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Covalently linked benzothiadiazole-fullerene adducts for organic optoelectronic devices: synthesis and characterization†

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Fullerene adducts have attracted attention in a variety of applications including organic optoelectronic devices. In this regard, we have designed a covalently linked donor–acceptor dyad comprising a fluorobenzothiadiazole-thiophene (BTF₂-Th) unit with the electron acceptor fullerene in an Acceptor–Donor–Acceptor (A–D–A) molecular arrangement. We synthesized and characterized two new covalently bonded benzothiadiazole-based fullerene molecules, mono-adduct, **7** (benzothiadiazole : PC₆₁BM = 1 : 1, anchored terminally via esterification reaction) and multi-adduct, **10-I** (benzothiadiazole : PC₆₁BM = $n \geq 1$, attached directly to the fullerene core via the Prato reaction) using different synthetic strategies. A broadening of the UV-visible spectra of the modified fullerene derivative with strong absorption from 350 to 500 nm and at low wavelengths is observed as compared to PC₆₁BM. A suitable bandgap, good electronic conductivity, and appreciable solubility in solvents suggest their utility in optoelectronic devices. The mono-adduct **7** showed two-order higher electron mobility as compared to bis-adduct **10-I** due to retention of extended conjugation in fullerene, as in the case of PC₆₁BM. Experimentally determined optical properties and energy levels of the fullerene adducts were found to be in good agreement and supported by theoretical calculations. The presence of BTF₂ affects the ground state dipole moments as well as the absorption strengths, most noticeable in the case of two attached BTF₂ moieties. The HOMO and LUMO levels are found to be localized on the fullerene cage with the extension of the HOMO to the BTF₂ unit more and the same is noticed in ground state dipole moment in the side-chain functionalized structure. Such structural arrangement provides easy charge transfer between acceptor and donor units to allow a concomitant effect of favorable optoelectronic properties, energy levels of the frontier orbitals, effective exciton dissociation, and charge transport which may reduce processing complexity to advance single material-based future optoelectronic devices.

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Introduction

With the world moving towards sustainable energy organic solar cells (OSCs) have become very popular and a promising method of harvesting solar energy in an affordable, scalable and

efficient manner.¹ OSCs utilize either polymer or small molecule organic semiconductors owing to their unique advantages, such as ease of device fabrication by efficient solution-based roll-to-roll printing, light-weight, and mechanical flexibility. Moreover, designing materials at the molecular level allows bandgap tunability, and high optical absorptivity which is desired to achieve high power conversion efficiency (PCE) of the devices. One approach to improve PCE involves optimization of the energy gap between the highest occupied molecular orbital (HOMO) of the donor and the lowest unoccupied molecular orbital (LUMO) of the acceptor to allow maximum energy conversion and to affect open-circuit voltage (V_{oc}). The second approach demands the exploration of strategies to maximize the intermolecular interactions between the donor and acceptor units.^{1c,2}

Bulk heterojunction (BHJ) photovoltaic devices revealed high PCE and are vastly based on the blending of a semiconducting polymer/organic small molecule donor with a soluble fullerene derivative as an acceptor, to improve the charge-carrier mobility

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and exciton separation.³ The fullerene derivatives constitute so far as one of the most explored class of acceptors for BHJ. The parent fullerenes C_{60/70} rings are tethered usually with organic functionalities to improve the solubility and processability. Such acceptors have vastly improved the device efficacy due to their advantageous properties such as (i) the strong electron affinity to accept and transport electrons in three dimensions, (ii) high electron mobilities, (iii) multiple reversible electrochemical reductions, and (iv) the ability to aggregate in bulk heterojunctions to form both pure and mixed domains of the appropriate length scale to enable charge separation.⁴ Among the fullerene derivatives, [6]-phenyl-C₆₁ butyric acid methyl ester (PC₆₁BM) acceptor is widely used due to its good solubility, well structurally characterized, further structurally tailored and revealed appreciable compatibility with many conjugated donors.⁵ Padinger *et al.*⁶ reported BHJ device based on a blend of poly(3-hexylthiophene) donor and PC₆₁BM, P3HT:PC₆₁BM, with a PCE of >3.5%, and it has been used as the reference cell in this field.^{1c,3b,7} Fullerene derivatives have been widely used in optoelectronic device applications. Recently, Karunakaran *et al.* reported D-A-D type molecules with PCBM as acceptor material for bulk heterojunction-based optoelectronic devices.⁸ Anthopoulos and others demonstrated PC₆₁BM based ultraviolet phototransistors with balanced electron and hole transport characteristics.⁹ Field-effect transistors were also reported by Dong *et al.* using fullerene derivatives having ethylene glycol side chains with different lengths.¹⁰ Self-assembled monolayer based on fullerene were also investigated by Lee *et al.*¹¹ that improved electron transfer at the organic-ZnO interface, without impacting the morphology of active layer. Higher fullerene analogues, like PC₇₁BM, have also been studied extensively as acceptors in OSCs, and recently, Hu and co-workers reported a record 15.3% PCE for a PC₇₁BM based-all small molecule OSCs.¹² Functionalization of fullerene raises the LUMO level¹³ and thus reduces electrochemical losses and simultaneous increase in V_{oc}.¹⁴ Multi-adducts of fullerene, more specifically, the bis-adducts have shown better performance than their mono-adducts.¹⁵ In general, the higher-adducts than bis-adducts of fullerene show a lower light-harvesting to power conversion performance due to decrease in electron mobility, the occurrence of electronic traps, and a minimal effect on LUMO level.^{15b,16} Recently multiple functionalized fullerene adducts gained interest for advanced materials and biological applications and thus interesting for exploration.¹⁷

Another electron-deficient unit of growing interest is 2,1,3-benzothiadiazole (BT), which exhibits low-lying energy levels that make it an important acceptor building block in materials design to modulate electron transport properties.¹⁸ An ease to introduce various functionalities in the benzothiadiazole unit offers rich structural designing flexibility to tune the molecule properties. The most acceptable and preferable methods involve the introduction of electron-withdrawing groups on the aryl ring of BT that allows structural modification using a wide range of reactions.¹⁹ Fluorination of BT ring was found to improve the solar cell performance as it lowers the energy of the HOMO level. Moreover, fluorine functionality improves interactions between the donor and acceptor units, thereby

enhancing the diffusive properties of the compound.²⁰ Another way of altering electronic properties of BT derivative is through the fusion of the benzothiadiazole unit with widely explored donor units, like thiophene, which has been reported to generate structures with enhanced light-harvesting efficiency.²¹ Molecules involving fluorobenzothiadiazole (BTF₂) and thiophene (Th) have been studied and the proximity of the two moieties allows tunability of electronic properties. The F···S interaction between the fluorobenzothiadiazole and thiophene induces planarity and such conformation of the molecule in the solid-state favors intermolecular stacking to form an array of units and thus assists better charge transportation.²² Such intra- and intermolecular interactions (*e.g.*, F···H, F···S, and F···π), favor the crystalline morphology at a nanoscale.²³

Use of variation in the arrangement of the acceptor (A)-donor (D) units became popular in light-harvesting materials as they allow better intramolecular charge transfer from the acceptor to the donor units, leading to improved performance of the device.²⁴ Alternating arrangement of units as A-D-A was first introduced by Schulze *et al.*²⁵ and found that the acceptor/donor interface assists the separation of excitons as electrons in the acceptor and holes in the donor. More recently, Chen *et al.*²⁶ reported that A-D-A type acceptors exhibit outstanding performance with a PCE of 17.3% in tandem geometry. Such structural arrangements of the acceptor/donor units result in enhanced charge transfer with a simultaneous broadening of the optical absorption range.²⁷

