (300 mg, 0.387 mmol) and 2,6-bis(trimethyltin)-4,8-bis(2-ethylhexoxy)benzo-[1,2-b:4,5-b’]dithiophene (ii) (380 mg, 0.852 mmol) in dry toluene (20 mL) was degassed performing two freeze/thaw cycles under Argon. To this solution was added Pd(PPh3)4 (44 mg, 0.04 mmol). After being stirred at 120 °C for 48 h, the reaction mixture was poured into water and extracted with CH2Cl2. The organic layer was washed with water and then dried over MgSO4. The solvent was removed under reduced pressure and the crude product was purified by column chromatography on silica gel using a mixture of CH2Cl2 and hexane (1:1) as eluent to afford compound BDT-CPDT-CHO (iii) (290 mg, 39%).

1H-NMR (300 MHz, CDCl3) (Figure S1 and S2 of supplementary information) δH: 9.86 (S, 2H), 7.59 (S, 2H), 7.56 (S, 2H), 7.19 (S, 2H), 4.23 (d, J=5.4 Hz) 1.92 (m, 2H), 1.69 (m, 8H), 1.46 (m, 4H), 1.26 (m, 44H), 0.81-0.90 (m, 24H).

13CNMR (100MHz, CDCl3) δ: 182.23; 162.73; 157.92; 149.69 147.37; 146.81; 143.87; 143.64; 141.64; 136.53; 135.46; 133.58; 132.26; 129.78; 128.85; 119.29; 115.92; 54.05; 40.47; 38.49; 33.75; 31.34; 30.24; 29.47; 29.39; 29.02; 28.06; 24.39; 24.23; 23.53; 22.96; 22.72; 22.37; 14.00; 11.15. MALDI: m/z calcld for C70H94O6S6 1190.55, found 1190.7 (Figure S3 of supplementary information)

**Synthesis of BDT(CDTRH)2:**

BDT-CPDT-CHO (iii) (200 mg, 0.167 mmol) was dissolved in a solution of dry CHCl3 (40 mL), a few drops of piperidine and then 3-ethylrhodanine (268 mg, 1.67 mmol) were added, and the resulting solution was refluxed and stirred for 24 h under argon. The reaction mixture was then extracted with CH2Cl2, washed with water, and dried over MgSO4. The solvent was removed under reduced pressure and the crude product was purified by column chromatography on silica gel using CH2Cl2 and hexane (1:1) as eluent and subsequent size exclusion chromatography using THF as eluent. The solid was further purified by two consecutive recrystallizations carried out as follows: The solid was dissolved in 15 mL of CHCl3, and poured by syringe through a 0.2 um PTFE filter in a 250 mL graduated cylinder. A buffer layer (10 mL) consisting of a 1:1 mixture of CHCl3:Hexane was carefully poured over the solution (filtered through 0.2um PTFE filter). Finally, pure hexane was carefully poured over the buffer layer (filtered through 0.2um PTFE filter) up to the top of the graduated cylinder and the solution was left in the dark undisturbed. BDT(CDTRH)2 precipitated as dark blue shiny crystals over a period of 2 to 3 weeks approximately, after which period they were recovered by filtration and washed with plenty of hexane (200 mL) (120 mg, 60%). 1H-NMR (300 MHz, CDCl3) (Figure S4 and S5 of supplementary information) δH: 7.84 (S, 2H), 7.45 (S, 2H), 7.21 (S, 2H), 7.14 (S, 2H), 4.19 (m, 8H), 1.90 (m,
2H), 1.61(m, 8H), 1.43(m, 4H), 1.16 (m, 50H), 0.80-
0.91 (m, 24H), 13C-NMR (100 MHz, CDCl3) δ: 191.60; 167.22; 161.87; 159.50; 145.94; 143.95; 141.95; 139.06; 136.74; 136.015; 132.47; 128.90; 126.22; 119.43; 115.89; 113.45; 54.36; 40.74; 39.96; 37.80; 31.60; 30.52; 29.62; 29.29; 24.70; 23.90; 23.21; 22.61; 14.25; 14.00; 12.28; 11.43. MALDI (Figure S6 of supplementary information): m/z calcd for C80H104N2O4S10 1476.52, found 1476.5196

