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Theoretical investigation of benzodithiophenebased donor molecules in organic solar cells: from structural optimization to performance metrics†

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Achieving high power conversion efficiency (PCE) remains a significant challenge in the advancement of organic solar cells (OSCs). In the field of organic photovoltaics (OPVs), considerable progress has been made in optimizing molecular structures to improve the PCE. However, innovative material design strategies specifically aimed at enhancing PCE are still needed. Here, we have designed BDTS-2DPPbased molecules and propose a molecular design approach to develop donor materials that can significantly improve the PCE of OSCs. Density functional theory (DFT) and time-dependent DFT (TD-DFT) methods have been adopted in both gas and solvent phases. Our newly designed molecule M1 shows the highest absorption value ($\lambda_{max} = 846$ nm), highest electron reorganization energy ($\lambda_{e} = 0.18$ eV), and the lowest energy gap ($E_q = 1.81$ eV) among all the designed molecules. M1 molecule also exhibits the highest dipole moment in both gas (10.62 D) and solvent phase (13.62 D), and their ground and excited state dipole moment difference is also higher ($\mu_e - \mu_g = 2.99$ D), which enhances its separation to make it a suitable candidate for charge transfer between HOMO-LUMO (97%). Newly designed molecule M3 is observed to have the highest voltage when the current is zero ($V_{oc} = 1.15$ V) highest PCE value (21.90%) and highest fill factor (FF) value (89.42%). The lowest excitation binding energy is estimated by newly designed molecule M2 ($E_{\rm b}=0.30$ eV), which indicates a higher rate of dissociation during the excitation as observed in transition density matrix (TDM) plots. Utilizing electron density difference maps, the newly designed molecules in dichloromethane solvent exhibited consistent intramolecular charge transfer (ICT). The designed molecules were evaluated against reference molecule R to determine if they exhibit superior optoelectronic capabilities. It is found that all designed molecules (M1-M5) exhibit reduced band gaps, are red-shifted in wavelength in comparison to a reference molecule R, and have remarkable charge motilities in terms of reorganisation energies.

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1. Introduction

A scarcity of energy resources has been a direct result of the continuous increase in the global population over the last few decades. 1,2 The world's fossil fuel reserves and other conventional energy sources are rapidly depleting. In addition to having a negative effect on the climate, burning fossil fuels is a major cause of environmental damage leading to a threatening level of global warming by the greenhouse effect. Renewable, greener, safer, and more eco-friendly options must be promptly pursued in order to solve the problem of limited energy supply and to control environmental pollution.3 Replacing carbon based fuels with nuclear power, biomass, solar power, hydropower, and wind power ensures little or no carbon emissions to the environment. Solar power offers the most promising potential among all these alternatives in terms of sustainable, long-term, and inexpensive energy sources for the globe. Modern society is tremendously encouraging solar

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is hugely boosted as a result of this replacement. Due to the elongation of $\pi{-}\pi$ stacking on sides of chains that include thiophene groups, the BDT unit with linear alkyl-thio groups experiences an enhanced rate of hole transfer. Charge transfer was further improved by adding strong electron-withdrawing groups like DPP, which is planar like BDT. 21

In this study, we have designed five new BDTS-2DPP²² (reference molecule (R)) based donor molecules named as M1, M2, M3, M4 and M5. The central unit in molecule R is a donor and the acceptor groups are attached at its terminal end to make it A- π -D- π -A. All the new molecules contain spacers in between to connect donor and acceptor moieties for efficient charger transfer. All these molecules are linked with five different acceptor groups on the end caps as represented in Fig. S1.† Molecule R serves as the reference point with a well-balanced donor-acceptor structure, providing a baseline for understanding the effects of structural modifications introduced in the other molecules. Molecule M1 features an extension of the conjugated system by adding a thiophene unit, which is intended to enhance electron delocalization. This modification aims to improve light absorption and charge transport properties by increasing the effective conjugation length. Molecule M2 introduces electronwithdrawing groups on the donor part of the molecule and extends the conjugated system through the incorporation of a benzothiadiazole unit. These changes are designed to lower the HOMO level, thereby increasing the V_{oc} and potentially improving the PCE. The additional conjugated units in M2 also contribute to better light-harvesting capabilities. Molecule M3 incorporates additional conjugated units, specifically a thieno[3,2-b]thiophene group, along with structural modifications that increase the planarity of the molecule. The enhanced planarity is expected to improve π – π stacking interactions, which are crucial for efficient charge transport, leading to better charge mobility and higher overall efficiency. Molecule M4 features a more complex structure with both electron-donating groups and an extension of the conjugated system by incorporating a fluorinated benzotriazole unit. These modifications further extend the π -system and are expected to enhance the charge separation process by optimizing the balance between electron donation and withdrawal. The structural changes in M4 are aimed at increasing the FF and overall device performance. Finally, molecule M5 represents the most advanced design, combining the beneficial aspects of M1 through M4. It includes strategic modifications to both the donor and acceptor units, along with additional conjugated systems such as the integration of a fused aromatic ring system (e.g., dithieno[3,2-b:2',3'-d]thiophene). These are designed to maximize light absorption, charge separation, and transport, leading to the highest theoretical PCE among the five molecules.