In the current work, we aim to modify the PC₆₁BM acceptor molecule with a covalently linked BTF₂-Th unit to obtain an A-D-A type arrangement. Three aspects are considered in the tuning of the acceptor molecule: (i) lowering of bandgap, (ii) efficient inter- and intra-molecular electron transport (iii) to improve optical absorptivity of fullerenes. As a design strategy to attain the former objective, fluorine atoms are incorporated in the acceptor molecule while for the latter objective, benzothiadiazole moiety was covalently tethered to PC₆₁BM to allow directionality for efficient electron transport and improve light absorbance characteristics. Fig. 1 shows the structures of two new covalently bonded benzothiadiazole-based fullerene molecules, mono-adduct (benzothiadiazole : PC₆₁BM = 1 : 1) and multi-adduct (benzothiadiazole : PC₆₁BM = n : 1, where n ≥ 1) which were synthesized and characterized. The thiophene π-bridge (D1, Th) linked difluoro-2,1,3-benzothiadiazole (A1, BTF₂) unit is anchored to the fullerene (A2) moiety *via* a phenyl linker (D2) to design acceptor molecules in D1-A1-D2-A2

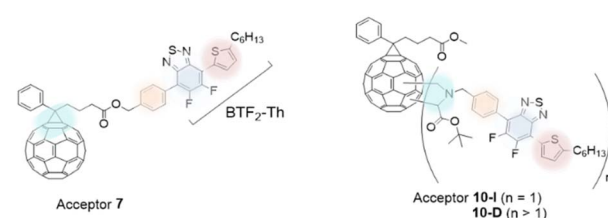


Fig. 1 The mono- and multi-adduct acceptors (7: PC₆₁BTF₂, 10-I: PC₆₁BM-BTF₂ and 10-D: PC₆₁BM-nBTF₂).



configuration. The fullerene moiety was introduced using PC_{61}BM as it is among the most widely used electron acceptor in BHJ devices and offers good solvent solubility and electronic properties.²⁸ The terminal thiophene donor (**D1**) unit contained a C_6 -long alkyl chain which not only assists solvent processability of the final molecule but is also known to affect inter- and intramolecular interactions.^{23a} The structural arrangement of the two-hybrid π -bridge units sandwiching **A1** unit may favor structural planarity to affect molecular interactions. Furthermore, variation in electron-donating power as provided by the difference in aromaticity between BTF_2 and fullerene ring may provide electron conduction gradient to ease the movement of electrons. Backbone modulation of **D1-A1-D2** is achieved by altering the mode of its linkage to the fullerene unit, one at the distal end (Acceptor **7**) and the other attached to the core (Acceptor **10**). The energy levels of the molecules are evaluated theoretically and experimentally.

Results and discussion

Synthesis and characterization of benzothiadiazole derivatives

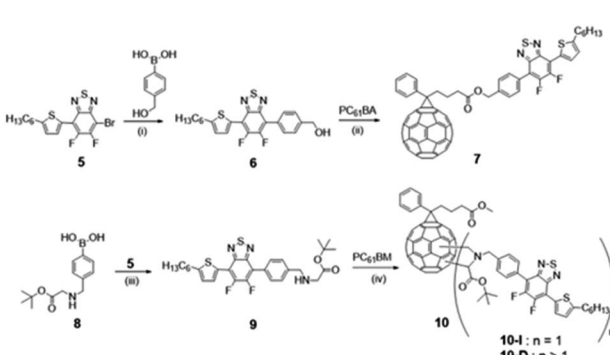
A multi-step synthesis was adopted to obtain the thiophene-benzothiadiazole-fullerene compounds (**7** and **10**) as shown in Scheme 1. The synthesis of compound **5** is shown in Scheme S1 (ESI†). The nitro group of commercially available 4,5-difluoro-2-nitroaniline was reduced to an amino group using *in situ* generated hydrogen by reaction of HCOONH_4 in the presence of Pd/C to form compound **2**. Intramolecular cyclization of **2** with SOCl_2 yielded difluorobenzothiadiazole, **3**. Bromination of **3** with bromine and iron powder afforded dibromo-difluorobenzothiadiazole, **4**.²⁹

Compound **4** underwent Stille coupling reaction with (5-hexylthiophen-2-yl)trimethylstannane, **1**, in the presence of $\text{Pd}_2(\text{dba})_3$ pre-catalyst and tri(*o*-tolyl)phosphine ligand to generate **5** as a monosubstituted product.³⁰ Due to the formation of unwanted disubstituted analogue, **5** is formed in a moderate yield.³¹ Structures of the compounds **1–5**, synthetic and characterization details (Fig. S1–S9†) are provided in the ESI.† Suzuki coupling reaction between **5** and 4-(hydroxymethyl)

phenylboronic acid and 4-aminomethylphenylboronic acid, **8**, using $\text{Pd}(\text{PPh}_3)_4$ as catalyst and sodium carbonate as base produced compounds **6** and **9**, respectively (Scheme 1).³² The structure of PC_{61}BM was modified using two different pathways, one as on the side chain other on to the fullerene ring. In the former case, benzothiadiazole derivative **6** was attached to the PC_{61}BM molecule *via* the ester group, with the replacement of methyl group in PC_{61}BM with the BTF_2 to form fullerene adduct **7**, $\text{PC}_{61}\text{BTF}_2$. Deprotection of the ester group was first carried out to form PC_{61}BA (synthetic and characterization details are provided in the ESI†), followed by a Steglich esterification reaction between **6** and PC_{61}BA in the presence of $\text{EDC}\cdot\text{HCl}$ and DMAP to form **7** as a brown solid in a yield 69%.³³

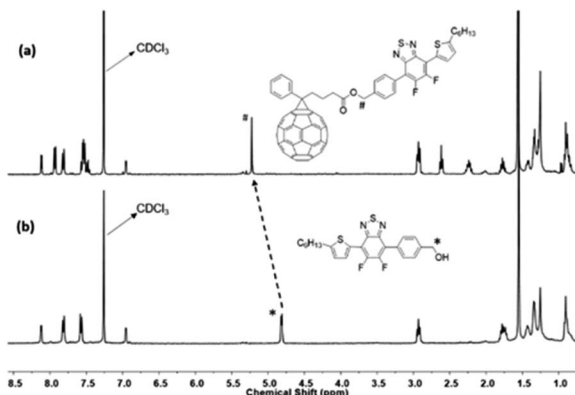
To understand the structure–activity relationship *i.e.*, attaching the Ph- $\text{BTF}_2\text{-Th}$ to PC_{61}BM at the fullerene core, in the latter case, a bis-adduct fullerene derivative (compound **10**, Scheme 1) was also synthesized. Functionalization of the C_{60} frame can be achieved by several reactions to form different fullerene derivatives.³⁴ Prato reaction is one of the most widely used reactions to functionalize fullerene. It proceeds *via* a 1,3-dipolar cycloaddition of the intermediate ylide, formed by the decarboxylation of the iminium salt, onto a [6,6] double bond of the fullerene ring to form fulleropyrrolidines.³⁵ Prato-type reaction between **9** and PC_{61}BM , using paraformaldehyde yielded compound **10**. Different bis-adduct isomers, as well as multi-adduct fullerenes, were obtained. From the purification by column chromatography, two fractions were collected, **10-D** and **10-I**, the latter being the major fraction. Fraction **10-I** is most likely the bis-adduct regio-isomers (abbreviated as $\text{PC}_{61}\text{BM-BTF}_2$) and **10-D** (abbreviated as $\text{PC}_{61}\text{BM-}n\text{BTF}_2$, where $n > 1$), the multi-adduct derivatives. The regio-isomers involved the addition of BTF_2 at the fullerene core. The bis-adduct isomers were not separated since it has been reported that the LUMO energy levels of the different isomers are comparable enough that their photovoltaic properties are not affected.^{15a,b}

