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Novel A- π -D- π -A type non-fullerene acceptors of dithienyl diketopyrrolopyrrole derivatives to enhance organic photovoltaic applications: a DFT study

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To boost the photovoltaic attributes of organic photovoltaic cells, seven dithienyl diketopyrrolopyrrole (TDPP) donor-based A- π -D- π -A (acceptor-bridge-donor-bridge-acceptor) type molecules (TM1–TM7) were formulated by modifying the electron accepting ends of the reference molecule (TMR). Optical and quantum chemical parameters of seven synthesized molecules were investigated using density functional theory with the MPW1PW91/6-31G(d,p) functional. Several parameters that can be used to measure and improve the efficiency of solar cells have been analyzed and summed up. These parameters include binding energy of exciton, excitation energy of electron, reorganization energies, dipole moment, molecular electrostatic potential, charge mobility, wavelength of maximum absorption, open circuit voltage, short circuit current, fill factor, density of states, transition density matrices, as well as iso-surface and non-covalent interactions. Thus, all of our proposed structures are perceived to be superior to the reference in terms of the maximum possible solar energy yield in solar cells with bulk heterojunctions, as determined by analyses of our designed molecules for the aforementioned parameters.

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1. Introduction

Considering the need to prevent the energy resources of the globe from being depleted, and to replace non-renewable energy sources with useful inexhaustible energy sources that could provide relief to society from undesirable and disease-creating pollution, there should be an alternative way of bringing into use a clean, continuous and renewable energy source. There is enough energy in the sun to power the entire planet for a year with just one hour of exposure. In order to take advantage of the beneficial aspects of solar energy, we

endeavored to use this competent energy source in a way that would be sufficient to meet the energy needs of the entire world. Moreover the tunability, flexibility, mechanical adaptability, ease of preparation, wide range of absorption, and low weight of organic photovoltaic/solar cells (OPVs/OSCs) with bulk heterojunctions (BHJs) have attracted both the commercial and academic sectors.^{1–6} OSCs are susceptible to the incorporation of a variety of functional groups, which enables them to exhibit intriguing and novel optoelectronic properties.^{7,8} Due to their adaptable optoelectronic properties and increased efficiency of power conversion, organic photovoltaic and other such devices have garnered considerable interest.^{9–11}

Since their invention in the mid-19th century, OPVs have been employed to transform solar energy into electricity.¹² Initially, solar cells with thin layers of acceptor and donor were utilized, followed by solar cells with thick layers of acceptor and donor, but both of these were surpassed by BHJ solar cells in which both acceptor and donor layers are in maximum contact with each other, allowing for maximum interaction of both at a large interfacial area. In BHJ-type OSCs, the absorption of light generates a large number of electron–hole pairs that are tightly coupled to one another. These electron–hole pairs dissociate at the acceptor–donor interface and proceed towards their respective electrodes where they generate electricity.^{13–15}

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Due to their low reorganization energy and by association high mobility, fullerene acceptors (FAs) provide a remarkable 11% efficiency in OSCs. However, after discovering the drawbacks of FAs, their use has been avoided, and thus NFAs (non-fullerene acceptors) have been researched to overcome the drawbacks of FAs. Their assets include the outstanding tunability of E_g (band gaps), superior light-absorbing properties, thermal and photochemical stability, and increased lifetime. Furthermore, they have enhanced open-circuit voltages, reproducibility, as well as fill factor.¹⁶ Therefore, numerous attempts have been made to swap FA-based OSCs with NFA-based OSCs, such as BTP-Cl, ITC-2Cl, *etc.*, which are currently being developed and utilized.^{17,18} In this regard, un-fused NFAs are acting as the premier contributors due to their two- to four-step and cost-effective synthetic routes, as well as high yield as opposed to their counterparts, the fused NFAs.¹⁹

The TDPP-RDN compound has already been engineered and scrutinized to evaluate the effectiveness and performance of OPVs.²⁰ We chose this compound as our reference standard for the following reasons: it has a terminal functional group that accepts electrons and a dithienyl diketopyrrolopyrrole (TDPP) core. The TDPP-RDN electron mobility reached $0.07 \text{ cm}^2 \text{ V}^{-1} \text{ s}^{-1}$. Diketopyrrolopyrrole (DPP), which possesses good planarity, strong intermolecular contact, and good charge carrier mobility, served as the fundamental component (building block).^{21,22} The levels of energy for the molecule were evaluated. Using cyclic voltammetry, the calculated energies of the highest occupied (HOMO) and lowest unoccupied (LUMO) molecular orbital for TDPP-RDN were -5.39 eV and -3.85 eV , respectively. In combination with cyclic voltammetric data, the calculated HOMO and LUMO energy levels for TDPP-RDN were -5.61 eV and -3.83 eV , respectively. Maximum absorption for the TDPP-RDN molecule was measured at 710 nm in chlorobenzene, with a band gap E_g of 1.54 eV . Due to the lack of a single crystal structure, the exact molecular packing and arrangement of TDPP-RDN in thin films is unknown.

Seven photovoltaic compounds, designated **TM1–TM7**, were formulated by performing side-chain engineering of different acceptor groups at the peripheries of the A- π -D- π -A type²³ reference **TMR**. These acceptor groups were selected for their prominent attributes in various computational studies and are known to have high electron-withdrawing abilities, along with significant conjugation in them, which aids in effective transfer of charge density towards them.²⁴ The acceptors utilized were 2-(5,6-difluoro-2-methylene-3-oxo-indan-1-ylidene)malononitrile (**TM1**), 1-dicyanomethylene-2-methylene-3-oxo-indan-5,6-dicarbonitrile (**TM2**), 6-cyano-1-dicyanomethylene-2-methylene-3-oxo-indan-5-carboxylic acid methyl ester (**TM3**), 1-dicyanomethylene-2-methylene-3-oxo-indan-5,6-dicarboxylic acid dimethyl ester (**TM4**), 2-(2-methylene-3-oxo-2,3-dihydrocyclopenta[*b*]naphthalen-1-ylidene)malononitrile (**TM5**), 2-(6,7-difluoro-2-methylene-3-oxo-2,3-dihydrocyclopenta[*b*]naphthalen-1-ylidene)malononitrile (**TM6**), and 2-(1-chloro-5-methylene-6-oxo-5,6-dihydrocyclopenta[*c*]thiophen-4-ylidene)malononitrile (**TM7**). When contrasted to **TMR**, these newly developed structures showed marked improvements in charge

mobility, band gap reduction, and maximum absorption. Utilizing these acceptor moieties broadened the spectrum of absorption. Thiophene spacers also facilitated the transfer of charge from the central core to terminals (electron-accepting moieties). This suggests the active layer of molecules that have been formulated to be more efficient than the reference molecule **TMR** which could be beneficial to OSCs.

2. Computational details

All of the molecular structures under consideration were drawn using ChemDraw. Quantum mechanical computations of all the designed molecules and the **TMR** reference compound were carried out using Gaussian 09 software.²⁵ To build, as well as to display, the 3D assemblies of all optimized compounds, we utilized GaussView 6.0.16.²⁶ The in-depth investigation of **TMR** was done with DFT computing with its sub-method TD-DFT (time-dependent DFT), which was engaged to study the excited state properties. To execute the λ_{max} (wavelength of maximum absorption), we used multiple approaches: B3LYP,²⁷ CAM-B3LYP,²⁸ MPW1PW91,²⁹ and ω B97XD²⁵ with 6-31G(d,p) basis set, in solvent phase. Spectral depiction of λ_{max} was achieved through Origin 6.0 software.³⁰ UV-visible absorption^{31,32} of **TMR** was contrasted with the cited (710 nm in chlorobenzene)³³ value to determine the best functional. Closeness to the experimentally determined maximum absorption of **TMR** was achieved by MPW1PW91/6-31G(d,p). Using the IEFPCM (integral equation formalism polarizable continuum model), the maximum absorption characteristics adjoining the solvent were analyzed.³⁴ Multiwfn 3.7 (ref. 35) and PyMolyze 1.1 (ref. 36) were used to obtain the spectra for the transition density matrix (TDM) analysis and the density of states (DOS), respectively.

The most notable factor, *i.e.* reorganization energy (RE), which influences the working efficiency of solar cells is also needed to be measured. The overall RE is distributed into two forms: internal (λ_{int}) and external (λ_{ext}) RE. λ_{ext} is related to the external deformations in the molecule, and is excluded because of the constant external environment of chloroform,³⁷ whereas λ_{int} is related to changes that take place in the internal environment, *i.e.* variations in geometry/structure.³⁸ We solely looked at the energy required for internal rearrangement in order to analyze charge transfer through electron–hole mobility. For the molecules, the Marcus rate equation³⁹ has been employed to quantify the electron and hole reorganization energy:

$$\lambda_e = [E_0^- - E_-] + [E_0^0 - E_0] \quad (1)$$

$$\lambda_h = [E_0^+ - E_+] + [E_0^0 - E_0] \quad (2)$$

E_0^- and E_0^+ are anionic and cationic energies used in these equations to describe the ground state energy of a neutral compound. At a single-point ground state, E_0 represents the energy of a neutral molecule, whereas E_- and E_+ reflect the anion and cation single-point ground states energies after optimization. E_0^0 and E_0^0 are the ground-level energies for anions and cations, correspondingly, after optimization.