The significance of studying these molecules lies in their potential to push the boundaries of organic photovoltaic performance. By systematically varying the molecular architecture, including the addition of specific conjugated units like thiophene, benzothiadiazole, thieno[3,2-*b*]thiophene, fluorinated benzotriazole, and fused aromatic rings, we aim to identify the key features that contribute to higher efficiency, stability, and scalability in organic solar cells. The insights gained from this study are crucial for guiding the design of next-

generation materials that could significantly advance renewable energy technologies. Density functional theory (DFT) and time-dependent density functional theory (TD-DFT) serve as powerful tools for conducting computational analysis and exploring the intricate optoelectronic properties of designed molecules. The designed molecules were evaluated against reference molecule (R) to determine if they exhibit superior optoelectronic properties. It was observed that all designed molecules have reduced band gaps, hence red-shifted in assessment to reference molecule R and have remarkable charge motilities.

2. Computational methodology

The calculations were conducted utilising the Gaussian09 simulation package, 23,24 and the results were visualised using Gauss-View 6.0. To confirm which DFT method would be the best for the designed molecules, reference molecule R was initially optimised using four different functionals: B3LYP,25 CAM-B3LYP,26 MPW1PW91,27 WB97XD.28 Once the optimisation was completed using DFT method, ²⁹ the λ_{max} of reference molecule (622 nm)³⁰ was found to be the best correlated with the MPW1PW91 functional (683 nm). We found that for further photovoltaic study, MPW1PW91 is the best functional because correlates with the experimental value of λ_{max} of reference molecule. So, all designed molecules were optimised using the MPW1PW91/6-31G(d,p) functional method. PyMOlyze-1.1 (ref. 31) was used for plotting and visualising the graphs of the density of states (DOS). We also computed other geometric factors like dipole moment (μ) ,³² binding energies $(E_b)^{33}$ and open circuit voltage $(V_{oc})^{34}$ In order to predict the nature of transitions for the reference and all designed molecules, the transition density matrix (TDM)35 was analysed using Multiwfn software.36 For analyzing the charge transfer in the molecules, a critical factor known as reorganisation energy has been used. 37,38 The reorganization energy has a very strong relation with the electron and hole motilities, and it is computed by using equations from Marcus theory.39 The equations for calculating the reorganisation energy of electrons and holes are as follows:

$$\lambda_{\rm e} = [E_{\circ}^{-} - E_{-}] + [E_{-}^{\circ} - E_{\circ}] \tag{1}$$

$$\lambda_{\rm h} = [E_{\circ}^{+} - E_{+}] + [E_{+}^{\circ} - E_{\circ}] \tag{2}$$

 E_{-} is the anionic energy that has been optimised, and E_{-}° is the anionic energy from the neutral molecule after optimisation. E_{+} is the cationic energy that has been optimised, and E_{+}° is the cationic energy from neutral molecules of optimised structures. E_{\circ} are single-point energy calculations for both the positively charged cation and the negatively charged anion whereas, E_{\circ}° , E_{\circ}^{-} , represents the cationic and anionic neutral energies from their optimised structures.

3. Optoelectronic characteristics

3.1 Structural optimization

In this work, we optimised BDTS-2DPP (reference molecule R) using four distinct functional such as; B3LYP, CAM-B3LYP, WB97XD, and MPW1PW91 with basis set 6-31G(d,p).

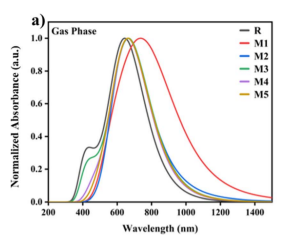
MPW1PW91 is the best chosen functional as the value of λ_{max} for this functional turns out to be 683 nm which matches well with the experimental value *i.e.* 622 nm as depicted in Fig. S2.† The reference molecule is altered by slight modifications on the terminal side with different acceptor groups connecting with spacers. These modifications are performed through the spacer groups. Therefore, all the molecules followed the acceptor-spacer–donor–spacer–acceptor combination. The reference BDTS-2DPP and all designed molecules (M1–M5) are optimised with the chosen functional MPW1PW91 to explore the opto-electronic properties.

3.2 UV-visible spectral characteristics

To explore more useful insight into DFT functions, the geometry of molecule R is optimised using a variety of approaches. In the solvent phase, the molecule R is optimised using different functionals such as B3LYP, MPW1PW91, CAM-B3LYP, and WB97XD. The values of different parameters like absorption profile (λ_{max}) , light harvesting efficiency (LHE), percentage assignments (S_0 – S_1), dipole moments (μ), excitation energy (E_X), oscillator strength (f) for gas phase and solvent phase are shown in Tables S1 and S2.† The change in λ_{max} values of all the designed molecules (M1-M5) in contrast with reference molecule R is due to the introduction of acceptor groups to the donor core of the material. All the designed molecules (M1-M5) have greater λ_{max} and are red-shifted in absorption spectra in contrast to the reference molecule R. The highest λ_{max} is shown by molecule M1 (846 nm) in dichloromethane (CH₂Cl₂). The λ_{max} values of all the designed molecules using dichloromethane came out to be M1 (846 nm), M2 (748 nm), M3 (710 nm), M4 (699 nm), and M5 (707 nm) respectively, and are shown in Fig. 1(a) and (b). As molecule M1 shows the maximum absorption strength of 846 nm and minimum band gap of 1.81 eV as depicted in Fig. S3[†] and tabulated in Table 1, hence it is the proficient donor material among all designed molecules.