**Device fabrication and characterization**

The BHJ organic solar cells were fabricated using the glass /ITO /PEDOT:PSS/ BDT(CDTRH)2:PC71BM/Al device structure. The ITO coated glass substrates were cleaned ultrasonically consecutively in aqueous detergent, de-ionized water, isopropyl alcohol and acetone and finally dried under ambient conditions. An aqueous solution of PEDOT:PSS (Heraeus, Clevios P VP, Al 4083) was spin cast on the ITO substrates to obtain film having a thickness of about 40 nm. The PEDOT:PSS film was then dried for 10 min at a temperature of 120° C in ambient conditions. Mixtures of BDT(CDTRH)2 with PC71BM with weight ratios of 1:0.5, 1:1, and 1:2 and 1:2.5 in chloroform (CF) were prepared and then spin-cast onto the PEDOT:PSS layer and dried overnight at ambient atmosphere. The devices were further fabricated by the use of 1 %, 2 %, 3 % and 4% CN processing additive during the spin casting step (spinning speed was about 2000 rpm). The concentration of the BDT(CDTRH)2:PC71BM blend was 20 mg/mL. The thickness of the active layer is in between 85-90 nm. Finally, the aluminium (Al) top electrode was thermally deposited on the active layer at a vacuum of 10⁻⁵ Torr through a shadow mask of area of 0.20 cm². All devices were fabricated and tested in ambient atmosphere without encapsulation. The hole-only and electron-only devices with ITO/ PEDOT:PSS/ BDT(CDTRH)2 :PC71BM/ Au and ITO/ Al/ BDT(CDTRH)2:PC71BM/ Al architectures were also fabricated in an analogous way, in order to measure the hole and electron mobility, respectively.

The current-voltage characteristics of the BHJ organic solar cells were measured using a computer controlled Keithley 238 source meter under simulated AM1.5G illumination of 100 mW/cm². A Xenon light source coupled with optical filter was used to give the stimulated irradiance at the surface of the devices. The incident photon to current efficiency (IPCE) of the devices was measured illuminating the device through the light source and monochromator and the resulting current was measured using a Keithley electrometer under short circuit condition.

**Results and discussion**
Photophysical and electrochemical properties

Figure 1 shows the UV-Visible absorption spectra of BDT(CDTRH)$_2$ in a chloroform solution and a thin film spin coated on a quartz substrate. In solution BDT(CDTRH)$_2$ exhibits an absorption peak around 582 nm with a molar extinction coefficient of $7.8 \times 10^4$ M$^{-1}$ cm$^{-1}$. However, the BDT(CDTRH)$_2$ film shows a broader absorption band covering from 400 to 800 nm with an absorption peak around 618 nm and also exhibits a vibronic shoulder at 662 nm, indicating an effective $\pi-\pi$ packing between the molecule backbones in solid state [26]. The optical bandgap of BDT(CDTRH)$_2$ was estimated from the onset absorption edge in its absorption spectra in thin film, and is about 1.72 eV. The low bandgap of BDT(CDTRH)$_2$ is attributed to the ethylrhodanine acceptor moiety [27]. The absorption spectra of BDT(CDTRH)$_2$ in thin film exhibits a broader absorption profile from 550 nm to 720 nm and low bandgap which increases light harvesting when used as donor in the active layer in organic solar cells. Cyclic voltammetry (CV) was used to estimate approximate HOMO and LUMO energy levels of BDT(CDTRH)$_2$. The HOMO and LUMO energy levels of the BDT(CDTRH)$_2$, calculated from the onset oxidation and reduction potential respectively, are -5.38 eV and -3.54 eV, respectively. The electrochemical bandgap of BDT(CDTRH)$_2$ is 1.80 eV. The optical and electrochemical band gap of BDT(CDTRH)$_2$ is similar to other small molecules reported in literature [20]. Considering that the PC$_{71}$BM acceptor has HOMO and LUMO levels of $-6.1$ eV and $-4.0$ eV, respectively, the LUMO energy difference ($>0.3$ eV) between the donor and acceptor is large enough for the separation of excitons [28], which is beneficial for increasing the short circuit current in BDT(CDTRH)$_2$ based organic solar cells.