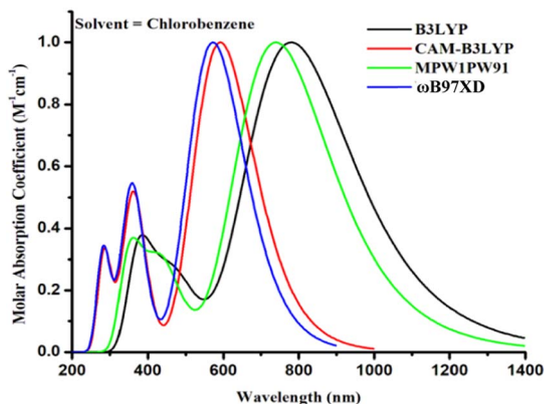


Fig. 1 UV-visible absorption of TMR for four different methods in chlorobenzene.

3. Results and discussion

3.1. Method selection and geometry optimization

The aforementioned four alternative and valuable computational approaches with a 6-31G(d,p) basis set were used to optimize the ground state geometry of **TMR**. The reason for using these functional is to investigate the functional that could lead to desirable results in terms of λ_{max} . Graphs depicting this behavior can be seen in Fig. 1(a) and (b). The MPW1PW91/6-13G(d,p) method provided an absorption maximum at 739 nm that was in close correlation with the experimental value (710 nm in chlorobenzene) as mentioned in the literature.³³ Therefore, this approach was chosen as the optimum method for further research of all the formulated compounds. In Fig. 2 and 3, the chemical structures and optimized geometries of the reference **TMR** and designed compounds are illustrated.

The bond lengths and dihedral angles (θ) of **TMR** and **TM1–TM7** are presented in Table 1. All of the optimal structures of the investigated molecules (**TMR** and **TM1–TM7**) were found to have comparable geometries, with bond lengths and dihedral angles within a reasonable range. All the molecules studied had bond lengths between 1.418 Å and 1.424 Å, with the C–C single bond being 1.54 Å and the C=C bond being 1.33 Å, which exhibited enhanced charge transfer properties and also enabled conjugation through delocalization of π -electrons. All of the investigated molecules had dihedral angles within the range of -0.653° to 0.007° , indicating the absence of any twisting between the non-fused thiophene bridges and the peripheral acceptors, thereby indicating that all structures have planar geometry.

3.2. Quantum chemical descriptor

Solar cells can be evaluated in general by analyzing the HOMOs and LUMOs derived from the optimal structures of the molecules under investigation. The collective name for these orbitals is “frontier molecular orbitals” (FMOs).⁴⁰ As we are familiar with transfer of electrons from HOMO to LUMO, this gives a clear indication that HOMO resides on the electron-donating portion

of the molecule, whereas LUMO resides on the electron-accepting portion.^{41,42} In other words, the LUMO is in the vicinity of the electrophilic part of the molecule, while the HOMO is close to the nucleophilic part.

Fig. 4 displays the FMO charge distribution. Here, the red color is an indicator of a constructive (positive) phase, while green color shows a destructive (negative) one. Our reference **TMR** and **TM1–TM7** molecules exhibited LUMO energies of -3.84 eV, -4.07 eV, -4.56 eV, -4.29 eV, -4.05 eV, -3.91 eV, -4.03 eV, and -4.08 eV, correspondingly. Whereas HOMO energies were -5.88 eV, -5.94 eV, -6.34 eV, -6.09 eV, -5.90 eV, -5.75 eV, -5.86 eV, and -5.91 eV, correspondingly (Table 2). Almost all the proposed compounds have lower HOMO and LUMO energies than the reference (**TMR**) due to the presence of strong peripheral electron-accepting entities. The FMO values of **TM2** are shown to be the lowest among all the molecules in both the ground and excited states compared to those of the **TMR** molecule. $\text{TM5} > \text{TM6} > \text{TMR} > \text{TM7} > \text{TM1} > \text{TM3} > \text{TM2}$ and $\text{TMR} > \text{TM5} > \text{TM6} > \text{TM4} > \text{TM1} > \text{TM7} > \text{TM3} > \text{TM2}$ are the orders of energy levels of HOMO and LUMO, respectively. The band gap (eqn (3)) plays a decisive role in maximizing charge transmission over the whole molecule and the molecule's overall efficiency. Band gap and power conversion efficiency (PCE) of OSCs are inversely correlated, as a molecule with a lower energy gap results in higher PCE and *vice versa*.

$$E_g = E_H - E_L \quad (3)$$

Here E_H and E_L are HOMO and LUMO energies, respectively. The minimum band gap can be achieved by increasing the HOMO of the donor, while decreasing the LUMO. The energy-gap values for the reference compound **TMR** along with **TM1** to **TM7** chromophores were 2.04 eV, 1.87 eV, 1.78 eV, 1.80 eV, 1.85 eV, 1.84 eV, 1.83 eV, and 1.83 eV, respectively (Table 2). All the planned structures have narrower band gaps than the standard compound **TMR**. The E_g values for all compounds fall in the range of 2.04–1.78 eV, as shown in Fig. 4. The compounds have band gaps in the following order: $\text{TMR} > \text{TM1} > \text{TM4} > \text{TM5} > \text{TM6} = \text{TM7} > \text{TM3} > \text{TM2}$. The results show that the presence of the most electron-deficient units (acceptors) enhances charge mobility by decreasing the energy-gap values between the FMOs.^{43,44}

The HOMO states of **TMR** and **TM1–TM7** have almost the same degree of charge consistency in both the spacer and donor segments. Acceptor moieties had a small amount of charge consistency, and the substituted methyl groups in the central donor fragment did not experience electron density dispersion, due to their vertical alignment with respect to the molecules' horizontal plane. While the electron density is uniformly distributed in the donor, spacer as well as acceptor regarding the LUMOs of **TMR**, **TM1**, **TM2**, and **TM7**, with the exception of the methyl substituents in the acceptors and donor (core) section, the LUMOs of **TM3**, **TM4**, **TM5**, and **TM6** revealed greater charge consistency over the donor core and spacer fragments but reduced charge consistency over the electron-accepting entities at the end-capped portions of the molecules. **TMR**, in contrast, shows almost equivalent electronic