3.3 The frontier molecular orbitals (FMOs)

When a bridge unit connects a donor and acceptor region, there is a less energy band gap that facilitates efficient charge transfer. The donor segment is positioned in the centre of the molecule, with end-capped acceptor segments located at the terminals representing two states of energy: one in which electrons are located and another in which they can be promoted. An important factor in a material's electrical and optical properties is its band gap. For efficient charge movement and light absorption, an ideal band gap is required. The linkage between the donor and acceptor units is established using thiophene bridges. 40 The band gap for reference molecule R is 2.24 eV, as reported by the FMOs, with HOMO energy of -5.05 eV and LUMO energy of -2.81 eV. When an electron density is excited, it moves from an electron donor to an electron acceptor region. The electron density of the HOMO and LUMO in reference molecule R is mostly concentrated on the whole molecule, including the donor, acceptor, and bridge fractions. The HOMO of the molecule M1 is formed by contributions from the donor,



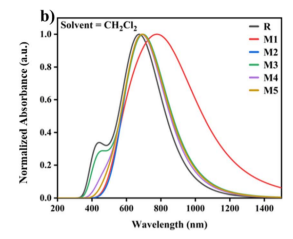


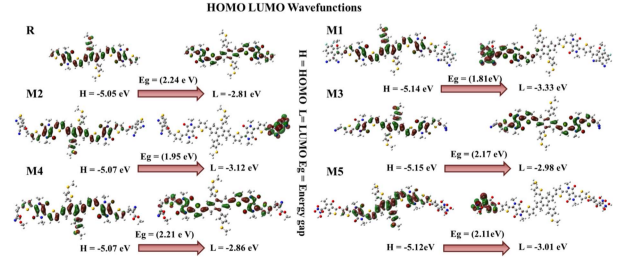
Fig. 1 Graphically illustration of UV-vis absorption of reference molecule R and designed molecules (M1-M5), in gas phase as well as in solvent

Table 1 The theoretically calculated HOMO-LUMO and energy gap $(E_{\rm g})$ values of reference molecule R and designed molecules (M1–M5) using the MPW1PW91/6-31G(d,p) level of theory

Molecules	E_{HOMO} (eV)	E_{LUMO} (eV)	$E_{\rm g}$ (eV)
R	-5.05	-2.81	2.24
M1	-5.14	-3.33	1.81
M2	-5.07	-3.12	1.95
M3	-5.15	-2.98	2.17
M4	-5.07	-2.86	2.21
M5	-5.12	-3.01	2.11

acceptor, and bridge. Conversely, the LUMO signal within the molecule M1 exclusively illuminates the acceptor bridge electron density. The electron density in the HOMO of molecules M2, M3, M4, and M5 is mostly generated by the donor and spacer units, with no contribution from the acceptor moieties in these molecules. Absorption of longer wavelengths of light is

possible when the energy gap is reduced. Having the capacity to absorb light in the red and near-infrared parts of the spectrum is a practical property of materials with a low energy gap. The energy levels in these zones are lower than those of blue or UV light. The LUMO of molecules M2 and M5 is determined by the electron density located on the acceptor moieties, whereas the LUMO of molecules M3 and M4 is influenced by the contribution of both the acceptor and the bridge unit. Fig. 2 illustrates that all the designed molecules have the capability to efficiently transmit charges. The E_g values of designed molecules M1-M5 and the reference molecule R are listed in Table 1, along with the HOMO and LUMO energies. M1, with its lowest bandgap (1.81 eV), has been selected as the most effective molecule among all molecules. The lowest bandgap is seen in molecule M1, suggesting that it may easily shift between the HOMO to the LUMO (Fig. S3†). Therefore, it may be inferred that M1 has the lowest band gap, as seen in Fig. S3.† It exhibits the smoothest transitions among all newly engineered molecules.



FMOs diagram of reference molecule R and all newly designed molecules (M1-M5)

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3.4 Density of state analysis (DOS)

The DOS gives information regarding the quantity of electronic states that are accessible at each energy level. The significance of this factor in OSCs lies in its direct influence on the creation of electron-hole pairs (also known as exciton) when light is absorbed, as well as their subsequent separation and the collection of charges at electrodes. In order to examine the electronic distribution and analyse the electron density contribution in both the conduction and valence bands, DOS calculations were conducted using the MPW1PW91/6-31G(d,p). The different colours in Fig. 3, are used to represent distinct donor and acceptor groups. Additionally, they provide the individual and collective contributions of each segment in the DOS spectrum. The elevated peaks in the DOS graphs, resulting from increased electron density, indicate a greater degree of attraction between the connected groups. The PyMOlyze-1.1 program was used to analyse DOS plots. Fig. 3 provides a clear picture about how the donor core transmits electron density to the acceptor units. The bridging units are what make this density transfer possible. The donor peaks exhibit strong intensity in the HOMO, whereas the acceptor intensity is notably high in the LUMO. This observation is most precisely seen in molecules M1, M2, and M5. The DOS spectra of the M1 molecule demonstrate the segregation of red peaks (representing the acceptor unit) from black peaks (representing the donor unit) in the LUMO region. This provides compelling evidence for the existence of a very potent acceptor unit in the M1 molecule. The examination of the spectra indicates that the primary factors that affect the transfer of charge involve the acceptor and donor components via the spacer group. The findings obtained from DOS spectra are consistent with the FMOs.