The formation of the different compounds was followed by ^1H , ^{13}C , and ^{19}F NMR. The synthesis of compound **5** was confirmed by comparing the ^1H NMR spectrum of product **5** and that of **1** (see Fig. S1 and S8†). In CDCl_3 , a significant downfield shift of the characteristic thiophene “a” proton from 7.04 ppm (in compound **1**) to 8.07 ppm (in compound **5**) is observed due to the replacement of the trimethylstannane moiety by fluorinated benzothiadiazole group. The highly electronegative fluorine atoms in the molecule led to the deshielding of the “a” proton and thus account for a downfield shift of the corresponding signals. From Fig. S10†, ^1H NMR spectrum of the desired Suzuki product **6**, obtained by coupling reaction of **5** with (hydroxymethyl)phenylboronic acid, is evident from the formation of new signals in the aromatic region at 7.80 ppm (“b” proton) and 7.57 ppm (“c” proton) and the benzylic (Ph-CH_2) “e” protons at 4.80 ppm. The successful esterification reaction between the alcohol, **6** and the acid, PC_{61}BA (NMR is provided in Fig. S12†), and formation of modified PC_{61}BM with the replacement of methyl ester with benzyl ester *i.e.*, fullerene linked with Ph- $\text{BTF}_2\text{-Th}$ (mono-adduct **7**) is also suggested by the prominent chemical shift of benzylic (Ph-CH_2) protons from 4.80 ppm (“*” proton) to



Scheme 1 Synthesis of **7** $\text{PC}_{61}\text{BTF}_2$, **10** $\text{PC}_{61}\text{BM-}n\text{BTF}_2$ where $n \geq 1$, reagents and conditions: (i) $\text{Pd}(\text{PPh}_3)_4$, Na_2CO_3 , THF, $66\text{ }^\circ\text{C}$, 18 h, 47% (ii) $\text{EDC}\cdot\text{HCl}$, DMAP, ODCB, $0\text{ }^\circ\text{C}$ to rt, 2 d, 69% (iii) $\text{Pd}(\text{PPh}_3)_4$, K_2CO_3 , THF, $66\text{ }^\circ\text{C}$, 18 h, 83% (iv) CH_2O , PhMe, $110\text{ }^\circ\text{C}$, 18 h, 42% (for **10-I**).



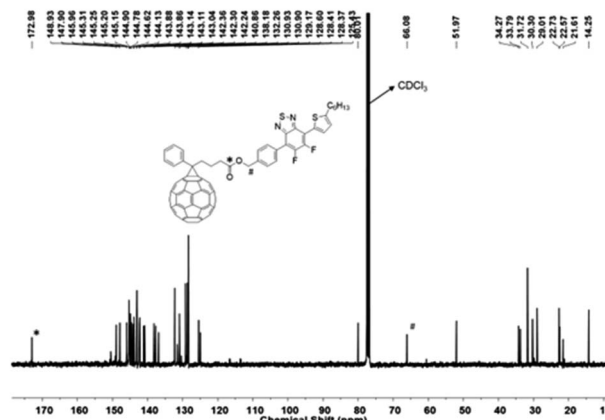
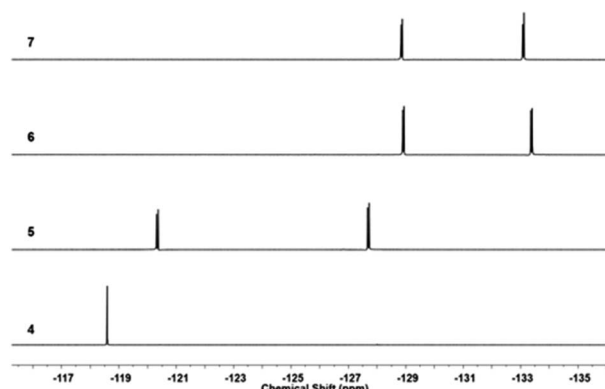
Fig. 2 ^1H NMR spectra of (a) 7 and (b) 6 in CDCl_3 .

5.22 ppm ("#" proton), as shown in Fig. 2. This resonance towards the downfield value is associated with the deshielding effect induced by the adjacent ester group. Detailed structure characterization to confirm the formation of 7 is provided in Fig. S13.[†]

Likewise, the formation of bis- and multi-adducts of PC_{61}BM with $\text{Ph-BTF}_2\text{-Th}$ was studied with NMR. The structures of precursor compound **8–9** were confirmed by NMR (Fig. S14–S17[†]). ^1H NMR confirms the formation of bis- and multi-adducts of PC_{61}BM . Fig. S18[†] shows the stacked ^1H NMR spectra of PC_{61}BM , **9**, **10-D**, and **10-I** and it can be seen that the aromatic region changes from well-defined signals in PC_{61}BM and **9**, to broad multiplets in **10-D** and **10-I**. Similarly, the aliphatic region shows additional signals, which may correspond to the splitting of signals due to the fullerene ring, as reported in the literature.³⁶

Further structural confirmation of successful synthesis of the compound was confirmed by ^{13}C NMR spectroscopy. The ^{13}C NMR spectrum of the Stille coupling product **5**, Fig. S9,[†] showed a shift in the carbon signal of C–Br after replacement of bromide with thiophene, from 99.39 ppm (Fig. S7[†]) in the dibromo compound **4** to 96.34 ppm in **5**. This carbon signal bound to bromine *i.e.*, C–Br at 96.34 ppm also confirms the Suzuki coupling reaction product, **6**, as indicated by the disappearance of resonance at 96.34 ppm in the ^{13}C NMR spectrum of **6** (Fig. S11[†]) due to bromine substitution. ^{13}C NMR spectrum in Fig. 3 supports the formation of the benzyl ester of fullerene linked with $\text{Ph-BTF}_2\text{-Th}$ to form $\text{PC}_{61}\text{BTF}_2$, **7**. A new signal at 172.97 ppm corresponds to the carbonyl carbon as well as several other resonances in the region 125 and 148 ppm due to the fullerene ring were observed. Additionally, a shift in the benzyl CH_2 signal from 65.21 ppm in alcohol, **6** (Fig. S11[†]) to 66.08 ppm in the ester product, **7** was recorded (Fig. 3). In the case of the bis- and multi-adducts, the ^{13}C NMR spectra show multiple complex signals, which remained inconclusive to confirm the structures of products, **10-I**, and **10-D**, as shown in Fig. S19 and S20,[†] respectively.

^{19}F NMR of the fluorine-containing compounds was recorded and each reaction undergone by the compounds was reflected by the shifts of the corresponding F signals. Fig. 4 shows

Fig. 3 ^{13}C NMR spectrum of 7 in CDCl_3 .Fig. 4 Stacked ^{19}F NMR spectra of **4**, **5**, **6**, and **7** in CDCl_3 .

the stacked ^{19}F NMR spectra of **4**, **5**, **6**, and **7** ($\text{PC}_{61}\text{BTF}_2$). Transformation of symmetrical structure dibromo-difluorobenzothiadiazole, **4** to asymmetrical structure bromothiophenyl-linked benzothiadiazole, **5** is confirmed by the appearance of initial one F signal at -118.59 ppm transformed to two doublets (-120.37 and -127.72 ppm). The substitution of the second bromine atom in **5** by the benzyl alcohol group is apparent by the shift in F resonance value from -127.72 and -120.36 ppm in **5** to -128.89 and -133.36 ppm in **6**, respectively. The esterification reaction of **6** did not significantly affect the magnetic environment of fluorine in **7**. As a result, no major change in the ^{19}F NMR was noticed, with the appearance of two doublets at a similar position (-128.88 and -133.12 ppm) in the $\text{PC}_{61}\text{BTF}_2$ structure.