Theoretical Calculations

We have additionally performed a theoretical study on the BDT(CDTRH)$_2$ molecular structure within the framework of density functional theory (DFT) and time-dependent density functional theory (TD-DFT). The initial geometry optimization calculations were performed employing the gradient corrected functional PBE [29] of Perdew, Burke and Ernzerhof. The triple-$\zeta$ quality TZVP basis set [30], was used for all of the calculations. At this stage of the calculations, to increase the computational efficiency (without loss in accuracy), the resolution of the identity method [31] was used for the treatment of the two-electron integrals. Subsequent geometry optimization were further performed using the hybrid exchange–correlation functional B3LYP [32] as well as Truhlar’s meta-hybrid exchange–correlation functional M06 [33], and the same basis set. Tight
convergence criteria were placed for the SCF energy (up to $10^{-7}$ Eh) and the one-electron density (rms of the density matrix up to $10^{-8}$) as well as for the norm of the Cartesian gradient (residual forces both average and maximum smaller than $1.5 \times 10^{-5}$ a.u.) and residual displacements (both average and maximum smaller than $6 \times 10^{-5}$ a.u.). Solvent effects were included for chloroform (CF) using the integral equation formalism variant of the Polarizable Continuum Model (IEFPCM), as implemented in the Gaussian package [34]. The first round of geometry optimization was performed using the Turbomole package [35]. All of the follow up calculations were performed using the Gaussian package [34].

The TD-DFT excited state calculations were performed to calculate the optical gap of BDT(CDTRH)$_2$ employing the same functionals and basis set on the corresponding ground state structures. The UV/Vis spectra were calculated using the B3LYP and M06 (that provides leveled performance over transition types [36, 37]) functionals.

For the geometry optimization of the BDT(CDTRH)$_2$ molecular structure several rotamers were examined as initial geometries, including aliphatic configurations (details are given in the supplementary information). Vibrational analysis revealed that the final optimized structure is a true local (if not global) minimum; none of the vibrational modes had imaginary eigen frequencies. The BDT(CDTRH)$_2$ structure is for the most part planar, with a tilting of the outer groups relative to the central benzodithiophene group by dihedral angles in the range of $17^\circ$–$27^\circ$ (depending on the functional used and the presence of solvent). We have calculated the HOMO and LUMO energy levels and the optical gaps, defined here as the energetically lowest allowed vertical electronic excitation, employing the PBE, M06, and B3LYP functionals. In Table 1, in addition to the frontier orbitals’ energy levels, we also provide the optical gap the main contributions to the first excitation as well as the wavelength of the first excitation and of the excitations with the largest oscillator strengths. In Table 1 we also provide the character of the first allowed excitations only for contributions larger than 4%. The first excitation, as calculated by each of the functionals, clearly exhibits a strong single-configuration character, with only marginal (if any) to moderate secondary contributions.

In Figure 2 we have plotted the isosurfaces (isovalue = 0.02) of the HOMO and LUMO, as well as the next nearest frontier orbitals, of the BDT(CDTRH)$_2$ structure. Both the HOMO and LUMO extend over the main, nearly planar, body of the structure. There is however a varying level of delocalization; the HOMO is more localized near the central part of the structure while the LUMO is more localized towards the outer (edge) parts. The HOMO–1 and LUMO+1 do not exhibit significant contributions from the central structure,
i.e. the benzodithiophene (BDT) moieties, but are rather localized mainly towards the outer parts, i.e. the cyclopentadithiophene (CPDT) and the Thiazole (TA) groups. To quantify the contributions of the moieties to the frontier orbitals we have calculated the total and partial density of states (PDOS). The PDOS for BDT(CDTRH)₂ is shown in Figure S7 of supplementary information. We partition BDT(CDTRH)₂ into the BDT, CPDT, TA and the aliphatic groups. The contribution of the BDT, CPDT, and TA groups to the HOMO is 39.0%, 45.3%, and 14.9% respectively, with only an 8.3% combined contribution from all the aliphatic groups. The corresponding contributions to the LUMO are 12.3%, 41.1% and 44.4%, respectively, and 2.2% combined from all the aliphatic groups. This is in agreement with our earlier observation on the localization of HOMO on the inner structure and the LUMO mainly on the outer structure. Taking into account that the first excitation is dominated by a HOMO to LUMO transition this indicates at least a moderate (due to overlaps) charge transfer from the corresponding inner to outer parts of the structure. The first significant contributions (16.2%) from the aliphatic groups are noted at lower energies, around −8.0 eV, which corresponds to the HOMO−15 level.