3.5 Reorganization energy (RE)

RE (λ) plays a vital role in defining the improved photovoltaic properties of materials designed for future applications in organic solar cells. It facilitates the investigation of the energies of electrons (λ_e) and holes (λ_h) during molecular excitations and has an inverse relationship with charge transfer. Determining the effect of molecular reorganization on the transport of charges, namely electrons and holes, inside organic solar cell materials is the main reason for calculating RE. The symbol λ can be separated into two components: λ_{int} , which specifies the interior geometries, and λ_{ext} , which defines the exterior polarisation environment. The value of λ_{int} can be determined using equations from Marcus' theory since the energies of anions, cations, and neutral molecules influence it. Marcus theory describes the rate at which an electron can transfer between two molecules, considering factors such as the reorganization energy (λ) and the driving force of the reaction (ΔG). The theory provides a relationship between the rate of electron transfer and these key parameters, predicting that the rate increases with decreasing reorganization energy and reaches a maximum at a specific point where the driving force equals the reorganization energy. The significance of Marcus theory in our study lies in its ability to explain the impact of molecular structure on electron transfer efficiency. By analyzing the reorganization energies of the molecules, we can predict their electron transfer rates and, consequently, their performance in photovoltaic applications. Lower reorganization energies, as suggested by Marcus theory, are indicative of more efficient charge transport, which is crucial for achieving higher power conversion efficiencies in organic solar cells.41 The reorganisation energies are shown in Table 2. RE for electron transfer of all the designed

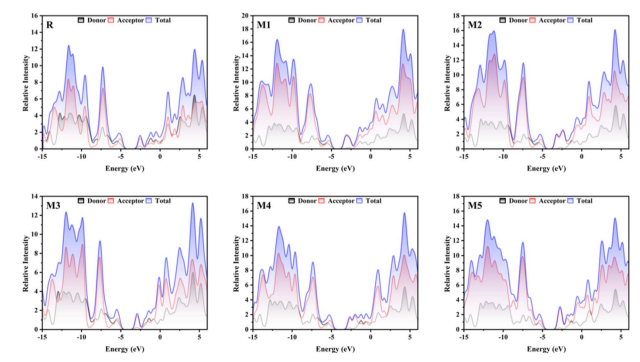


Fig. 3 DOS graphs for reference molecule R and designed molecules.

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Table 2 The reorganization energies of holes (λ_h) and electrons (λ_e) for the reference molecule R and designed molecules (M1-M5) using the MPW1PW91/6-31G(d,p) level of theory

Molecules	$\lambda_{ m h}$	λ_{e}
R	0.143	0.146
M1	0.185	0.113
M2	0.157	0.099
M3	0.164	0.078
M4	0.139	0.099
M5	0.119	0.108

molecules is lower than reference molecule R. Therefore, it is evident that all designed molecules exhibit excellent electron mobility. The molecule M3 has the lowest RE value for electron transfer, λ_{ext} indicating that it may effectively transfer charges. RE for electron transfer exhibits an ascending trend as: M3 < M2 = M4 < M5 < M1 < R.

3.6 Dipole moment (μ)

The dipole moment serves as a promising factor in assessing the material's solubility in organic solvents and plays a crucial role in enhancing the effectiveness of OSCs. 42 The energy levels of materials can be changed by molecular dipole moments, particularly the HOMO and LUMO values. This may improve energy balance at donor-acceptor interfaces in OSCs. If a material's polarisation is strengthened by a strong dipole moment, it is feasible to achieve a rise in the dielectric constant. This improved polarisation may lead to an increase in charge mobility and a lower recombination rate of separate charges (electrons and holes), hence decreasing the coulombic affinity between them. For the charges to be transferred between the materials and collected at the electrodes efficiently, this alignment is crucial. A direct relation between the dipole moment and solubility means that the higher the dipole moment of any material, the higher will be its solubility. Higher mobility improves molecular symmetry, which increase charge mobility and facilitates charge movement. The dipole moment of all designed molecules is determined by DFT-MPW1PW91/6-31G(d,p). Table 3 shows the μ values in the ground and excited states of the solvent. Molecule M1 shows the highest μ in both states i.e. ground state as well as in excited state, hence it will be the best molecule for its use in film formation because higher μ leads to enhanced crystallinity.⁴³ All the designed

Table 3 The dipole moments of reference molecule R and designed molecules (M1–M5) at the ground state ($\mu_{\rm q}$) and the excited state ($\mu_{\rm e}$) using the MPW1PW91/6-31G(d,p) level of theory

Molecules	$\mu_{ m g}$	$\mu_{ m e}$	$\mu_{ m e} - \mu_{ m g}$
R	3.143	3.942	0.799
M1	10.625	13.624	2.999
M2	4.211	5.688	1.477
M3	7.723	9.786	2.063
M4	4.041	4.228	0.187
M5	6.360	6.360	0.000

molecules show greater μ values than the reference molecule R. All of the designed molecules and the reference molecule R have μ values in the order of: M1 > M3 > M5 > M2 > M4 > R. Hence, M1 will show better photovoltaic stability as well as solubility in dichloromethane solvent.