^{19}F NMR was also helpful to confirm the successful linking of compound **9**, $\text{Ph-BTF}_2\text{-Th}$ anchoring on to PC_{61}BM (Fig. 5) to form **10**. Similar to the mono-adduct fullerene, **7**, the ^{19}F NMR spectra of the multi-adduct derivatives only showed a slight shift in the fluorine signals when comparing **9** to the multi-adduct products, **10-I** and **10-D**. The signal at -128 ppm appeared, as a doublet in all cases, whereas the doublet at -133 ppm in **9** undergoes an upfield shift and now appears as broad peaks in **10-I** and **10-D** ^{19}F NMR spectra. This change may account for the formation of a mixture of isomers during the



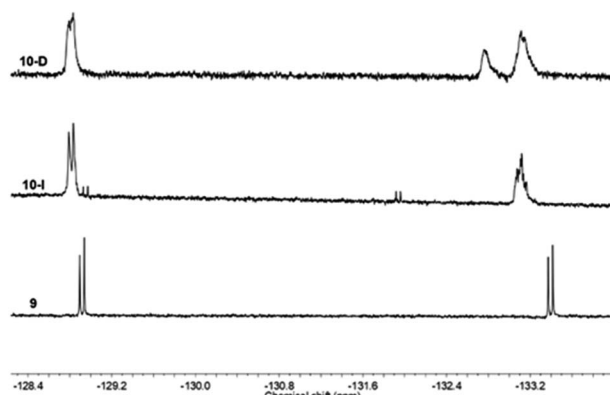


Fig. 5 Stacked ^{19}F NMR spectra of **9**, **10-D**, and **10-I** in CDCl_3 .

Prato reaction and was recently reported *via* light-controlled regioselective synthesis of C_{60} adducts.³⁷ In the case of **10-D**, an additional signal at -132 ppm is noticed, suggesting a different magnetic environment of the fluorine atoms. This supports the addition of multiple benzothiadiazole units onto the fullerene ring to form multiple adducts, $\text{PC}_{61}\text{BM-}n\text{BTF}_2$.

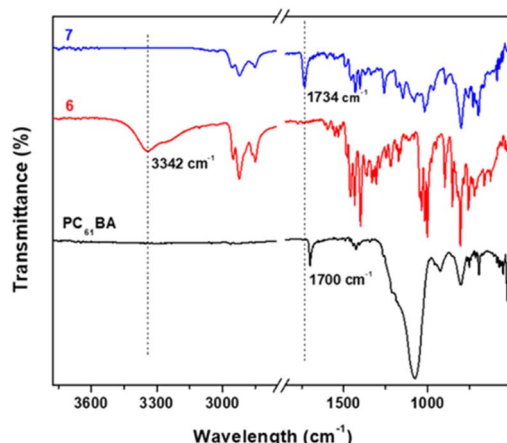


Fig. 6 Stacked FTIR spectra of **7**, **6**, and PC_{61}BA .

FTIR spectroscopy is additionally used to probe the successful esterification reaction between PC_{61}BM and **6**. Fig. 6 shows the stacked FTIR spectra, with **7** ($\text{PC}_{61}\text{BTF}_2$) having characteristic bands of both reactants PC_{61}BA and **6**. The O-H stretching peak at 3342 cm^{-1} in **6** is absent in the spectrum of **7**, indicating successful co-reaction. The weak broad O-H stretching in PC_{61}BA at 3270 cm^{-1} also disappears in **7**. Moreover, the carbonyl C=O stretching undergoes a shift from 1700 cm^{-1} in PC_{61}BA to 1734 cm^{-1} in **7** after covalently linking to the benzothiadiazole unit.

Optical and electrochemical properties

UV-visible absorption spectra of PC_{61}BM (instead of PC_{61}BA due to its poor solubility in ODCB), benzothiadiazole alcohol **6**, and their corresponding ester, **7** $\text{PC}_{61}\text{BTF}_2$, in ODCB are represented in Fig. 7A. From the UV/vis spectra, fullerene benzothiadiazole mono-adduct **7** shows the combined absorptions of the alcohol, **6**, and PC_{61}BM . At 300 nm , all three compounds show absorption peaks corresponding to the localized $\pi \rightarrow \pi^*$ transition in the phenyl rings, similar to that reported in the literature.³⁸ The sharp band at 332 nm in PC_{61}BM , which is associated with the $\pi \rightarrow \pi^*$ transition in the fullerene ring, undergoes a slight hypsochromic shift of 6 nm in **7** ($\lambda_{\text{max}} = 326\text{ nm}$). The absorption observed at 420 nm in the spectrum of **6** corresponds to the characteristic intramolecular charge transfer (ICT) transition between electron acceptor (benzothiadiazole moiety) and electron donor (aromatic group) units in **6**.^{38,39} The existence of ICT transition is also present in $\text{PC}_{61}\text{BTF}_2$ **7**, along with an additional absorption observed in the 375 to 480 nm region due to the fullerene ring. In the spectrum of **7**, a very weak and broad absorption band in the range 445 to 635 nm is also noticed, which is assigned to the first-order forbidden transitions characteristic of fullerenes.⁴⁰ The UV-visible spectra of compounds **9**, **10-D**, and **10-I** were also recorded in ODCB, as shown in Fig. 7B (Fig. S21,† as recorded). Due to their very close structural similarity, compounds **6** and **9** have similar UV-visible spectra, with absorptions centered at 300 and 420 nm . Along with the retention of absorption features of **9** in $\text{PC}_{61}\text{BM-}n\text{BTF}_2$, the presence of fullerene in **10-D** and **10-I** is supported by

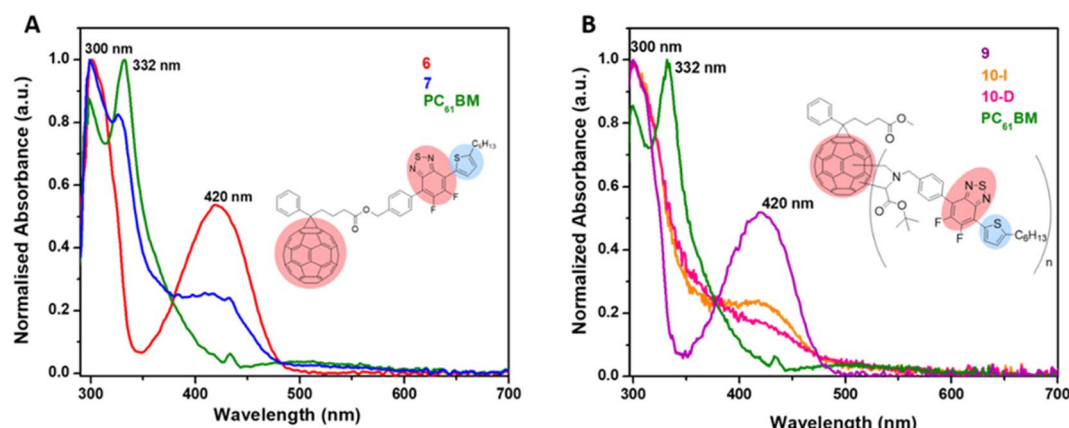


Fig. 7 Normalized UV-visible spectra of (A) **6**, **7**, and PC_{61}BM and (B) **9**, **10-I**, **10-D**, and PC_{61}BM in ODCB (concentrations of 10^{-5} M).



broadening of the bands from 315 to 375 nm due to $\pi \rightarrow \pi^*$ transition. Likewise in the case of $\text{PC}_{61}\text{BTF}_2$ 7, a lower intensity absorption at 420 nm is additionally observed in both **10-D** and **10-I**, which extends to up to \sim 600 nm. The optical band gap of monoadduct 7 and bisadduct **10-I** was calculated by Tauc plot using the solution data and found to be 2.58 and 2.52 eV respectively (Fig. 8).

To further probe the effect of BTF_2 substitution on fullerene at the core (**10-D** and **10-I**) or side chain (7), electrochemical properties of the fullerene derivatives were performed (Fig. 9A and B). Cyclic voltammetry was employed to determine the electrochemical energy gap. The potentials were calibrated by a ferrocene/ferrocenium (Fc/Fc^+) redox couple under the assumption that the Fc/Fc^+ energy level was -4.8 eV below the vacuum level. The estimated HOMO and LUMO energy levels from the onset oxidation and reduction potentials are determined to study the effect of covalent modification of the fullerene derivatives as compared to PC_{61}BM . Fig. 9 shows a total of four reversible/quasi-reversible reduction potential peaks for both compounds 7 and **10-I**. In the negative region, the minima show the reduction events in which electrons are being added to the LUMO of the molecule whereas the maxima in the positive region represent oxidation in which electrons are being removed.