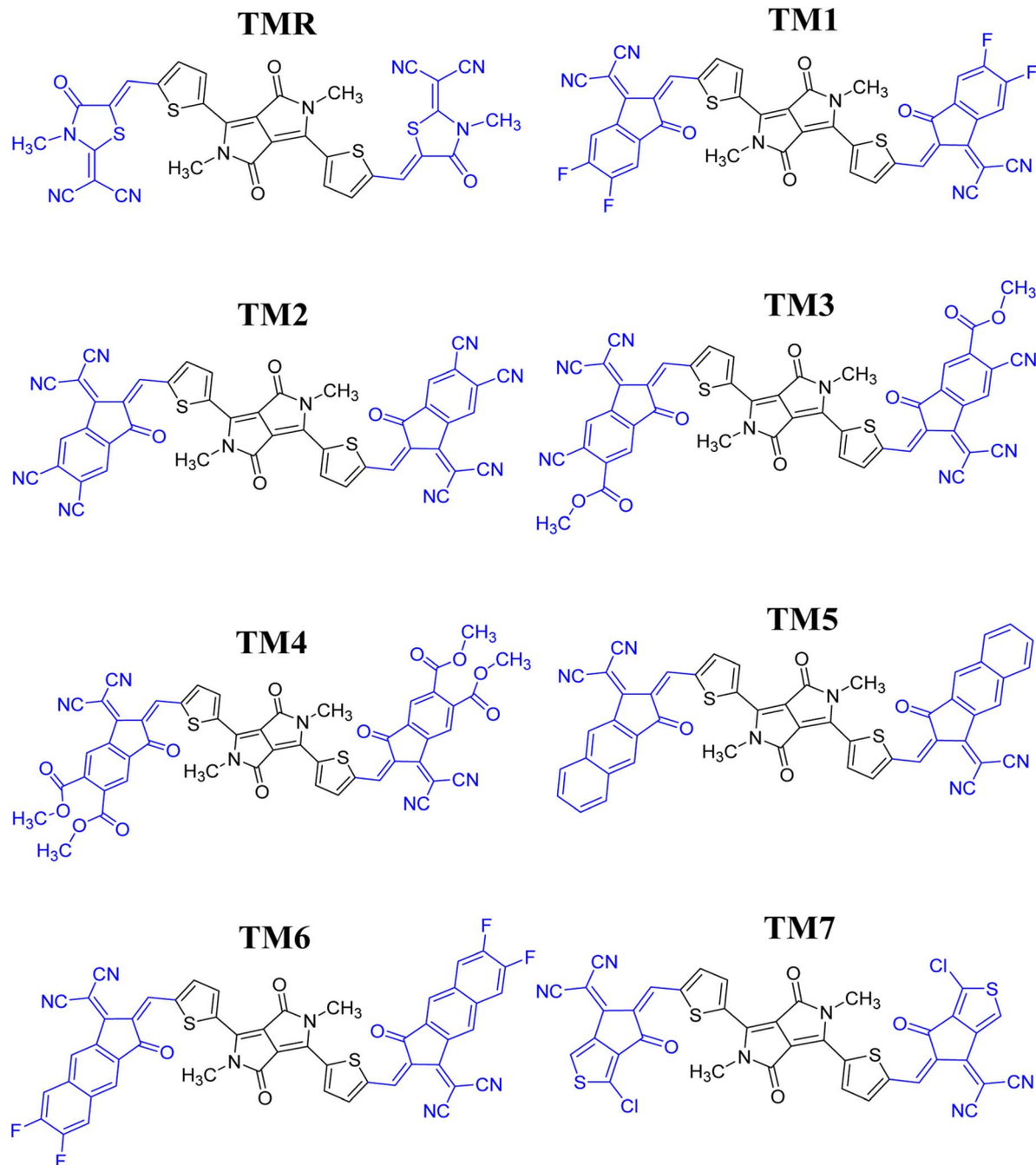


Fig. 2 ChemDraw structures of reference TMR and TM1–TM7 chromophores.

transitions between FMOs for all atoms excluding those in the methyl substituents. Because of the high density of charges in the LUMO of the acceptor and the HOMO of the donor, **TM1** and **TM7** are the most efficient in the interface of the OSC active layer. Also, the minute degree of charge density in the LUMO of donor and HOMO of acceptor could be due to the fairly planar geometry of the molecules.

3.3. Ionization potential and electron affinity

Ionization potential (IP) and electron affinity (EA) are useful tools for analyzing the charge-transfer efficiency of solar cells.

The former is the energy necessary to eject an electron from an atom's outermost orbit, while the latter is the energy given by an atom when an electron is introduced to its outermost shell.⁴⁵ But why we are discussing both of these in solar cells? This is for the reason that these variables work together in alternative ways to enhance the PCE of OSCs. Here IP values correspond to the energy needed by the HOMO in order to transfer its electron to the LUMO, hence they should be minimized so that electron will be removed with greater ease. EA, on the other hand, is linked to the LUMO values and so has to be greater. As groups that tend to donate electrons destabilize the HOMO by



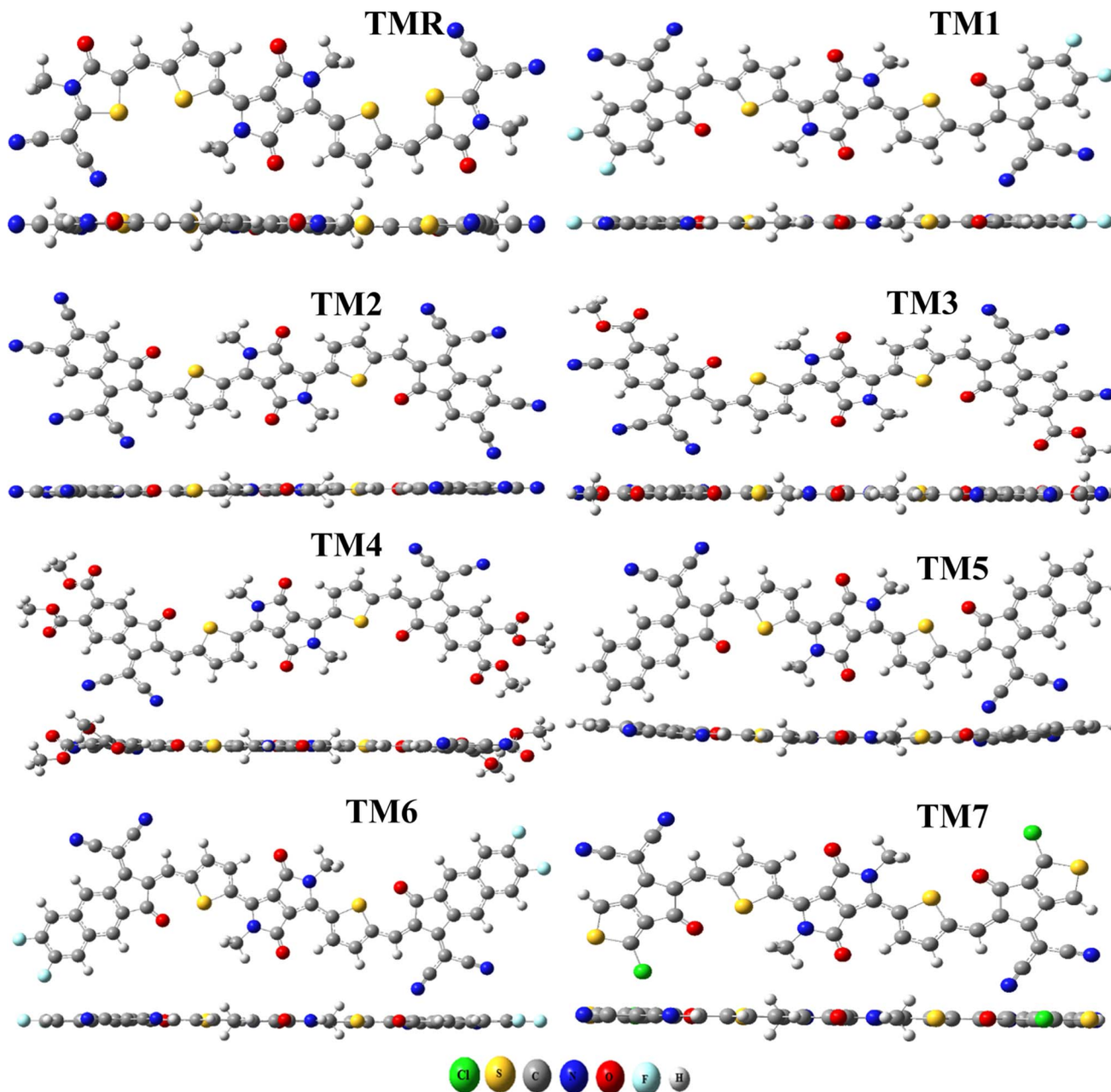


Fig. 3 Structural representation after optimization of TMR and tailored compounds TM1–TM7.

Table 1 Computed bond length and dihedral angle values for TMR and the modified chromophores TM1–TM7

Molecule	Bond length (Å)	Dihedral angle (θ)
TMR	1.424	−0.007
TM1	1.422	−0.003
TM2	1.418	0.007
TM3	1.420	−0.002
TM4	1.421	−0.428
TM5	1.422	−0.653
TM6	1.421	0.001
TM7	1.422	0.004

promoting electron transport, molecules with low IPs and high EAs are considered as good for passing charges. In contrast, the IPs of compounds with strong electron-withdrawing groups are higher, as the HOMO is more stable and electron removal is rather difficult. In accordance with Koopman's theorem^{46,47} the IP and EA of all the molecules under consideration were determined through eqn (4) and (5),⁴⁸ and the outcomes are presented in Table 2.