In Figure 3a we show the UV/Visual absorption spectra of the BDT(CDTRH)₂ structure calculated at the TD-DFT/M06 level of theory, both accounting for solvent effects for CF and in gas phase. The spectra have been produced by convoluting Gaussian functions with HWHM = 0.25 eV centered at the excitation wave numbers. In Figure 3b, we also provide the corresponding spectra calculated using the B3LYP functional, which is in very good agreement with both that using M06 and the experimental spectra. A characteristic absorption spectrum in found that displays a main band centered at 618nm and three lower intensity bands at smaller wavelengths, specifically centered at 462nm, 376nm, and 307nm. The calculated spectrum is in good agreement to the experimental one, with slightly overestimated wave numbers by about 5–35 nm, depending on the region (more overestimated at larger wavelengths). The wavelengths of the excitations with the largest oscillator strengths within these bands are given in Table 1.

**Photovoltaic properties**

In BHJ organic solar cells, the relative amounts of the donor and acceptor materials used in blend active layer is of great importance for their photovoltaic performance, since there should be a balance between the absorption and charge transporting network in the active layer. When the acceptor content is too low, the electron transporting ability will be limited, while when the acceptor content is too high, the absorbance and hole transport ability
in active layer will be decreased. BHJ devices were fabricated and optimized using different:

\[ \text{BDT(CDTRH)}_2 : \text{PC}_{71} \text{BM} \] ratios with device configuration being ITO/PEDOT:PSS/\n\[ \text{BDT(CDTRH)}_2 : \text{PC}_{71} \text{BM}/\text{Al} \] using conventional solution spin coating process. Various
ratios of \[ \text{BDT(CDTRH)}_2 : \text{PC}_{71} \text{BM} \] were investigated and we found that the optimum weight ratio
was 1:1. The normalized absorption spectra of \[ \text{BDT(CDTRH)}_2 : \text{PC}_{71} \text{BM} \] blend thin film is shown Figure 4. It can be seen from this figure that the absorption spectra of blend is the combination both \[ \text{BDT(CDTRH)}_2 \] and \[ \text{PC}_{71} \text{BM} \] absorption features indicating that both
components participate into exciton generation and photocurrent of the device.

The current-voltage \( J-V \) characteristics, under sun-stimulated illumination (AM1.5, 100 mW/cm\(^2\)) of the BHJ organic solar cell with the \[ \text{BDT(CDTRH)}_2 : \text{PC}_{71} \text{BM} \] blend of 1:1 weight ratio processed from chloroform (CF) is displayed in Figure 5a and the corresponding photovoltaic parameters are compiled in Table 2. The device shows an overall PCE of 4.58 % with \( J_{SC} =8.66 \text{ mA/cm}^2 \), \( V_{OC} = 0.98 \text{ V} \) and FF= 0.54. The high value of \( V_{OC} \) is attributed to the
deeper HOMO level of \( V_{OC} \), since the maximum theoretical \( V_{OC} \) of the BHJ organic solar cells
is directly related to the energy difference between the HOMO of donor and LUMO of
acceptor materials used in active layer.

To further improve the performance of these devices we used CN as the solvent
additive and varied the volume concentration in the blend solution from 1 to 4 % and found
that 3% v/v gave the best photovoltaic performance. The \( J-V \) characteristics of the best
performing device are shown in Figure 5a and photovoltaic parameters are summarized in
Table 2. It achieved and overall PCE of 6.07 % with \( J_{SC} \) of 10.42 mA/cm\(^2\), \( V_{OC} = 0.94 \text{ V} \) and
FF = 0.62.