Open circuit voltage (V_{oc})

PCE of OSCs is defined by a critical factor named open-circuit voltage. It is the highest voltage by a device when the device is present at zero current. A high $V_{\rm oc}$ requires an acceptor molecule with a high LUMO energy level and a donor molecule with a low HOMO energy level.44 The energy disparity between an acceptor material's LUMO and HOMO, as compared to the substance of the material being applied, is the primary determinant of a material's V_{oc} . To enhance charge separation and extraction, the position of these energy levels can be tuned to achieve a greater V_{oc} . All the designed molecules show properties of electron donation hence their HOMO values are correlated with some acceptor LUMO values. The acceptor used in the current work is PCBM. The equation used to find the values of open circuit voltage is as follows:

$$V_{\rm oc} = \left(\frac{\left|E_{\rm HOMO}^{\rm D}\right| - \left|E_{\rm LUMo}^{\rm A}\right|}{e}\right) - 0.3\tag{3}$$

In the above equation, energies are represented by E, the charge on the discrete molecule is represented by e, and 0.3 is the empirical factor. The $V_{\rm oc}$ values of reference molecule R and designed molecules are shown in Fig. 4. The $V_{\rm oc}$ values of reference molecule R and designed molecules (M1-M5) are 1.05 V, 1.14 V, 1.07 V, 1.15 V, 1.07 V and 1.12 V respectively. The molecular structures have variations in conjugation and the presence of electron withdrawing or electron donating groups, which directly affect these energy levels. For instance, molecules M1 and M3, which incorporate additional conjugated units or electron withdrawing groups, exhibit higher V_{oc} values. These structural modifications result in a larger energy gap between the HOMO and LUMO, thereby increasing the $V_{\rm oc}$ compared to the reference molecule R. All designed molecules have more $V_{\rm oc}$

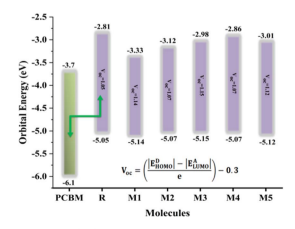


Fig. 4 $V_{\rm oc}$ values of reference molecule R and all designed molecules (M1-M5) calculated with regard to the PCBM.

in comparison to reference molecule R, hence they are better donor materials than reference molecule R. M3 exhibits the highest $V_{\rm oc}$ (1.15 V) value, hence it is the most efficient molecule among all designed molecules and it has the potential to serve as donor molecules for solar cells applications.

3.8 Transition density matrix (TDM)

The electron-hole and excitation motilities are effectively elucidated through a comprehensive analysis of TDM graphs. ⁴⁵ The charge density distribution in fragments as well as in associations between donor and acceptor subunits can be investigated using TDM graphs. ⁴⁶ The hydrogen atoms have been excluded because of their minor contribution. In order to analyse the TDM graphs properly, the fragments of our designed molecules have been generated *i.e.* A (end-capped acceptor groups) and D (donor core unit) as shown in Fig. 5. TDM graphs of reference molecule R and designed molecules are depicted in Fig. 5. In reference molecule R and designed molecules, the electron density resides on the donor unit and is distributed efficiently towards the acceptor units through the bridge unit. The charge transfers and exciton dissociation vary inversely with the interaction coefficient. The designed

molecule M2 exhibits better charge distribution hence, gives better results by the reference molecule R.

3.9 Exciton binding energy (E_b)

The exciton binding energy, which is the energy required to separate an electron hole pair into free charge carriers, plays a crucial role in determining the performance of photovoltaic materials. When the exciton binding energy is reduced, the probability of exciton dissociation increases, leading to a higher generation of free charge carriers. This can improve the $J_{\rm sc}$ and overall PCE of the material. Conversely, if the exciton binding energy is high, excitons are less likely to dissociate, which can result in significant recombination losses, thereby reducing the device's efficiency. The dissociation of exciton potential can be assessed by exciton binding energy denoted by $E_{\rm b}$. Both electron and hole pairs have a direct relation with $E_{\rm b}$. For that reason, it is the superior method to study the photophysical and optoelectronic characteristics of designed molecules. 32,48 The output of a device will be good if the E_b is lower. The lower E_b will consequently make the coulombic interactions lower as it has a direct relation and will give higher charge dissociation because charge dissociation is inversely related to the binding energy. $E_{\rm b}$ is calculated using the equation below⁴⁹