$$\text{IP} = [E_0^+ - E_0] \quad (4)$$

$$\text{EA} = [E_0 - E_0^-] \quad (5)$$



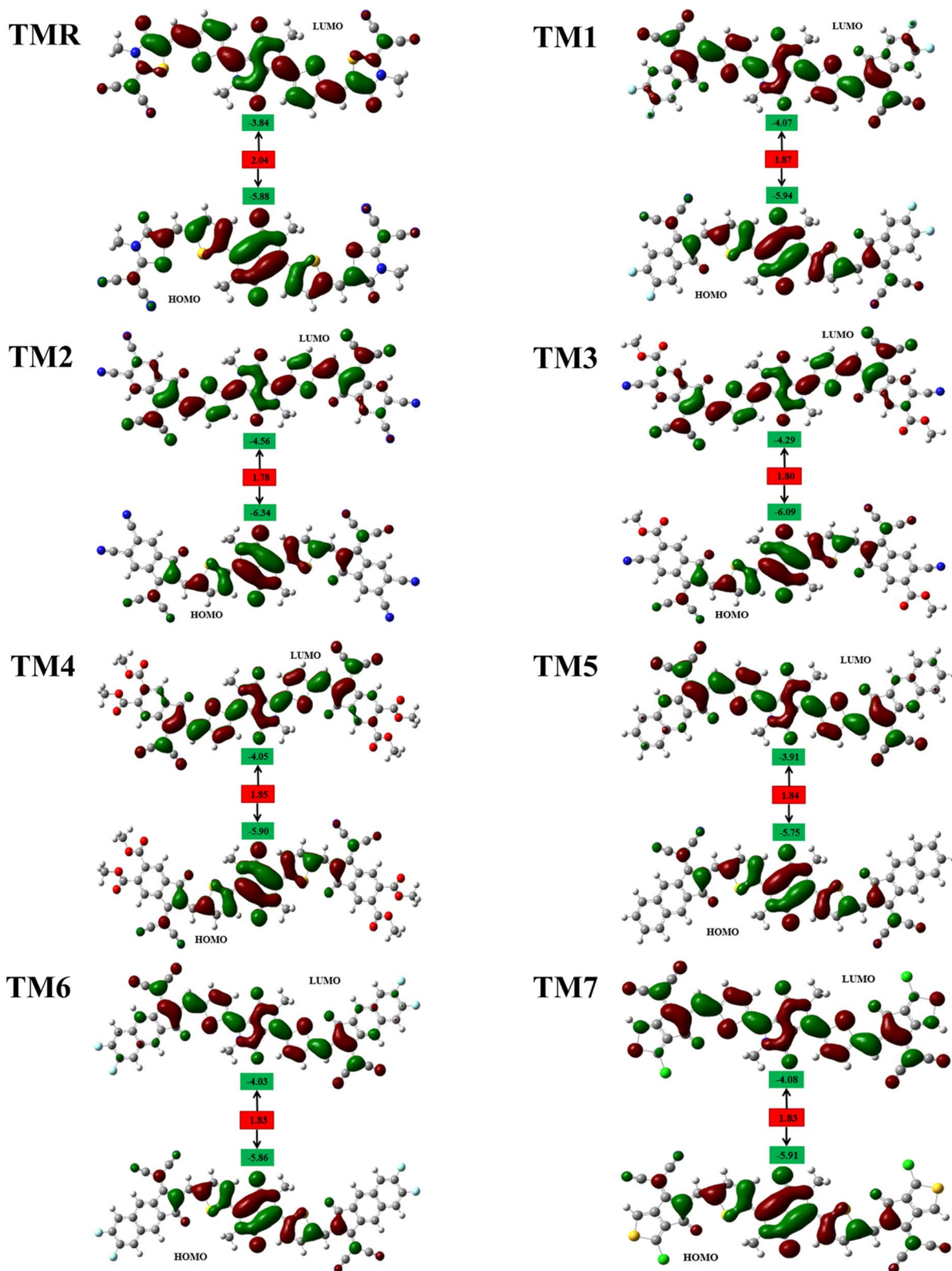


Fig. 4 FMO representation of TMR and tailored compounds TM1–TM7.

Here, IP denotes ionization potential and EA symbolizes the electron affinity of the system. The neutral molecule has cation with energy that is characterized *via* E_0^+ and anion with energy

which is characterized *via* E_0^- . The energy of a neutral molecule is denoted by the symbol E_0 .⁴⁰ Table 2 displays the calculated IP as well as EA values for the compounds under study. Almost all



Table 2 Computed values for HOMO, LUMO, E_g , IP, and EA of TMR along with TM1–TM7 in eV

Molecule	HOMO	LUMO	E_g	IP	EA
TMR	−5.88	−3.84	2.04	6.65	3.15
TM1	−5.94	−4.07	1.87	6.67	3.42
TM2	−6.34	−4.56	1.78	7.04	3.93
TM3	−6.09	−4.29	1.80	6.80	3.67
TM4	−5.90	−4.05	1.85	6.60	3.43
TM5	−5.75	−3.91	1.84	6.45	3.29
TM6	−5.86	−4.03	1.83	6.56	3.42
TM7	−5.91	−4.08	1.83	6.64	3.43

of the optimized molecules were found to have higher charge transfer efficiency than **TMR**. This was determined by the findings that almost all IP values for the optimized molecules were lower than those for **TMR**. **TM5** has the lowest IP (6.45 eV) and highest charge extraction from HOMO. The IP value of the **TM2** molecule is high indicating the poor charge transfer. Each of the synthesized molecules **TM1–TM7** also had higher electron affinity than the standard reference molecule **TMR**.

3.4. Photovoltaic properties

The absorptive characteristics of the molecules in the gaseous as well as solution states were estimated using the pre-set functional and basis set at TD-DFT. The absorption spectra⁴⁹ of molecules in both phases under investigation were acquired with the help of Origin 6.0, allowing us to clearly distinguish between the values of maximum absorption of the various compounds under research³⁰ as shown in Fig. 5(a) and (b). The molar extinction coefficient was standardized on the y-axis with wavelength on the x-axis. Additionally, bar graphs comparing the experimental and the calculated values of λ_{\max} are shown in Fig. 5(a) and (b) to visualize the discrepancy between the two sets of results. Maximum absorption, oscillator strengths, excitation energies, light-harvesting efficiencies, *etc.* are shown in Tables 3 and 4.

The absorption maxima of **TMR** and the tailored compounds **TM1–TM7** were 690 nm, 757 nm, 799 nm, 787 nm, 770 nm, 771 nm, 778 nm, and 771 nm in gas phase and in the presence of the chlorobenzene solvent these were 739 nm, 820 nm, 878 nm, 861 nm, 840 nm, 835 nm, 842 nm, and 837 nm, respectively. All of the tailored compounds showed a clear red shift in absorption relative to **TMR** in gas phase, and significantly in solvent (near IR) as the polar excited state was stabilized by the polar solvent. In Fig. 5(a) and (b), even if the calculated absorption value of all the designed molecules has shifted to a greater extent in both gas phase and solvent, maximum wavelength is given by solvent, which is owing to higher conjugation in the solvent phase. This increased conjugation may result from the compounds' increased solubility in the solvent. The tightly conjugated and electron-accepting component of **TM2** is responsible for its highest λ_{\max} (799 nm and 878 nm in both examined phases). In the gaseous state, the sequence of absorption is **TM2** > **TM3** > **TM6** > **TM5** = **TM7** > **TM4** > **TM1** > **TMR**, while in the solvent (chlorobenzene) phase,

the sequence is **TM2** > **TM3** > **TM6** > **TM4** > **TM7** > **TM5** > **TM1** > **TMR**.

The excitation energy (E_x) is a crucial determinant of whether or not a compound can be used in OPVs. The energy utilized by an electron to jump from its ground state (S_0) to its excited state (S_1) is denoted by E_x .⁵⁰ When E_x is small, it becomes easier for an electron to be excited, and for charge to move smoothly.⁵¹ The ability of the proposed molecules to efficiently transport electrons is revealed by a decrease in the E_x values of **TM1–TM7** in both examined phases in contrast to **TMR** as presented in Tables 3 and 4. The smallest value for **TM2** of first excitation energy, narrower band gap (1.78 eV), and greatest absorption maximum of 799 nm in gas phase and 878 nm in solvent indicated that the molecule has a higher charge transport ability. This is because of the existence of a potent end-capped electron-withdrawing acceptor entity. Because of its low E_x (1.54 eV in the gas phase and 1.41 eV in the solvent phase), **TM2** has the potential to serve as an active layer in efficient OSCs.

Oscillator strength (f_{os})⁵² is the probability of electronic transition from the ground HOMO levels to the excited LUMO levels in photovoltaic cells. Higher absorption in the UV-visible zone means a higher f_{os} , which in turn shows a greater charge transfer rate and hence a stronger oscillator strength. **TMR** and the constructed chromophores **TM1–TM7** had oscillator strengths in the range of 1.97–1.67 in gas phase and 2.18–1.87 in solvent (Tables 3 and 4). The **TM5** and **TM6** structures with their strongest electron-deficient acceptor units at the terminal position have highest f_{os} and as a result enhanced absorption and charge transformations.

Light harvesting efficiency (LHE) measures the ability of a molecule to absorb photons. According to the following eqn (6),⁵³ its value is exactly proportional to the oscillator strength (f_{os}):

$$\text{LHE} = 1 - 10^{-f} \quad (6)$$

Here, LHE represents light harvesting efficiency, and f represents the oscillator strength, describing the relationship between the two variables. We make use of solvent-phase oscillator strength to evaluate LHE. High levels of LHE need a robust oscillator. The f_{os} has no units. Due to their more planar conformation, **TM5** and **TM6** displayed the greatest f_{os} values in both phases (0.89 and 0.93 respectively), as seen in Tables 3 and 4.