Details of electrochemical data are summarized in Table 1. Monomer 7 reduces at a lower potential than bis adduct **10-I**. Covalent functionalization of PC_{61}BM raises the LUMO levels of both the adducts, however, compound **10-I** shows the uplifted LUMO as well HOMO levels compared to 7.⁴¹ The uplifted LUMO level of molecule **10-I** compared to molecule 7 suggests better electron-accepting properties of the latter which may be prudential for electron mobility. The HOMO energy level of 7 is found to be deeper than both **10-I** and PC_{61}BM as can be seen in

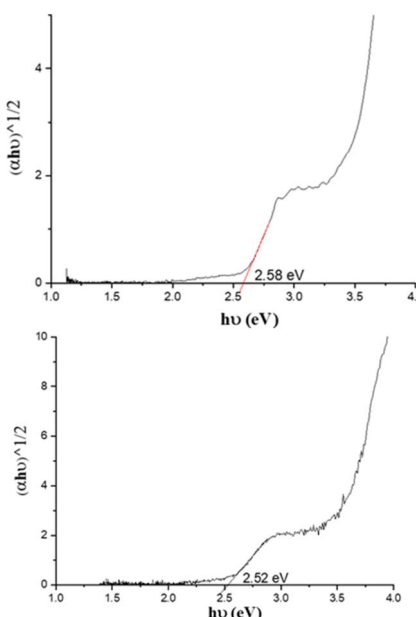


Fig. 8 Optical band gap calculated by Tauc plot of monoadduct 7 (top) and bisadduct **10-I** (bottom).

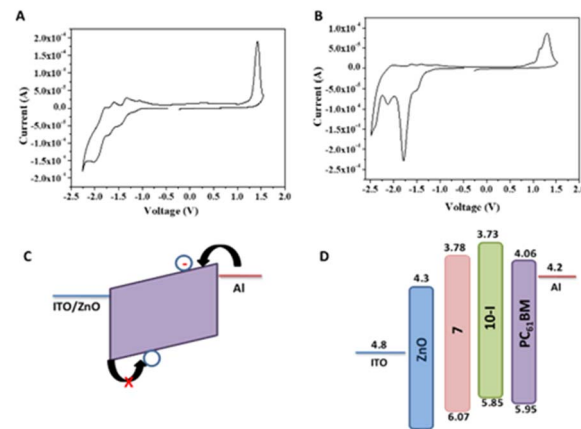


Fig. 9 Cyclic voltammetry curves of (A) 7 and (B) **10-I** (v/s Fc/Fc^+) with $0.1 \text{ M } n\text{-Bu}_4\text{NPF}_6$ as a supporting electrolyte and scan rate 100 mV s^{-1} . (C) Schematic representation of fabricated electron-only device and (D) energy level diagram with HOMO and LUMO values.

Table 1. Both optical and electrochemical bandgap values follow the same trend $\text{PC}_{61}\text{BM} < \text{PC}_{61}\text{BM-}\text{BTF}_2 < \text{PC}_{61}\text{BTF}_2$ range. The observed discrepancy in the calculated value of bandgap from two different techniques, the electrochemical vs. optical measurements, is accounted to the variation in key aspects of the physical process *viz.* adiabatic or vertical, solvation, exciton binding energy, *etc.*⁴²

Charge transport properties

The electron-only devices based on 7 and **10-I** were fabricated for studying the charge transport properties using the space-charge-limited current (SCLC) method.⁴³ To investigate detailed electron mobility properties, we fabricated electron only device with configuration: ITO/ZnO/material/Al. SCLC is observed in electron/hole only devices in which using right contact of electrodes either only electrons or hole can be injected. The current so obtained is only space charge limited at higher voltage due to ohmic contact and trap free transport.

ITO was deposited on a simple glass slide to form an ITO substrate. On to ITO substrate, the ZnO layer was deposited to block hole injection from ITO. Fullerene adduct (7/**10-I**) film was spin-coated using a 40 mg mL^{-1} solution in chlorobenzene. The coated substrate was annealed at $\sim 120^\circ\text{C}$ for 15 min. Finally, the Al layer was deposited above the fullerene derivative layer. The n-type semiconductor properties were measured using a Keithley 2420 power source. The electron mobility of the products (7/**10-I**) was calculated using the Mott–Gurney eqn (1).

$$J = \frac{9}{8} \mu \epsilon \frac{V^2}{L^3} \quad (1)$$

where J is current density, V is the voltage drop across the device, L is the active layer thickness, μ is the mobility of charge carrier and ϵ is the permittivity of the current transporting medium. The current density–voltage (J – V) characteristics of the electron-only diodes are measured based on the geometry as shown in Fig. 9c. At lower applied voltage, ohmic region is observed and after it, trap-controlled space charge limited conduction (SCLC)

Table 1 Optical and electrochemical data of fullerene adducts

Entry	$\lambda_{\text{max}}^{\text{abs}}$ (nm)	$\text{Red}_{\text{onset}}^a$ (V)	Red^{-1} (V)	Red^{-2} (V)	Red^{-3} (V)	Red^{-4} (V)	LUMO (eV)	$\text{Ox}_{\text{onset}}^a$ (V)	Ox^{-1} (V)	HOMO (eV)	$E_{\text{g}(\text{cv})}$ (eV)	$E_{\text{g}(\text{opt})}^b$ (eV)
PC₆₁BM	332	-0.74	-0.91	-1.17	-1.40	-1.95	-4.06	1.15	—	-5.95	1.89	2.12
7	326, 385 sh	-1.02	-1.14	-1.45	-1.70	-2.01	-3.78	1.27	—	-6.07	2.29	2.58
10-I	326, 385 sh	-1.07	-1.50	-1.78	-2.12	-2.40	-3.73	1.05	1.14	-5.85	2.12	2.52

^a Potentials are measured relative to a Fc/Fc⁺ redox couple. ^b Optical bandgap was estimated from Tauc plot; sh stands for the shoulder.

is observed. The SCLC region in each J - V characteristic is fitted as shown in Fig. 10. The electron mobility thus determined is $\sim 7.1 \pm 0.76 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ for 7, which is nearly comparable to the electron mobility of $10.0 \times 10^{-4} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ of PC₆₁BM.⁴⁴ The electron mobility is $\sim 4.6 \pm 0.60 \times 10^{-5} \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$ for **10-I**. PC₆₁BT₂ 7 showed higher electron mobility than PC₆₁BM-BT₂ **10-I** due to better electron-accepting ability as evident from cyclic voltammetry measurements. Structurally, the fullerene core has retained the extended conjugation in PC₆₁BT₂ due to functionalization at the arm with the replacement of -OMe group in PC₆₁BM to BT₂ in PC₆₁BT₂ 7. While in **10-I**, a structure involved substitution at the fullerene core thus decreasing the degree of effective conjugation length accounting for lower electron mobility than 7.

Theoretical calculations

To probe the effect of functionalization with BT₂ on energy properties theoretical calculations were performed for geometry PC₆₁BM, PC₆₁BT₂ 7, PC₆₁BM-*n*BT₂ **10-I** (*n* = 1), and **10-D** (*n* = 2, and 3, shown in the ESI Fig. S22 and S23[†]). Structures were constructed for PC₆₁BM with phenyl-butyric-acid-methyl groups latched on neighboring carbon atoms. Attempt for latching on next neighboring C₆₀ carbon atoms was also examined for completeness but was not showing an energetically favorable configuration. For adduct **10**, functionalization with one (*n* = 1) BT₂ group was performed on two distinct PC₆₁BM locations, specifically, (a) with the BT₂ group positioned at a *para* position to phenyl-butyric acid methyl ester (PBM) moiety, and (b) with BT₂ located at a (near *meta*) position that was found suitable for the later construction of the doubly functionalized PC₆₁BM, to accommodate comparisons. For functionalization with *n* = 2 BT₂ groups, two near meta locations (analogous to the aforementioned configuration of the *n* = 1 case) were chosen so that the distance from PBM is the same as the distance between them, specifically, with latching carbon atoms at five hops

(PC₆₁BM C atoms) from each other. Finally, for functionalization with *n* = 3 BT₂ groups, the situation was complicated due to the spatial extent of the BT₂ groups, so latching locations were chosen in a manner to avoid steric effects.