The improvement in the overall PCE of the device processed with CN/CF solvent was
mainly due to a significant increase in the \( J_{SC} \) as well as the FF. The \( J_{SC} \) is directly related to the
light absorption ability of the active layer as well as better collection of charges at the
electrodes, which is in direct relationship with IPCE response as well as the film morphology.
The \[ \text{BDT(CDTRH)}_2 : \text{PC}_{71} \text{BM} \] film cast from the CN/CF displays stronger absorption than
the film cast from CF (Figure 5). As shown in the absorption spectra of the blend cast from
the CN/CF solvent the absorption band of \[ \text{BDT(CDTRH)}_2 \] shifts slightly toward longer
wavelength and broadens. The absorption coefficient also increases. The vibronic shoulder peak is clearly observed at around 668 nm. The observed red shift in the absorption band of
\[ \text{BDT(CDTRH)}_2 \] is attributed to the increase in order in the active layer film. The higher
degree of crystallinity as indicated by the red shift and vibronic shoulder in the absorption
spectra of the blend, also attributes to the enhanced \( \pi-\pi^* \) conjugation. The increase in
crystallinity of the active layer has been confirmed by XRD measurements and discussed later. The enhanced value of FF may be ascribed to high crystallinity and high carrier mobility of BDT(CDTRH)$_2$ in the blend film [38].

The IPCE spectra of the devices processed with CF and CN/CF are shown in Figure 6b. The IPCE curves of the device covered a broad wavelength range from 360 nm to 720 nm. The shape of IPCE spectra of the devices closely resembles the optical absorption spectra of the active layers (Figure 4). Moreover, the IPCE spectrum of BDT(CDTRH)$_2$:PC$_{71}$BM solar cell processed with CN/CF was over 64 % at 622 nm, whereas that of the device based on CF processed was about 56 % (Figure 5b). Additionally, the values of IPCE of the device processed with CF/CN at all absorption wavelengths were higher leading to the enhanced value of $J_{SC}$.

The surface morphology of the BDT(CDTRH)$_2$:PC$_{71}$BM blend films processed from CF and CN/CF were investigated by atomic force microscopy (AFM). Figure 6 shows the topographic images of the BDT(CDTRH)$_2$:PC$_{71}$BM active layers processed with CF and CN/CF solvents. The film cast from CF shows apparent homogeneously mixed domains of BDT(CDTRH)$_2$ and PC$_{71}$BM with a surface root mean square (RMS) roughness of 2.45 nm. The incorporation of 3 % v/v CN in the blend solution results in a significant change in the surface morphology of the film with the RMS roughness of the film decreasing to 1.24 nm. Although the AFM images show that the morphology of the active layer has improved with the incorporation of CN it is difficult to conclude that these features originate from the crystallization of the donor phase. Therefore we have further investigated the domain sizes of the active layer films processed with CF and CN/CF solvents using TEM. In the TEM images (Figure 7) well interpenetrating structures are formed with domain sizes 30-40 nm and 15-20 nm for the active layer processed with CF and CN/CF solvents, respectively.

Figure 8 shows the XRD pattern of the pristine BDT(CDTRH)$_2$ and BDT(CDTRH)$_2$:PC$_{71}$BM thin films cast from CF and CN/CF solvents. In the case of pristine BDT(CDTRH)$_2$ cast from both CF and CN/CF solvents, a band centered at 20=5.64° was observed in the XRD pattern. However, the intensity of the band is more pronounced for the film cast from CN/CF, indicating an increase in crystallinity. It can be seen from the Figure that when the BDT(CDTRH)$_2$ :PC$_{71}$BM blend is cast from CF, the diffraction peak becomes wider and the intensity decreased. However when the blend is cast from CN/CF solvent, the intensity of the diffraction peak observed at 20=5.64°, increases. Since the boiling point of CN is higher than CF, the film cast from the CN/CF solvent evaporates slowly and results in
an increase in crystallinity. This change in crystallinity of the films agrees with the observed change in the absorption spectra. The increase in crystallinity of the blend helps to increase the hole mobility in the blend.