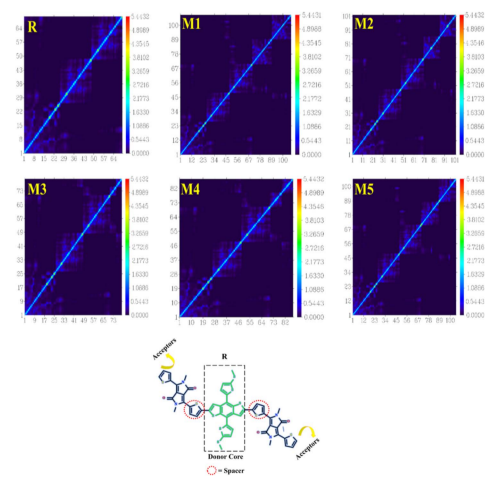


Fig. 5 TDM plots of reference molecule R and designed donor molecules (M1-M5) at the same level of theory.

$$E_{\rm b} = E_{\rm g} - E_{\rm opt} \tag{4}$$

where, $E_{\rm g}$ is the energy bandgap, and $E_{\rm opt}$ is the minimum energy required for the first transition. Table S3† displays the theoretically computed $E_{\rm b}$ of reference molecule R and each of the proposed molecules.

We observed that molecule M1 has a slightly lower binding energy compared to the reference molecule R, suggesting that the exciton in M1 is less tightly bound. This could facilitate better charge separation upon excitation, potentially enhancing device efficiency by increasing the generation of free charge carriers. Molecule M2 exhibits the lowest binding energy among the molecules studied, indicating the weakest exciton binding. This characteristic is generally favorable for photovoltaic applications, as it promotes the efficient dissociation of excitons into free carriers, which could significantly boost device performance. On the other hand, molecule M4 has the highest binding energy (0.44 eV), indicating very strong exciton binding. While this could lead to lower photovoltaic efficiency due to reduced free carrier generation, it might be advantageous in systems where controlled exciton diffusion is required before charge separation. Overall, the variation in exciton binding energy across the molecules demonstrates its significant impact on their photovoltaic performance, with lower binding energies generally promoting better charge separation and higher efficiency. As molecule M2 has the smallest E_b value out of the five molecules that were designed; hence, it improves charge separation more than reference molecule R and the other molecules. The increasing E_b order is as M2 < M1 < M5 < M3 = R < M4.

3.10 Non-covalent interaction (NCI)

Covalent bonds are used to examine the disposition of all bonded atoms inside the primary structure of a molecule. On the other hand, NCI like hydrogen bonds, aerogen bonds, chalcogen bonds, π - π and n- π * stacking, hydrophobic interactions, and π anion and cation are used to regulate aggregation, conformation, stabilisation, tertiary structure, and quaternary structures. 50,51 Non-covalent interactions are affected by the local environment created by coordinated atoms. These interactions help in maintaining the structure and functioning of molecules. 52,53 Steric hindrance, hydrogen bonds (HBs), and van der Waals interactions (vdW) are examined in the NCI plot.54 Kinetic energy and electronic densities are used for analysing NCI. NCI plots rely on electron density (ρ) and reduced density gradient (s). 55,56 ρ , and s combine to give an estimation of bonding regions. Low ρ and low s give NCI. Interaction can be repulsive or attractive according to the sign of Hessian eigenvalue (λ_2). This distinguishes steric hindrance or non-bonding interactions ($\lambda_2 > 0$) from bonding interactions. Cut-off values are necessary to obtain a 3D isosurfaces plot. Reduced-gradient Iso-value is referred to as the first cut-off. 57 Mapping ρ against s-isosurfaces results in two cut-off values referred to as ρ^- cut and ρ^+ cut. These cut-offs are required for setting the scale of colours. The blue scheme represents ρ^- cut (attractive) while the red-green-blue scale combined with red denotes ρ^+ cut (repulsive) as shown in Fig. 6. Electron density itself can differentiate hydrogen bonds from other intermolecular interactions. The peak of the hydrogen bond is observed at approximately $\rho \approx -0.05$. It appears as a blue localised NCI domain. Peaks appearing within the ρ^- region indicate weaker interactions and are depicted in green colour. These interactions lie in the centre of the NCI plot. Highly dense localised blue region of hydrogen bond shows strong intermolecular interaction as compared with wide regions of other weaker interactions.58 By default, RDG isosurfaces are kept at 0.5. A high-quality grid (0.016 Bohr) is used because the number of

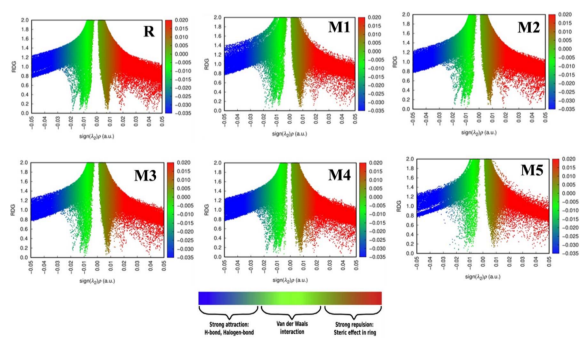


Fig. 6 Non-covalent interactions in reference molecule R and all designed molecules (M1–M5).