The isosurfaces of the frontier orbitals for the PC₆₁BM (reference) and PC₆₁BT₂ structures are shown in Fig. 11a-d.

From the isosurface plots, the HOMO and LUMO are found to be localized in the fullerene core in the case of PC₆₁BM as can be seen in Fig. 11a and b. Interestingly in the case of PC₆₁-BT₂, the HOMO was significantly localized over the difluorobenzothiadiazole (BT₂) and LUMO extended over the fullerene structure than over the thiadiazole moieties.

The isosurfaces of the frontier orbitals for the three singly functionalized PC₆₁BM structures with BT₂ are shown in Fig. S22a-f.[†] The HOMO and LUMO of all three structures are localized on the fullerene cage, whereas for all three cases the HOMO extends over the latching carbon of the BT₂ moiety. From Table 2 a clear dependence can be seen of the HOMO eigen energies to the number of attached BT₂ functional groups. This trend is consistent throughout *n* = 1-3. For the singly functionalized groups, the HOMO of the *para* and @conf3 functionalized structures are energetically higher when

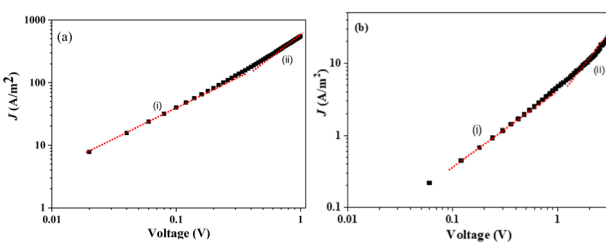


Fig. 10 J - V plots for the electron-only devices based on adduct (a) 7 and (b) **10-I**.

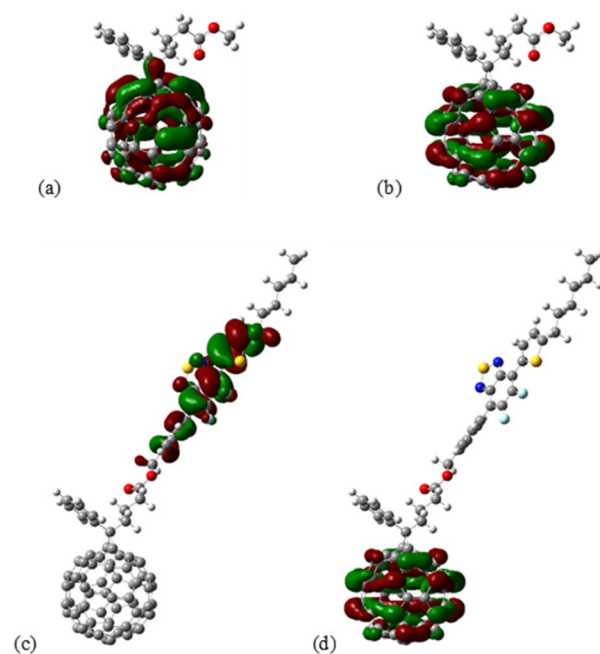


Fig. 11 Isosurfaces of the (a) HOMO and (b) LUMO of PC₆₁BM, as well as the (c) HOMO and (d) LUMO of 7 PC₆₁BT₂ structure. Computed at the M06/def2-SVP level of theory.



Table 2 Energy and optical properties of the pure and functionalized PC_{61}BM structures are determined by theoretical calculations (frontier orbital eigen energies, energy gaps, onset of absorbance, oscillator strength, and ground-state dipole moment, μ_g)

Structure	LUMO (eV)	HOMO (eV)	E_g (eV)	Optical gap (eV)	λ_{\max} (nm)	f , oscillator strength	Dipole moment μ_g (D)
PC₆₁BM	−3.25	−6.11	2.86	1.96	631	0.002	5.50
PC₆₁BT₂, 7	−3.35	−6.08	2.73	1.97	628	0.002	5.54
PC₆₁BM-1xBT₂-<i>para</i>, 10-I	−3.14	−5.88	2.75	2.71	458	0.180	2.86
PC₆₁BM-1xBT₂, 10-I	−5.97	−3.14	2.83	1.93	644	0.002	3.43
PC₆₁BM-1xBT₂@conf3, 10-I	−3.07	−5.86	2.79	1.91	649	0.002	5.70
PC₆₁BM-2xBT₂, 10-D	−2.70	−5.76	3.07	2.39	519	0.011	1.28
PC₆₁BM-2xBT₂@conf3, 10-D	−3.38	−5.42	2.04	1.17	1063	0.002	2.72
PC₆₁BM-3xBT₂, 10-D	−3.21	−5.32	2.11	1.20	1030	0.005	2.17

the functional group is closer to the PBM, however, the LUMO has the same energy for both structures.

The isosurfaces of the frontier orbitals for the doubly functionalized PC_{61}BM structure with BT_2 are shown in Fig. S23a–d.† The HOMO and LUMO are localized on the fullerene cage. As in the singly functionalized cages, the HOMO extends over the latching carbon of the BT_2 moiety for the **PC₆₁BM-2xBT₂**. Orientation of BT_2 and latching position have an impact on the HOMO and LUMO delocalization, as can be seen in the case of **PC₆₁BM-2xBT₂@conf3** where the LUMO is noticeably displaced towards the BT_2 latching positions while remaining over the fullerene cage.

The structure and the frontier orbitals of the triply functionalized structure **PC₆₁BM-3xBT₂** are shown in Fig. S24.† To alleviate any steric hindrances the **PC₆₁BM-3xBT₂** was designed with the two functional groups that are closest to the PBM with a different (inversed) orientation compared to the structures with $n = 1$ and 2. The HOMO of $n = 3$ structure is higher than the others, affirming the trend that the HOMO energy increases with the number of attached BT_2 functional groups. Interestingly, the effect of the number of attached BT_2 has a stronger effect on (rising) the HOMO level than the effect that the proximity of the BT_2 group to the PBM group. Structure **PC₆₁BM-3xBT₂** offers an example of both of these actions and exhibits the highest HOMO eigen energy. Both HOMO and LUMO values determined theoretically and experimentally for various structures are in an appreciable agreement. The theoretical HOMO value was found to decrease with an increase in substitution in the PC_{61}BM core. The frontier orbital values provided in Table 1 suggest that 7, **10-I** and **10-D** can be used in conjunction with BHJ donors in cases that offer preferred level alignment.

The UV-visible spectra of all the structures under study are provided in Fig. S25.† In analogy with experimental UV-visible spectra, all the structures (**7**, **10-I**, and **10-D**) containing BT_2 exhibit significantly strong UV-visible absorbance theoretically too. In the legends of the plots, the relative scale of the absorbance is given to that of the PC_{61}BM spectrum. The strongest absorbance is noted for the **PC₆₁BM-2xBT₂** structure that is functionalized with two BT_2 moieties at equal distances from the PC_{61}BM . Regardless of the absorbance magnitude, the main peak for all the functionalized structures is centered at nearly a similar wavelength range (458–460 nm). Of the three singly functionalized structures, the one with BT_2 closer to PBM

moiety has a wider and stronger absorbance profile at low wavelengths, near 350 nm. The widest absorbance is noted for the triply functionalized structure that reaches up to large wavelengths, with a low intensity (oscillator strength) onset at $\lambda_{\max} = 1030$ nm. To examine the contributions to the excitations from specific groups of the structures we have partitioned the structures into groups, specifically the fullerene cage, PBM, and BT_2 (except the first structure). In the ESI,† we have provided the main contributions to the excitations, the charge density as well as the change in charge density on each group (Table S1a–h†). Excitations that result in considerable charge transfer are denoted as bold. We noted that specifically for **PC₆₁BM-1xBT₂-*para*** and **PC₆₁BM-2xBT₂** the optical gap listed in Table 2 is the 10th and 3rd computed excitation, respectively, that correspond to the first excitations with non-negligible oscillator strengths.