3.5. Dipole moment

Dipole moment (μ) is another valuable parameter for assessing the effectiveness of solar devices. It arises in a molecule due to the difference in electronegativity of atoms contained in it. It is directly linked to the molecule's polarity.⁵⁴ According to the fundamental tenet that "like dissolve like," this increase in polarity results in better solubility in a polar solvent (for example, chlorobenzene). In contrast, symmetry results in a reduced μ , which in turn decreases solubility.⁵⁵ An increase in dipole moment also has the benefit of increasing the electron mobility amongst HOMO and LUMO. Thus in this way the reactivity of a molecule increases which in turn increases the



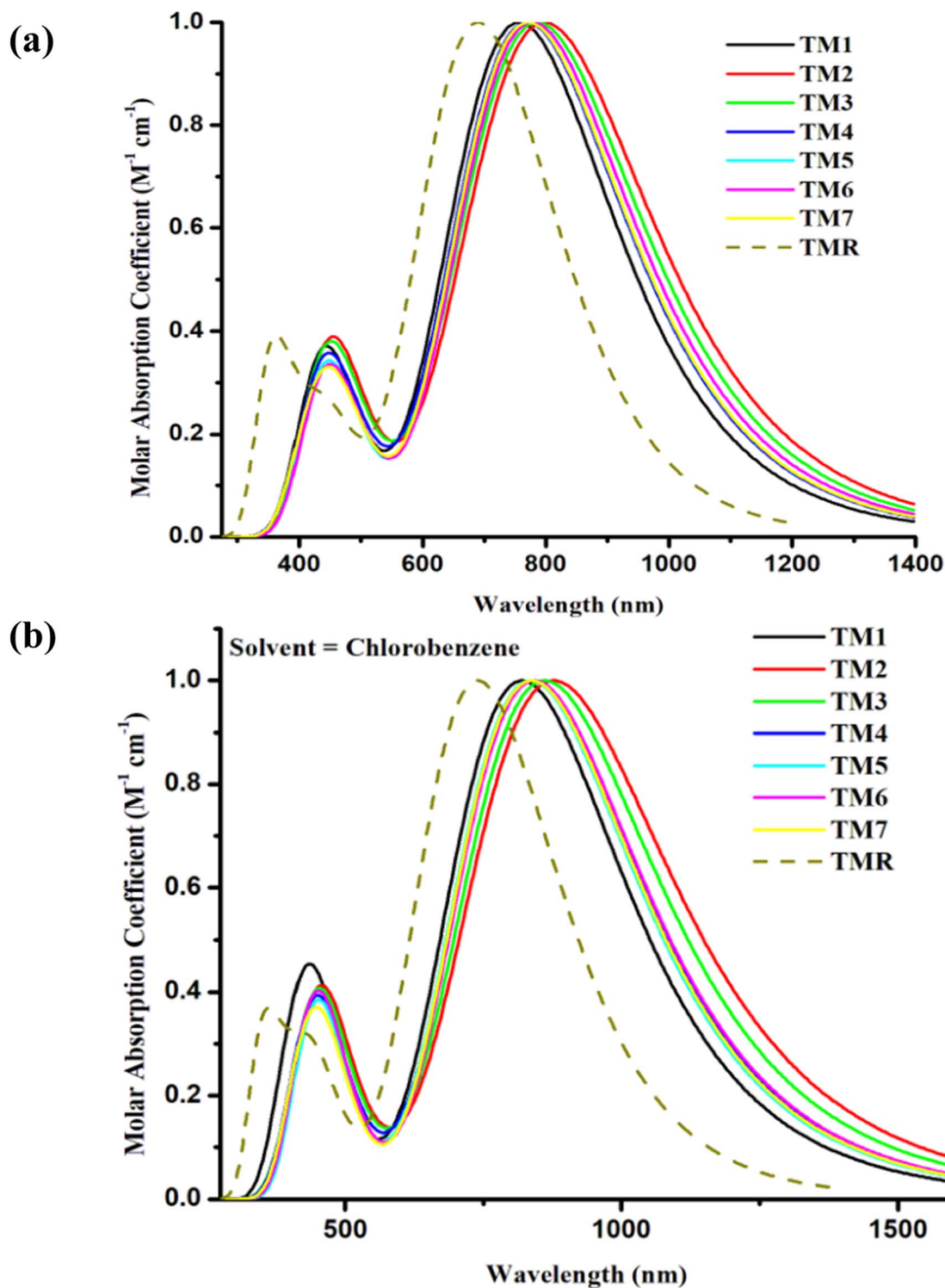


Fig. 5 Comparison plots for UV-visible absorption of TMR and tailored compounds TM1–TM7 in (a) gas phase and (b) chlorobenzene.

PCE of solar cells. Furthermore, the acceptors used in this research work are known to have a prominent effect on the dipole moment. The two fluoro-substituted malononitrile (TM1) groups significantly enhance the dipole moment due to enhanced π - π stacking found in them.⁵⁶ Similarly, the acceptors of TM3 due to their polar character tend to increase the

dipole moment as well.⁵⁷ However, it should be kept in mind that the dipole moment of any symmetrical molecule despite a strongly electron-withdrawing group can reduce significantly due to the cancellation of opposite poles.⁵⁸ However, regardless of their prominent symmetry, all the molecules except TM5 demonstrate better dipole moment than TMR. Table 5 displays

Table 3 Estimated results of λ_{max} , LHE, excitation energy, oscillator strength, and transition character of TMR together with TM1–TM7 in the gas phase

Molecule	Calculated λ_{max} (nm)	E_x (eV)	f_{os}	LHE	Assignment	Transition character (H \rightarrow L)
TMR	690	1.80	1.67	0.79	H > L	71%
TM1	757	1.64	1.76	0.83	H > L	71%
TM2	799	1.55	1.81	0.85	H > L	71%
TM3	787	1.57	1.83	0.85	H > L	71%
TM4	770	1.61	1.86	0.86	H > L	71%
TM5	771	1.61	1.97	0.89	H > L	71%
TM6	778	1.59	1.97	0.89	H > L	71%
TM7	771	1.61	1.78	0.83	H > L	71%

the dipole moment of the reference TMR and modified molecules TM1–TM7 in the gaseous state μ_g , in the solvent (chlorobenzene) μ_e , and their difference $\Delta\mu$. Both of the examined states (solvent and gas) have almost comparable order of dipole moments: TM5 > TM2 > TM6 > TM3 > TM4 > TM1 > TM7 > TMR and TM5 > TM3 > TM6 > TM2 = TM4 > TM1 > TM7 > TMR, correspondingly. The highest dihedral angle and the somewhat out-of-plane geometry of TM5 are responsible for its enhanced dipole moment. Because it is seen that dipole moment greatly depends on asymmetry, the lower the symmetry the greater the dipole moment.⁵⁹ Similarly, TM2 has a nitrogen along with oxygen and TM6 has fluorine and oxygen, which increase their μ . These findings suggest that the modification of molecules with strong acceptor fragments may be advantageous in the future for more efficient OSCs.

3.6. Density of states

The DOS for each of the compounds under study were calculated and analyzed to verify the results of FMO analysis. Donor cores, acceptors, and spacers are represented in the DOS plots by blue, red, and green lines, correspondingly (Fig. 6). These lines indicate the extent of involvement of various fragments in FMO enhancement. The entire DOS is shown as a black line. The HOMO is represented by the peak to the left of the central planar area, while the LUMO is represented by the peak to the right.⁶⁰ This core planar empty area is essentially the band gap between the FMOs.²⁴

Table 5 Computed values for gas-phase μ_g as well as solvent-phase μ_e dipole moment along with their difference $\Delta\mu$ in Debyes, for TMR and TM1–TM7

Molecule	μ_g	μ_e	$\Delta\mu$
TMR	0.0005	0.0005	0.0000
TM1	0.0009	0.0011	0.0002
TM2	0.0011	0.0039	0.0028
TM3	0.0018	0.0019	0.0001
TM4	0.0011	0.0013	0.0001
TM5	0.8102	0.9807	0.1705
TM6	0.0017	0.0020	0.0003
TM7	0.0008	0.0009	0.0001

Table 6 displays the results of a quantitative analysis of the three examined parts. It was discovered that the donor core made significant contributions towards the HOMO's improvement, spacers showed moderate and acceptors least involvement in HOMO elevation in the case of reference TMR and constructed structures TM1–TM7. For all our designed molecules (TM1–TM7) the percentage of donor involvement in the HOMO is greater than that for TMR. The results of FMO and the DOS are highly consistent with one another; the percentage of acceptors participating in the LUMO of TM1–TM7 is higher than in TMR, and it is lower in the case of the donor core. So, TM2 is the best because its donor moieties contribute the most (59.6%) to the HOMO. Similarly, TM7's acceptor fraction contributes 55.3% to the LUMO. Therefore, TM2 and TM7 stand out as the most notable molecules.