atoms in designed molecules is large and is necessary to obtain satisfactory graphical quality. In molecule R, a dense blue colour region indicates a hydrogen bond while a dense green region at -0.01 au indicates vdW interactions as depicted in Fig. 6. These interactions predominate over the scattered steric clashes which appear between 0.02 and 0.05 au. In M1, the less dense blue region shows slightly weak hydrogen bonds. Redcoloured steric clashes are much lesser when compared with the reference molecule R. Also, less dense green region indicates scattered vdW interactions. In M2, both steric clashes and vdW interactions are less dense where hydrogen bond predominates. M3 shows a similar trend to R in all three regions. There is a slight difference in the density of green region in M3. M4 shows a solid dense blue region indicating stronger hydrogen bonds. Regions of steric hindrance and vdW interactions are also scattered more. M5 shows a less dense blue region of hydrogen bonds. A less dense green region shows scattered vdW interactions. These interactions do not show a clear peak of vdW interactions. Steric clashes are also scattered and do not show clear peaks. If we compare these regions among reference molecules and all other designed molecules, M1 and M5 do not show dense blue regions of hydrogen bonds. There are many scattered areas of green region showing scattered vdW interactions. Reference molecule R and M4 show

a more similar region of steric clashes lying between 0.01 and 0.05 au. Reference molecule R and M3 show a similar region of vdW interactions. Hydrogen bonds in blue region dominate over all other interactions but this approach is not well defined in molecule M5 which shows much scattered regions. Reference molecule R, M1, M2, M3 and M4 show vdW interactions in the range of -0.01 to 0.02 au. There is no clear spike of these interactions in M5. In M1 and M5 more scattered vdW interactions are present. Fig. 6 shows all these interactions in reference molecule R and all designed molecules.

3.11 Electron density differences maps (EDDMs)

Essential for understanding the stability of molecules, the EDDM quantifies the electronic charge transfer. Disparities in the accumulation and decline in charge density in different regions may explain why the studied molecules exhibit different levels of intermolecular charge transfer (ICT). Reference molecule R and proposed molecules 2D EDD maps are highlighted in Fig. 7. The molecules under study have holes only and electrons at their acceptor and donor sites, respectively. Consequently, we used dichloromethane as a solvent to simulate EDD maps of the investigated compounds at the MPW1PW91/6-31G(d,p) at a theoretical level. The cubegen and cubeman tools from the G09 package were used for this analysis. Whereas different

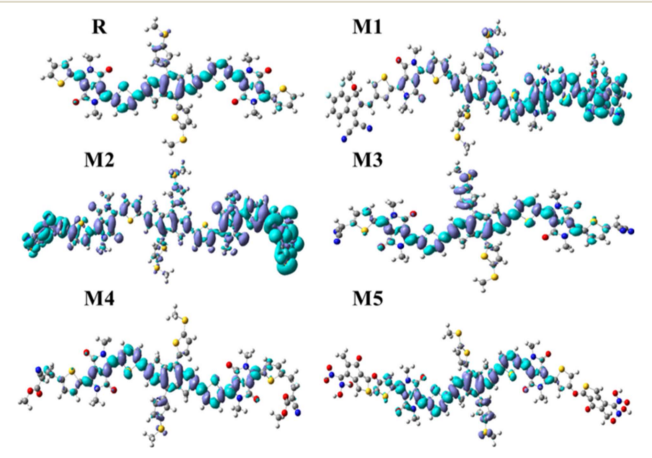


Fig. 7 Electron density differences maps of reference molecule R and designed donor molecules (M1–M5). All maps are constructed using the MPW1PW91/6-31G(d,p) level of theory at $A-\pi-D-\pi-A$ combination in dichloromethane solvent (purple = loose electronic cloud, cyan-blue = qain electronic cloud).

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colours (like cyan-blue and purple) appear on EDD maps, which represent different electron densities. The purple area represents a loss in electron density resulting from charge transfer, whereas the cyan-blue area represents an increase in electron density. The purple area acquires its density from the cyan-blue zone. All of the EDD map plots showed that there was constant ICT. All of these points to the same ICT *via* bridges between the donor core in the middle and the acceptor fragment at the end as the likely source of the red shift. So, the studied compounds might be used as efficient acceptor materials in organic photovoltaic cells, leading to better efficiency.

3.12 Fill factor (FF)

The FF plays a significant role in influencing the power conversion efficiency of an organic solar cell. FF is calculated by the following formula:⁵⁹

$$FF = \frac{\frac{eV_{oc}}{k_{B}T} - \ln\left(\frac{eV_{oc}}{k_{B}T} + 0.72\right)}{\frac{eV_{oc}}{k_{B}T} + 1}$$
(5)

where, V_{oc} , k_B , T, and e represent the open-circuit voltage, Boltzmann constant, the temperature at 300 K, and the elementary charge of a molecule respectively. Table 4 shows the calculated FF of all molecules. The calculated FF value of the reference molecule R is 88.66%. Furthermore, the estimated FF value of designed molecules ranges from 88.82% to 89.42%. It describes that the designed molecules exhibit a higher FF value than the reference molecule R. The planarity and rigidity of the molecular structures significantly influence π - π stacking interactions, which in turn affect charge mobility. Molecules M1 and M3, which show the highest FF values benefit from enhanced π - π stacking due to their more planar and rigid structures. These characteristics reduce charge recombination and allow for more efficient charge transport, leading to higher FF values. In contrast, molecules with less optimized stacking, like R, display slightly lower FF due to less efficient charge mobility. Higher FF values indicate the efficacy of newly designed molecules assembled as future solar devices. Hence, the FF increasing pattern is as follows an increasing order of R < M2 = M4 < M5 < M1 < M3. Among the designed molecules, M1 exhibits the second highest FF and PCE values which make it a suitable candidate to be used as a potential future solar cell device.