Incidentally, the transition dipole moments for **PC₆₁BM-1xBT₂-*para*** listed in Table S2† is comparatively considerably higher and corresponds to the transition with the first non-negligible oscillator strength that is not the first computed excitation. In the ESI,† we provide detailed values for the transition dipole moments for all excitations on all structures. The dipole moment of the adduct in the ground state (μ_g) was found to slightly increase with the substitution of BT_2 in place of the methyl group in PC_{61}BM , suggesting a marginal increase in polarity of the structure. It is noteworthy that when BT_2 is attached to the core at a para position to the PBM moiety (**10-I**) the μ_g value was found to decrease by nearly 2-fold, suggesting a significant cancellation of dipole moments. This alone suggests that the μ_g value may increase when BT_2 units are attached to the fullerene core. It turns out that this variation is manifested when 2 vs. 3 BT_2 units are attached to the fullerene core. This confirms that the existence of a BT_2 unit in the direction of PBM has a significant controlling effect on the overall dipole moment of the structure. Such variation in dipole moment may induce polarization which is desired to provide directionality to charge carrier mobility and aid charge separation.

Experimental

Materials

PC_{61}BM was purchased from Lumtec, trimethyltin chloride, triethylamine, tri(dibenzylideneacetone)dipalladium



(Pd₂(dba)₃), tri(*o*-tolyl)phosphine, tetrakis(triphenylphosphine) palladium(0) (Pd(PPh₃)₄) from Sigma-Aldrich, 2-hexylthiophene, ammonium formate, Pd/C (10%) and 4-(hydroxymethyl)phenylboronic acid from Alfa Aesar, thionyl chloride, bromine, carbon disulfide, and *N*-(3-dimethylaminopropyl)-*N*'-ethylcarbodiimide hydrochloride (EDC·HCl) from Specrochem, 4,5-difluoro-2-nitroaniline from TCI, iron powder from SRL, hydrochloric acid (35%) from Merck Millipore, glacial acetic acid from Fisher Scientific and 4-(dimethylamino)pyridine (DMAP), *n*-butyllithium and sodium sulfate from Chemlabs. All the chemicals were used without further purification. All solvents used were AR grade. Tetrahydrofuran (THF) was distilled over sodium using benzoquinone as an indicator under nitrogen atmosphere. Literature procedures were followed in the preparation of compounds **1**,⁴⁵ **2**,⁴⁶ **3**,⁴⁷ **4**,²⁹ **5**,³¹ **8** (ref. 48) and PC₆₁BA⁴⁹ (see ESI†).

Measurements

Proton (¹H), carbon (¹³C), and fluorine (¹⁹F) NMR spectra of the compounds were recorded on a Bruker AV400 NMR spectrometer at a proton frequency of 400 MHz and corresponding carbon, fluorine frequencies at room temperature, using CDCl₃ and DMSO-d₆ as solvents and tetramethylsilane (TMS) as an internal standard and plotted using MestReNova software. Fourier Transform Infrared (FTIR) spectra were recorded on a Nicolet iS5 spectrometer equipped with the iD5-ATR accessory, in the range of 4000 to 400 cm⁻¹. Mass spectrometry analysis was carried out using Agilent HRMS Q-ToF 6540 Series equipped with ESI mode. UV-visible spectra were collected using a Thermo Scientific Evolution 201 spectrophotometer. Samples were dissolved in *o*-dichlorobenzene (ODCB) and spectra were recorded from 200 to 800 nm with a resolution of 1 nm, using a 1.0 cm UV quartz cuvette. Three electrode system was used while performing cyclic voltammetry (CV) where, the platinum disc was used as a working electrode, silver-wire, and a platinum-wire as reference and counter electrode respectively in a 0.1 M *n*-Bu₄NPF₆(tetra-*n*-butylammonium hexafluorophosphate) supporting electrolyte in anhydrous acetonitrile solution. CV of the prepared fullerene adducts and PC₆₁BM was performed by drop-casting film (from a solution in dichloromethane) on Pt disc working electrode and dried in vacuum for two hours. Current *vs.* voltage was measured on an Autolab potentiostat. The energy of the highest unoccupied molecular orbital levels (*E*_{HOMO}) and the lowest unoccupied molecular orbital levels (*E*_{LUMO}) were calculated according to the following eqn (2).

$$E_{(\text{LUMO or HOMO})} = -[4.8 + E_{\text{onset(red or ox)}}] \text{ eV} \quad (2)$$

where *E*_{ox} and *E*_{red} were estimated from cyclic voltammetry curve using a ferrocene/ferrocenium redox couple as an internal standard. Electron mobility of the synthesized molecules was determined by the space charge limited current (SCLC) method for electron-only devices with structure: ITO/ZnO/material/Al. A zinc oxide (ZnO) layer of 30 nm thickness was deposited on a pre-cleaned ITO/Glass substrate using the spin coating method. The materials were dissolved in ODCB with

a concentration of 40 mg mL⁻¹. The solution-processed thin film of both the materials **7** and **10-I** with a thickness of ~200 nm each were deposited through spin coating for fabrication of electron-only diodes (ITO/ZnO/7 or **10-I**/Al) in sandwich geometry. A thermal evaporated Al metal the thin film was used as top metal electrodes for electron-only diodes. The effective device area of these devices was 6 mm².

All of the structures were examined theoretically *via* computations based on density functional theory (DFT) and time-dependent density functional theory (TD-DFT).⁵⁰ The M06 (ref. 50d) functional was used throughout for both ground- and excited-state computations. The results reported are all from computations performed using the Gaussian package.⁵¹ The ESI† includes further technical details on the computational methods and methodologies employed.

Synthetic procedures

Synthesis of (4-(5,6-difluoro-7-(5-hexylthiophen-2-yl)benzo[c][1,2,5]thiadiazol-4-yl)phenyl)methanol, 6.³² To a solution of 5 (200 mg, 0.48 mmol) in anhydrous THF (50 mL) under nitrogen was added a nitrogen purged aqueous solution of Na₂CO₃ (2 M, 10 mL), followed by 4-(hydroxymethyl)phenylboronic acid (80 mg, 0.53 mmol) and Pd(PPh₃)₄ (28 mg, 0.02 mmol). The mixture was stirred under reflux for 16 h under inert atmosphere. The solvent was evaporated, and diethyl ether (10 mL) was added to dissolve the residue. Water (10 mL) was added, and the organic layer was extracted in diethyl ether (2 × 10 mL). The combined organic phase was dried over anhydrous Na₂SO₄ and solvent was removed *in vacuo*. Purification of the crude product by silica gel chromatography (eluent CH₂Cl₂) resulted in a yellow solid. Yield: 100 mg (47%).

¹H NMR (CDCl₃, 400 MHz, ppm): 8.12 (d, 1H, *J* = 3.72 Hz, C-C-CH), 7.81 (d, 2H, *J* = 7.56 Hz, ArH), 7.56 (d, 2H, *J* = 8.12 Hz, ArH), 6.95 (d, 1H, *J* = 2.80 Hz, CH₂-C-CH), 4.81 (d, 2H, *J* = 5.56 Hz, HO-CH₂), 2.93 (t, 2H, *J* = 15.44 Hz, SC-CH₂), 1.80–1.33 (m, 8H, AlkH), 0.90 (t, 3H, *J* = 13.64 Hz, -CH₃).

¹³C NMR (CDCl₃, 100 MHz, ppm): 150.65, 150.57, 150.38, 150.32, 149.18, 149.09, 148.23, 148.03, 141.91, 131.45, 131.36, 130.88, 130.85, 129.78, 127.09, 124.92, 65.21, 31.72, 30.29, 29.01, 22.73, 14.24.