Table 4 Estimated results of λ_{max} , LHE, oscillator strength, excitation energy, and transition character of TMR together with TM1–TM7 in chlorobenzene along with exp. λ_{max}

Molecule	Calculated λ_{max} (nm)	Exp. λ_{max} (nm)	E_x (eV)	f_{os}	LHE	Assignment	Transition character (H \rightarrow L)
TMR	739	710	1.68	1.87	0.87	H > L	71%
TM1	820	—	1.51	1.98	0.90	H > L	71%
TM2	878	—	1.41	2.00	0.90	H > L	71%
TM3	861	—	1.44	2.02	0.90	H > L	71%
TM4	840	—	1.47	2.03	0.91	H > L	71%
TM5	835	—	1.48	2.18	0.93	H > L	71%
TM6	842	—	1.47	2.18	0.93	H > L	71%
TM7	837	—	1.48	2.01	0.90	H > L	71%



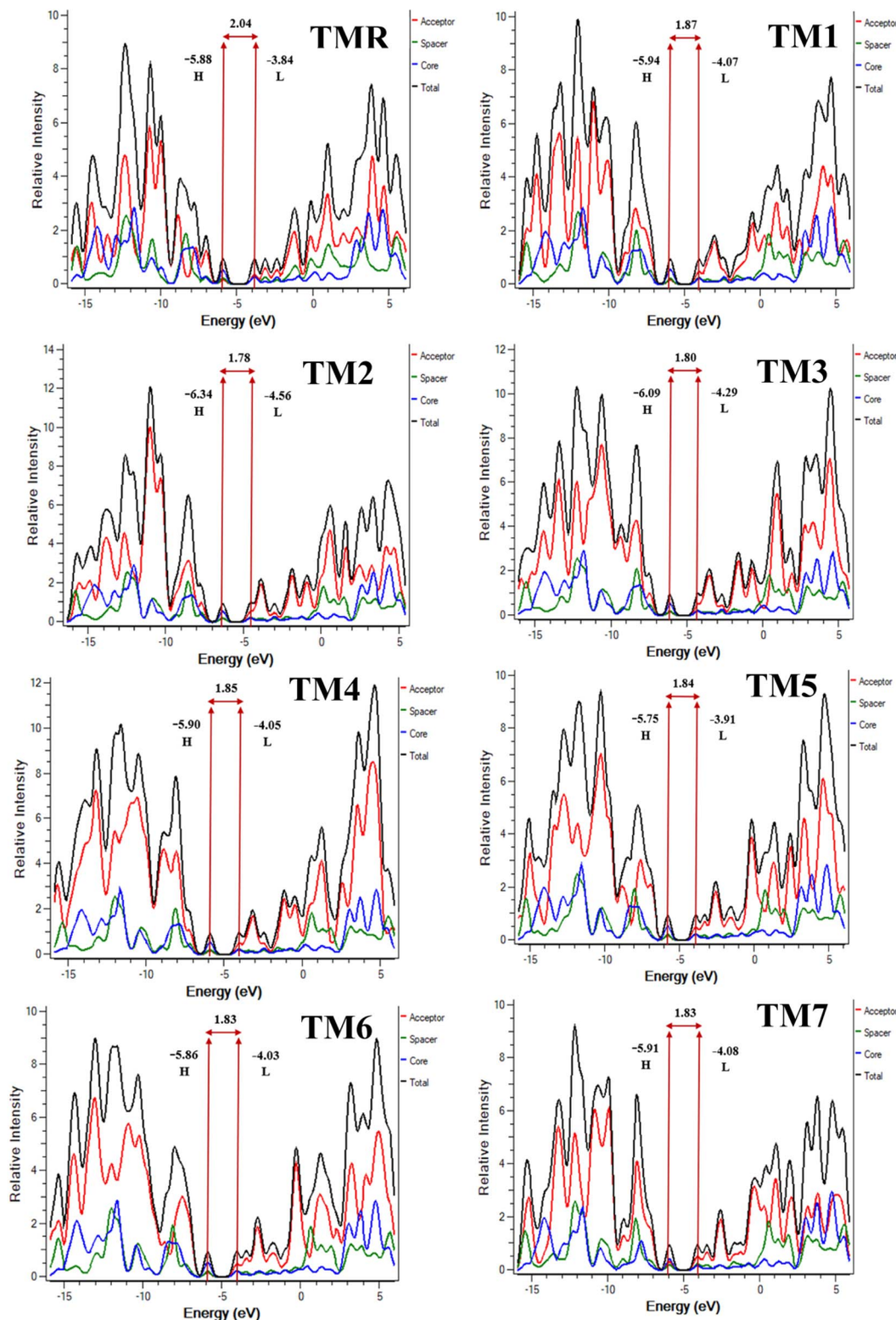


Fig. 6 DOS plots of TMR along with those of tailored compounds TM1–TM7.

3.7. Reorganization energy

Rearrangement energy analysis is a key factor in elucidating OSC performance. The RE of a molecule is the energy necessary to rearrange its constituent atoms and bonds followed by the

transport of electrons and holes inside the molecule. Competent materials for OSCs are driven by the appropriate RE, which corresponds to the charge content which has to be transferred easily from donating to accepting fragments and is concerned



Table 6 Estimated values for percentage contributions of acceptor, donor core as well as spacer of TMR and TM1–TM7 in FMO formation

Molecule	FMO	Acceptor (%)	Spacer (%)	Core (%)
TMR	HOMO	19.0	24.1	56.9
	LUMO	38.2	31.1	30.7
TM1	HOMO	18.1	22.5	59.4
	LUMO	49.0	26.1	24.9
TM2	HOMO	19.2	21.3	59.6
	LUMO	53.3	24.5	22.2
TM3	HOMO	19.0	21.7	59.3
	LUMO	52.5	24.7	22.9
TM4	HOMO	18.6	22.3	59.2
	LUMO	50.5	25.4	24.1
TM5	HOMO	19.3	22.7	58.0
	LUMO	50.8	25.0	24.1
TM6	HOMO	19.4	22.5	58.2
	LUMO	51.5	24.8	23.7
TM7	HOMO	33.8	22.2	44.0
	LUMO	55.3	25.4	19.3

with the hole–electron mobility.⁶¹ Thus this energy density (RE) is inversely linked to charge mobility. The more efficiently the charge transfers, the lower will be the RE value.⁶² The shape of cations and anions is just one of several important parameters that have an impact on RE. Table 7 displays the computed electron and hole RE values for all of the investigated compounds using eqn (1) and (2). RE of all of the molecules (TM1–TM7) including the reference (TMR) for positive (hole) and negative (electron) charges was examined through MPW1PW91/6-31G(d,p). Almost all of the seven altered molecules (TM1–TM7) had lower REs when compared with TMR proving that they are efficient electron transporters.

The RE (λ_h) values for holes of TMR and all modified compounds TM1–TM7 were 0.00967 eV, 0.00958 eV, 0.00917 eV, 0.00946 eV, 0.00981 eV, 0.00914 eV, 0.00930 eV, and 0.00921 eV, correspondingly. Similarly, the RE values for electrons of all the studied molecules were 0.01144 eV, 0.01060 eV, 0.00912 eV, 0.00977 eV, 0.01060 eV, 0.00977 eV, 0.00985 eV, and 0.01001 eV, correspondingly. These results for RE showed that all the designed molecules (with the exception of TM4) have lower hole RE compared to TMR, while the electronic RE (λ_e) for all the tailored structures TM1–TM7 are less than that of the reference TMR. Hole RE (λ_h) values determined *via* calculation are

Table 7 Estimated values for hole RE λ_h along with electron RE λ_e of TMR and TM1–TM7

Molecule	λ_e (electron)	λ_h (hole)
TMR	0.01144	0.00967
TM1	0.01060	0.00958
TM2	0.00912	0.00917
TM3	0.00977	0.00946
TM4	0.01060	0.00981
TM5	0.00977	0.00914
TM6	0.00985	0.00930
TM7	0.01001	0.00921

according to the order: **TM4** > **TMR** > **TM1** > **TM3** > **TM6** > **TM7** > **TM2** > **TM5**, while λ_e are in the order: **TMR** > **TM1** = **TM4** > **TM7** > **TM6** > **TM3** = **TM5** > **TM2**. **TM2** has the lowest electron RE and **TM5** has lowest hole RE of all the designed molecules. Thus **TM2** is a good electron transporter, while **TM5** is a good transporter of holes as they both provide the lowest RE values for both types of charges, respectively. The results show that the introduction of additional acceptor groups to the **TM2** and **TM5** molecules has improved the molecular charge transfer properties.

3.8. Molecular electrostatic potential

Another parameter which has proved effective to evaluate the degree of charge transfer between donor and acceptor sites is molecular electrostatic potential (MEP).⁶³ MEP is the 3-D visualization of charge density at different positions of a molecule of interest. The MEP correlates with a molecule's reactive potential by highlighting its electrophilic and nucleophilic centers. Herein we make use of green, blue, and red shades to represent the location of different charges (positive, negative or neutral) over the molecule. Here the red color represents negative charge density which is certainly over the acceptor region where there may be the electronegative atoms present. Alternatively the blue color represents positive charge which is mainly over the donor portion (central core) and the neutral area with zero charge is revealed by a green shade.