The charge carrier mobility in the BHJ active layer is critical for the BHJ organic solar cell. This is mainly due to the fact that after the exciton dissociation at the D/A interface the amount of photogenerated charge carriers extracted at the electrode depends on the ratio of the charge carriers swept and those lost by recombination during their transport to the electrodes [3e, 39]. In order to calculate the hole and electron mobility in BDT(CDTRH)$_2$:PC$_{71}$BM active layer, we measured the dark current of hole only and electron only devices, respectively and then analyzed the results using the space charge limited current (SCLC) model [40]:

$$J = \frac{9}{8} \varepsilon_r \varepsilon_0 \mu_h \left( \frac{V_{\text{appl}} - V_{\text{bi}}}{L^3} \right)^2$$

where $\varepsilon_r, \varepsilon_0$ is the dielectric permittivity of the active layer and permittivity of free space, respectively, $V_{\text{appl}}$ is applied voltage and $V_{\text{bi}}$ is built in potential, $\mu_h$ is the hole mobility, $L$ is the thickness of the active layer, and $\beta$ is the field activation factor. Figure 9 shows the experimental $J$-$V$ characteristics of hole only device with configuration ITO/PEDOT:PSS/ BDT(CDTRH)$_2$:PC$_{71}$BM/Au. The hole mobilities were estimated from fitting the experimental results (solid lines in Figure 10) and are about 2.3 x10$^{-5}$ cm$^2$/Vs and 6.4 x10$^{-5}$ cm$^2$/Vs, respectively for active layer processed with DIO/CF and CF. The increase in hole mobility with CN additive may be attributed to the increase in crystallinity and domain size. The electron mobilities were estimated from the $J$-$V$ characteristics of the electron only devices ITO/Al/ BDT(CDTRH)$_2$:PC$_{71}$BM/Al and are almost identical i.e. 2.52 x10$^{-4}$ cm$^2$/Vs and 2.48 x10$^{-4}$ cm$^2$/Vs, with and without CN additive, respectively. The difference between electron and hole mobilities is reduced in the BDT(CDTRH)$_2$:PC$_{71}$BM film processed with DIO/CF leading to a more balanced charge transport [41].

**Conclusion**

In summary, we have synthesized an A-D-D'-D-A small molecule, BDT(CDTRH)$_2$, with a low bandgap (~1.72 eV) and investigated its optical and electrochemical properties (both theoretically and experimentally). The photophysical and electrochemical investigations of BDT(CDTRH)$_2$ and PC$_{71}$BM films proves that the BDT(CDTRH)$_2$:PC$_{71}$BM blend can effectively harvest photons from visible to near infrared region of solar spectrum and transfer the electrons to PC$_{71}$BM molecules, resulting in a
photovoltaic effect. The solution processed BHJ solar cell based on BDT(CDTRH)$_2$:PC$_{71}$BM processed from CF displayed a PCE of 4.58%. In order to improve the PCE, the BDT(CDTRH)$_2$:PC$_{71}$BM layer was spin cast from a mixture of 3% v/v DIO/CF and the device showed a PCE of 6.07%, attributed to the enhancement in the $J_{SC}$ and FF. This is a result of the enhanced IPCE response, increase in the crystalline nature and more favourable nanoscale morphology of active layer with solvent additive and increase in hole mobility, which leads to more balanced charge transport, and enhanced hole mobility in the BHJ active layer.

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References


35. TURBOMOLE (version 5.6); Universitat Karlsruhe, 2000


Table 1 Calculated properties of BDT(CDTRH)$_2$. Specifically HOMO and LUMO energies (eV), HOMO–LUMO gap (eV), $HL$, Optical gap (eV), $OG$, with corresponding oscillator strengths, $f$, the wavelengths of the first excitation and excitations with the largest oscillator strengths, the main contributions to the first excited state, and the dipole moment (D), $\mu$