¹⁹F NMR (CDCl₃, 376 MHz, ppm): -128.93 (d), -133.40 (d). MS (ESI(+)): [(M + H)⁺]: 445.1 (444.1 calculated). FTIR (cm⁻¹) = 3348 ($\nu_{\text{O-H}}$).

Synthesis of [6,6]-phenyl-C₆₁-butyric acid(4-(5,6-difluoro-7-(5-hexylthiophen-2-yl)benzo[c][1,2,5]thiadiazol-4-yl)phenyl)methanoate, 7.⁵² PC₆₁BA (80 mg, 0.08 mmol) was added to ODCB (15 mL), sonicated for five minutes and kept under nitrogen atmosphere. To this solution, compound **6** (40 mg, 0.09 mmol) was added and reaction mixture stirred in an ice bath at 0 °C. After attainment of temperature, EDC·HCl (34 mg, 0.17 mmol) and DMAP (22 mg, 0.18 mmol) were added and stirring was continued at 0 °C for 3 h. Reaction was allowed to warm to room temperature and stirred for further two days. Reaction mixture was purified over a silica bed using hexane as eluent to remove the ODCB solvent. The product was then isolated by silica gel chromatography (eluent hexane/toluene 1 : 1



then pure toluene) as a brown solid. Yield: 50 mg (37%). ^1H NMR (CDCl_3 , 400 MHz, ppm): 8.11 (d, 1H, $J = 3.64$ Hz, C–C–CH), 7.92 (d, 2H, $J = 7.16$ Hz, ArH), 7.81 (d, 2H, $J = 7.52$ Hz, ArH), 7.57–7.47 (m, 5H, ArH), 6.95 (d, 1H, $J = 3.36$ Hz, CH_2 –C–CH), 5.22 (s, 2H, HO–CH₂), 2.93 (t, 4H, $J = 14.56$ Hz, SC–CH₂), 2.62 (t, 2H, $J = 14.68$ Hz, OOC–CH₂), 2.28–2.20 (m, 2H, AlkH), 1.77 (quintet, 2H, $J = 30.04$ Hz, AlkH), 1.47–1.33 (m, 4H, AlkH), 0.89 (t, 3H, $J = 9.64$ Hz, –CH₃). ^{13}C NMR (CDCl_3 , 100 MHz, ppm): 172.98, 150.55, 150.50, 149.21, 148.93, 147.90, 145.96, 145.31, 145.20, 145.15, 144.90, 144.81, 144.78, 144.62, 144.54, 144.13, 143.88, 143.86, 143.20, 143.14, 143.11, 143.04, 142.36, 142.30, 142.24, 141.11, 140.86, 138.18, 138.01, 137.70, 136.86, 132.26, 131.55, 131.46, 130.93, 129.17, 128.60, 128.41, 128.37, 125.43, 124.95, 80.01, 66.08, 51.97, 34.27, 33.79, 31.72, 30.30, 29.01, 22.73, 22.57, 21.61, 14.25. ^{19}F NMR (CDCl_3 , 376 MHz, ppm): –128.88 (d), –133.12 (d). MS (ESI(+)): [(M + H)⁺]: 1323.1 (1322.1 calculated). FTIR (cm^{-1}): 1735 ($\nu_{\text{C=O}}$).

Synthesis of *tert*-butyl-2-((4-(5,6-difluoro-7-(5-hexylthiophen-2-yl)benzo[*c*][1,2,5]thiadiazol-4-yl)benzyl)amino)acetate, 9.³² To a dry reaction tube under nitrogen was added compound 8 (63 mg, 0.24 mmol) and dry THF (1 mL). Aqueous solution of K_2CO_3 (2 M, 0.056 mL), purged with nitrogen gas, was added to the above solution and mixture stirred under reflux for 15 min. 5 (50 mg, 0.12 mmol) and $\text{Pd}(\text{PPh}_3)_4$ (14 mg, 0.01 mmol) were added to the stirring mixture and reaction was continued under reflux for 18 h. After evaporating the solvent under vacuum, ethyl acetate (5 mL) was added to dissolve the residue followed by the addition of water (10 mL). The aqueous layer was separated, and the organic layer was extracted in ethyl acetate (2 × 5 mL). The organic layers were combined, dried over anhydrous Na_2SO_4 and the solvent was evaporated. The crude sample was purified over silica column chromatography. The first column was run using a gradient of hexane in CH_2Cl_2 as eluent (from 100–0% hexane) and the fragment isolated was purified by a second column in hexane : CH_2Cl_2 (1 : 1) eluent. Product 9 was obtained as a yellow solid. Yield: 55 mg (83%).

^1H NMR (CDCl_3 , 400 MHz, ppm): 8.11 (d, 1H, $J = 3.72$ Hz, C–C–CH), 7.77 (d, 2H, $J = 7.16$ Hz, ArH), 7.52 (d, 2H, $J = 812$ Hz, ArH), 6.95 (d, 1H, $J = 3.56$ Hz, CH_2 –C–CH), 3.90 (s, 2H, Ar–CH₂–NH), 3.37 (s, 2H, NH–CH₂–CO), 2.93 (t, 2H, $J = 15.32$ Hz, SC–CH₂), 1.77 (quintet, 2H, $J = 30.20$ Hz, AlkH), 1.48 (s, 9H, C(CH₃)₃), 1.36–1.25 (m, 8H, AlkH), 0.90 (t, 3H, $J = 14.04$ Hz, –CH₃).

^{13}C NMR (CDCl_3 , 100 MHz, ppm): 171.79, 150.67, 150.59, 150.31, 150.25, 149.10, 148.28, 140.82, 131.40, 131.31, 130.76, 128.52, 124.91, 81.48, 53.14, 51.10, 31.71, 30.28, 29.01, 28.29, 22.73, 14.24.

^{19}F NMR (CDCl_3 , 376 MHz, ppm): –128.94 (d), –133.46 (d).

Synthesis of *tert*-butyl(N-ethyl(4-(5,6-difluoro-7-(5-hexylthiophen-2-yl)benzo[*c*][1,2,5]thiadiazol-4-yl)benzyl)fullerenopyrrolidine carboxylate addition on PC₆₁BM, 10.³⁵ 9 (45 mg, 0.08 mmol, PC₆₁BM (73 mg, 0.08 mmol) and paraformaldehyde (13 mg, 0.4 mmol) were weighed in a reaction tube under nitrogen atmosphere. Dry toluene (2.5 mL) was added, and the tube was sealed with silicone septum. The reaction was carried out at 110 °C for 18 h in dark. Upon completion of the reaction, solvent was evaporated, and the crude product was purified by silica column chromatography

using gradient elution with hexane : CH_2Cl_2 (100 : 0 to 90 : 10). Two fractions, compounds 10-D and 10-I were isolated from the column and dried under vacuum to give two shiny brown solids (19 mg for 10-D and 54 mg for 10-I).

Conclusions

In conclusion, two new fullerene adducts with A–D–A type arrangement were designed, synthesized, and characterized, where the BTF₂–Th unit is covalently anchored to the fullerene either as a terminal group or on to the fullerene core. The molecular structure confirmation of the two fullerene adducts was supported by NMR spectroscopy. The resultant modified structures exhibit wide and strong absorption covering the range from 300 to 600 nm and shown relatively higher absorption in the 350–500 nm region which was nearly absent in PC₆₁BM. Terminal attached BTF₂ unit led to a deeper HOMO energy level and higher electron mobility of the adduct than the one with the modified core. Both energy levels and electron mobility are affected due to different placement of BTF₂ on fullerene, with HOMO energy is deeper than PC₆₁BM for terminally modified structure. Additionally, the HOMO and LUMO were found to be affected in fullerene adducts with dependence on the location of BTF₂ at the periphery *vs.* core of the fullerene. Computational results show that the presence of BTF₂ affects the ground state dipole moments as well as the absorption strengths, most noticeable in the case of *n* = 2. Current work demonstrates that the fluorinated benzodithiazole unit linked to fullerene has tremendous potential as a building block for high-performance material for optoelectronic applications.

Conflicts of interest

There are no conflicts to declare.

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