Fig. 7 illustrates that the thiophene spacers are neutral parts of the molecules revealed by green color and the terminal acceptor groups are red indicating that they have a higher negative charge due to the presence of unsaturated electronegative atoms: oxygen, nitrogen, and sulphur. The tailored **TM2** compound expresses more of a blue color over the acceptors and methyl groups of the donor part which reveals that these parts have more positive IP. Because electronegative atoms lie at a peripheral site the proposed compound **TM3** showed more of a red color (negative charge) than any of the others, which shows that this molecule has greater negative IP. These results show that all the designed compounds promise an effective PCE of OSC systems compared with the reference **TMR**.

3.9. Exciton binding energy (E_b) TDM

The exciton binding energy (E_b) is a key parameter in the study and evaluation of OSCs.⁶⁴ It represents the power required to bring the charge carriers apart once they have been created as a linked couple (exciton) This E_b is proportional to the coulombic interaction strength between the hole and the electron. Therefore, charge carriers with reduced E_b can travel conveniently towards their respective electrodes. Using eqn (7), we can hypothetically determine the binding energy of all the compounds that were studied:⁶⁵

$$E_b = E_g - E_x \quad (7)$$

Here, the excitation energy of the changed molecules is designated as E_x and the band gap as E_g , which is derived from eqn (3). The determined values for E_b of **TMR** and all other



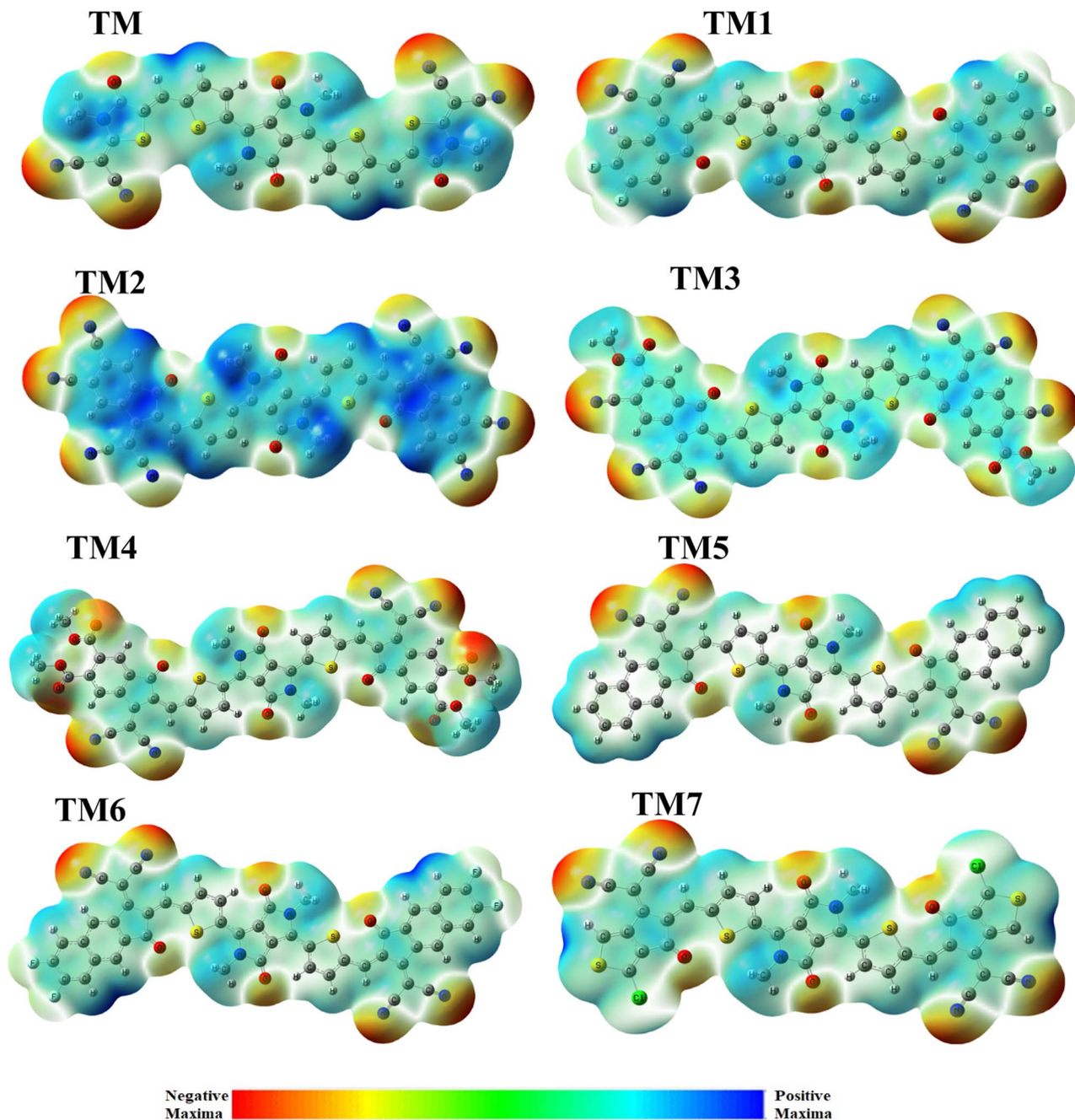


Fig. 7 Maps of MEPs of TMR along with those of tailored compounds TM1–TM7.

formulated structures **TM1–TM7** are 0.24 eV, 0.23 eV, 0.23 eV, 0.23 eV, 0.24 eV, 0.23 eV, 0.24 eV, and 0.22 eV in the gas phase, while 0.36 eV, 0.36 eV, 0.37 eV, 0.36 eV, 0.38 eV, 0.36 eV, 0.36 eV, and 0.35 eV in solution phase (chlorobenzene), correspondingly. Based on the data shown in Table 8, the binding energies of all the compounds follow the sequence: **TMR = TM4 = TM6** > **TM1 = TM2 = TM3 = TM5** > **TM7** in gas phase and **TM4** > **TM2** > **TMR = TM1 = TM3 = TM5 = TM6** > **TM7** in chlorobenzene. It is clear from this sequence that both **TM4** and **TM6** have the same binding energy as **TMR** in gaseous state, while **TM1**, **TM3**, **TM5**,

Table 8 Computed values for gas-phase as well as solvent (chlorobenzene)-phase exciton binding energy E_b of TMR and TM1–TM7

Molecule	E_b (eV) (gas phase)	E_b (eV) (solvent phase)
TMR	0.24	0.36
TM1	0.23	0.36
TM2	0.23	0.37
TM3	0.23	0.36
TM4	0.24	0.38
TM5	0.23	0.36
TM6	0.24	0.36
TM7	0.22	0.35



and **TM6** show this similar behavior in solution phase (chlorobenzene).

TM7, the molecule with the lowest E_b values, has the greatest potential to enhance the working of OSCs in terms of charge transfer and PCE.

TDM has been shown to be the most convincing tool among all others when a variety of processes, including electronic excitation, charge production, separation, recombination, and diffusion, are analyzed sequentially.⁶⁶ Through TDM, we can see how the quantum geometry of excited molecules works in practice. TDM determines the connection between excited