<table>
<thead>
<tr>
<th>Model</th>
<th>HOMO (eV)</th>
<th>LUMO (eV)</th>
<th>HL (eV)</th>
<th>OG (eV)</th>
<th>$\lambda_{\text{max}}$ (nm)</th>
<th>$f$</th>
<th>Main contributions</th>
<th>$\mu$ (D)</th>
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<tr>
<td>PBE</td>
<td>-4.60</td>
<td>-3.52</td>
<td>1.08</td>
<td>1.57</td>
<td>938</td>
<td>1.572</td>
<td>H→L (92.2%), H-1→L+1 (4.8%)</td>
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<tr>
<td>B3LYP</td>
<td>-5.24</td>
<td>-3.07</td>
<td>2.17</td>
<td>1.93</td>
<td>641, 492, 390, 337, 684, 509, 398, 331</td>
<td>2.558</td>
<td>H→L (98.3 %), H→L (97.2)</td>
<td>1.215</td>
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<tr>
<td></td>
<td>-5.21$^a$</td>
<td>-3.11$^a$</td>
<td>2.10$^a$</td>
<td>1.81$^a$</td>
<td>641, 492, 390, 337, 684, 509, 398, 331</td>
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<td>H→L (98.3 %), H→L (97.2)</td>
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<td>2.12</td>
<td>585, 454, 368, 318, 619, 468, 375, 302</td>
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<td>H→L (90.0%), H-1→L+1 (5.8%), H→L (88.3 %), H-1→L+1 (7.9%)</td>
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<td>-5.47$^a$</td>
<td>-2.92$^a$</td>
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<td>2.00$^a$</td>
<td>585, 454, 368, 318, 619, 468, 375, 302</td>
<td>3.356$^a$</td>
<td>H→L (90.0%), H-1→L+1 (5.8%), H→L (88.3 %), H-1→L+1 (7.9%)</td>
<td>1.498$^a$</td>
</tr>
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$^a$ Values when solvent effects are taken into account for chloroform

Table 2 Photovoltaic parameters of the BDT(CDTRH)$_2$:PC$_{71}$BM (1:1) based solar cells

<table>
<thead>
<tr>
<th>Solvent</th>
<th>$J_{sc}$ (mA/cm$^2$)</th>
<th>$V_{oc}$ (V)</th>
<th>FF</th>
<th>PCE (%)</th>
</tr>
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<tr>
<td>CF cast</td>
<td>8.66</td>
<td>0.98</td>
<td>0.54</td>
<td>4.58</td>
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<tr>
<td>CN/CF cast</td>
<td>10.42</td>
<td>0.94</td>
<td>0.62</td>
<td>6.07</td>
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</table>
Scheme 1 Synthetic route of BDT(CDTRH)$_2$. (Reaction conditions): (i) Pd(PPh$_3$)$_4$, Toluene, 48 h, Reflux (ii) dry CHCl$_3$, piperidine, 12 h, Reflux
Figure 2 Absorption spectra of BDT(CDTRH)$_2$ in dilute solution of CF and thin film cast from CF

Figure 2 Frontier and near frontier orbitals of BDT(CDTRH)$_2$
Figure 3 Theoretical UV-Vis absorption spectrum of BDT(CDTRH)$_2$ (a) calculated using the M06 functional and (b) B3LYP functional.
Figure 4 Normalized absorption spectra of BDT(CDTRH)$_2$:PC$_{71}$BM (1:1) films cast from CF and 3% v CN/CF solvents.
Figure 5 (a) Current–voltage (J-V) characteristics under illumination and (b) IPCE spectra of solar cells based on BDT(CDTRH)$_2$:PC$_{71}$BM (1:1) blend cast from CF and 3% v CN/CF solvents
Figure 6  AFM tapping mode height images (5 μm x5 μm size) of \( \text{BDT(CDTRH)2:PC}_{71}\text{BM} \) blend thin films cast from CF and 3 % v CN/CF

Figure 7  TEM images of \( \text{VC90:PC}_{71}\text{BM} \) blend thin films cast from CF and 3 % v CN/CF, Scale bar is 200 nm
Figure 8 XRD patterns of (a) pristine $\text{BDT(CDTRH)}_2$ and (b) $\text{BDT(CDTRH)}_2: \text{PC}_{71}\text{BM}$ blend thin films cast from CF and 3 % v CN/CF
Figure 9 Current-voltage (J−V) characteristics in dark for hole-only devices, using CF and 3 % v CN/CF processed BDT(CDTRH)$_2$:PC$_{71}$BM active layer blends