Table 4 Theoretically calculated value of open circuit voltage ($V_{\rm oc}$), fill factor (FF), and power conversion efficiency (PCE) of reference molecule R and designed molecules (M1–M5)

Molecules	$V_{\rm oc}$ (V)	FF (%)	PCE (%)
R	1.05	88.66	19.82
M1	1.14	89.33	21.69
M2	1.07	88.82	20.24
M3	1.15	89.42	21.90
M4	1.07	88.82	20.24
M5	1.12	89.18	21.27

3.13 Power conversion efficiency (PCE)

The PCE of a solar cell is measured by adding up all of its output results to see how well the material is made for practical purposes. The equation is provided below:

$$PCE = \frac{V_{\text{oc}} J_{\text{sc}} FF}{P_{\text{in}}}$$
 (6)

where $P_{\rm in}$ stands for the power of the incident rays. The $J_{\rm SC}$ is an experimental value of 21.3 mA cm⁻² obtained from previous research.60 Fig. S4† shows the graphical representation of the PCE value of all investigated molecules concerning the reference molecule. The increase in PCE value results from alterations at the terminal side, where various electron withdrawing groups are attached. These groups help disperse the electronic cloud between the donor core and the acceptor part of the molecule. The molecular structures that enhance $V_{\rm oc}$ and FF also contribute to higher PCE. Molecules like M1 and M3, with their extended conjugation and optimized electronic properties, show higher PCEs due to their ability to better harness light, efficiently separate charges, and transport them with minimal losses. The structural modifications in these molecules enhance their ability to absorb a broader spectrum of light and facilitate charge transport, leading to improved overall efficiency.

The M3 molecule has the highest estimated PCE value of 21.90%, followed by M1 with a value of 21.69%. Therefore, by making appropriate alterations to the reference molecule at the end, the new molecules exhibit the highest PCE, FF, and $V_{\rm oc}$ values compared to the reference molecule. These new molecules can be utilised in future solar devices to achieve the highest PCE value through practical application, as determined by experimentalists.

Conclusions

In summary, our study presents a comprehensive study into the molecular design and characterisation of A- π -D- π -A endcapped structured molecules for improving the photovoltaic efficiency of organic solar cells (OSCs). We have successfully designed five BDTS-2DPP-based donor molecules and employed theoretical approaches, including DFT and TD-DFT methods, to evaluate their performance in OSCs. Our study demonstrates that molecule M1 exhibits exceptional photovoltaic characteristics, including maximum absorption at 846 nm, and highest λ_e of 0.18 eV, with a narrow energy band gap of 1.81 eV. Furthermore, our findings reveal that M1 displays a high dipole moment in both the gas phase (10.62 D) and solvent phase (13.62 D) and efficient charge transfer capability between the HOMO and LUMO states with efficiency of up to 97%, making it a promising candidate for enhancing charge separation and transport in OSCs. Molecule M3 also shows potential with its high $V_{\rm oc}$ of 1.15 V, highest PCE value (21.90%) and highest fill factor (FF) value (89.42%), indicating its potential for efficient charge collection. The minimum excitation binding energy determined by molecule M2 ($E_b = 0.30$ eV) suggests a higher dissociation occurring during excitation. The high electron density seen in the newly formed M2 molecule when dissolved

in solvent is likely a result of its strong dissociation in a solvation environment. The comparison of the designed compounds with the reference molecule R reveals that all newly designed molecules (M1–M5) exhibit a reduced band gap, are red-shifted towards longer wavelengths relative to reference molecule R, and demonstrate remarkable charge motilities. Overall, our findings contribute to the advancement of organic photovoltaic and offer potential avenues of device engineering for the development of next-generation OSCs with improved efficiency and stability.

Data availability

Data for this article obtained from density functional theory (DFT) and time-dependent density functional theory (TD-DFT) methods, including geometry optimisation, density of states (DOS) analysis, natural transition orbitals (NTO), transition density matrix (TDM) calculations, and total energy computations, are available at Mendeley Data at https://doi.org/10.17632/64tf9v78n3.2.

Author contributions

S. Chang and F. Abbas conceived and directed the project; S. M. K. A. Naqvi, M. K. Shehzad, S. Bibi, and Y. Zhu prepared the models, carried out the simulations and prepared the figures; S. M. K. A. Naqvi, N. Alhokbany, Zahid Nazir, H. Long, R. B. Vasiliev and S. Chang analysed the data, interpreted the results and wrote the manuscript.

Conflicts of interest

No potential conflicts of interest have been disclosed by the authors.

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