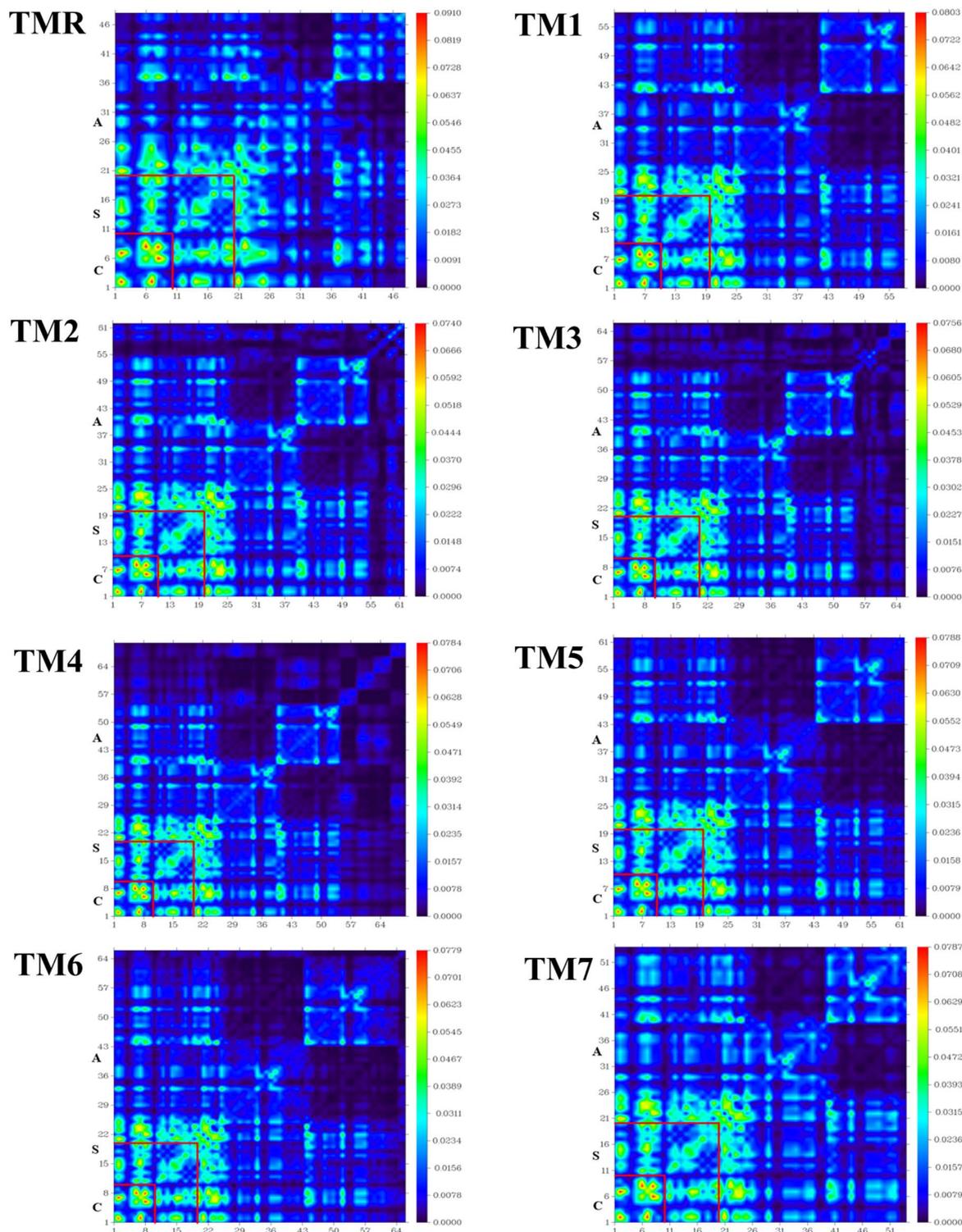


Fig. 8 TDM plots of TMR along with those of tailored compounds TM1–TM7.



donors and acceptors of molecules.^{67,68} TDM study of **TMR** and the designed molecules (**TM1–TM7**) was carried out using the desired hybrid DFT functional. Fig. 8 shows the TDM plots of **TMR** and the targeted molecules (**TM1–TM7**). Donor core (C), spacer (S) and acceptor (A) were the three divided slots for both the model molecule (**TMR**) as well as manufactured molecules (**TM1–TM7**). TDM plots depict atoms along the left and baseline axes, and electron density along the right axis. Hydrogen atoms have negligible impact, so they have been ignored.

A more stable and localized charge is found in the donor region (core) of the model (**TMR**) along with all the constructed **TM1–TM7** compounds. Advantageous localized electron density (diagonal) and charge transfer (off-diagonal) properties are present in the newly formulated molecules (**TM1–TM7**), which also have a more uniform charge on the acceptor section. In the case of molecules **TM3** and **TM4**, the donor core and spacer mainly reveal diagonal as well as off-diagonal behavior, while the acceptor fragment mainly demonstrated off-diagonal behavior. Both the localized electron density and charge transfer were shown by all the segments (acceptor, spacer, and core) of the **TM5**, **TM6**, and **TM7** chromophore. As faster exciton dissociation is necessary for OSCs to be built, the TDM plots demonstrated significant diagonal as well as off-diagonal charge transfer from electron-rich center (donor) to electron-poor center (acceptor). This indicates that coupling of electrons and holes is weaker in the modified molecules. As shown in Fig. 8, we see a visual depiction of the interaction coefficient of the studied molecules in chlorobenzene. TDM analysis and a declining trend in the interaction coefficient show that the tailored compounds are superior at transferring charge from the donor moiety to the acceptor fragment through thiophene spacers. As a result, compared to **TMR** the formulated structures **TM1–TM7** show remarkable charge dissociation capacity.

3.10. Open circuit voltage (V_{OC})

Open circuit voltage (V_{OC}) provides a significant contribution in determination of the photovoltaic features of NFA-based organic photovoltaic devices. It is the greatest voltage that can be drawn from a photovoltaic device at zero current level.^{69–71} The open circuit voltage is influenced by several factors such as device temperature, molecule energy levels, light intensity, and charge-carrier recombination.⁷² The maximum value for V_{OC} is

calculated by comparing the HOMO value with LUMO value. To attain a value of V_{OC} , the value of energy of HOMO of the donor must be smaller and the LUMO of the acceptor should be higher, which results in reduced band gap which is the desired condition for solar cells from any aspect. Here smaller value of HOMO corresponds to negative values which when becoming smaller, the actual HOMO value becomes higher and *vice versa*. Similar is the case with LUMO. A smaller band gap maximizes the absorption, which in return enhances the V_{OC} and light harvesting efficiency. The maximum value for V_{OC} is associated with success of NFA-based OSCs in terms of improved PCE. The V_{OC} has direct relation with the effectiveness of OSCs: the higher the V_{OC} the greater will be the efficiency.⁷³ Visual representation of a comparison between the LUMO of the acceptor molecule with the HOMO of the donor polymer PTB7-Th to ascertain V_{OC} is shown in Fig. 9. The V_{OC} is determined mathematically through the use of the following eqn (8):⁷⁴

$$V_{OC} = \frac{E_{LUMO \text{ of acceptor}} - E_{HOMO \text{ of donor}}}{e} - 0.3 \quad (8)$$

Here, e represents molecular charge having constant value of 1, 0.3 corresponds to empirical factors, which is also constant for every molecule, and E denotes energy of the utilized FMOs. PTB7-Th has values of -5.20 eV and -2.60 eV for HOMO and LUMO, respectively.^{44,75} The calculated values of open circuit voltage for reference molecule **TMR** and constructed compounds (**TM1–TM7**) are 1.06 V, 0.83 V, 0.34 V, 0.61 V, 0.85 V, 0.99 V, 0.87 V, and 0.82 V, respectively, as presented in Table 9. Amongst all the newly developed structures, **TM5** (0.99 V) and

Table 9 Computed values for V_{OC} , normalized V_{OC} along with the fill factor of **TMR** and **TM1–TM7**

Molecule	V_{OC} (V)	Normalized V_{OC}	Fill factor
TMR	1.06	41.01	0.89
TM1	0.83	32.11	0.86
TM2	0.34	13.15	0.74
TM3	0.61	23.60	0.83
TM4	0.85	32.88	0.87
TM5	0.99	38.30	0.88
TM6	0.87	33.66	0.87
TM7	0.82	31.72	0.86

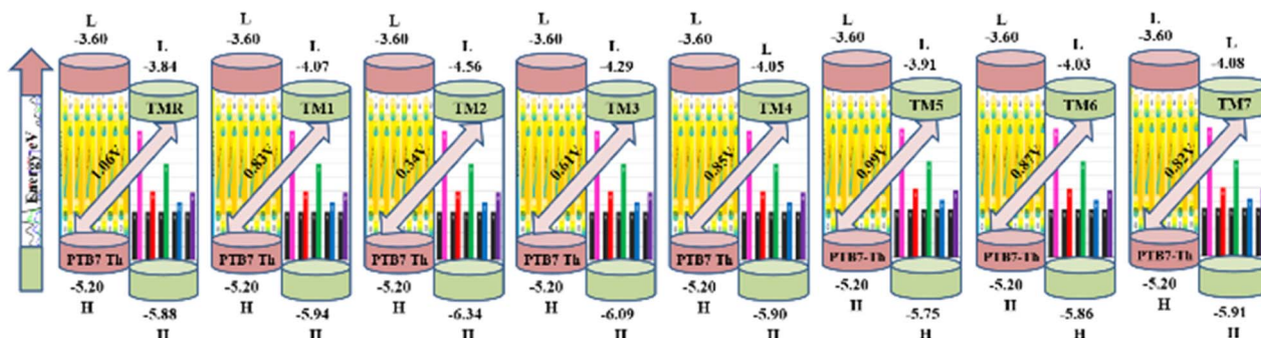


Fig. 9 Open circuit voltage of **TMR** and **TM1–TM7** molecules with PTB7-Th.



TM6 (0.87 V) have been proved as molecules with highest V_{OC} due to the involvement of the end-capped electron-withdrawing acceptor entities, from which it is concluded that **TM5** and **TM6** exhibit more charge delocalization, conjugation, and higher PCE than **TM1**, **TM2**, **TM3**, **TM4**, and **TM7** molecules.

3.11. Fill factor (FF)

The PCE and other photovoltaic properties of NFA-based OSCs can also be estimated through the utilization of a convincing tool, *i.e.* fill factor. The fill factor of reference **TMR** and other

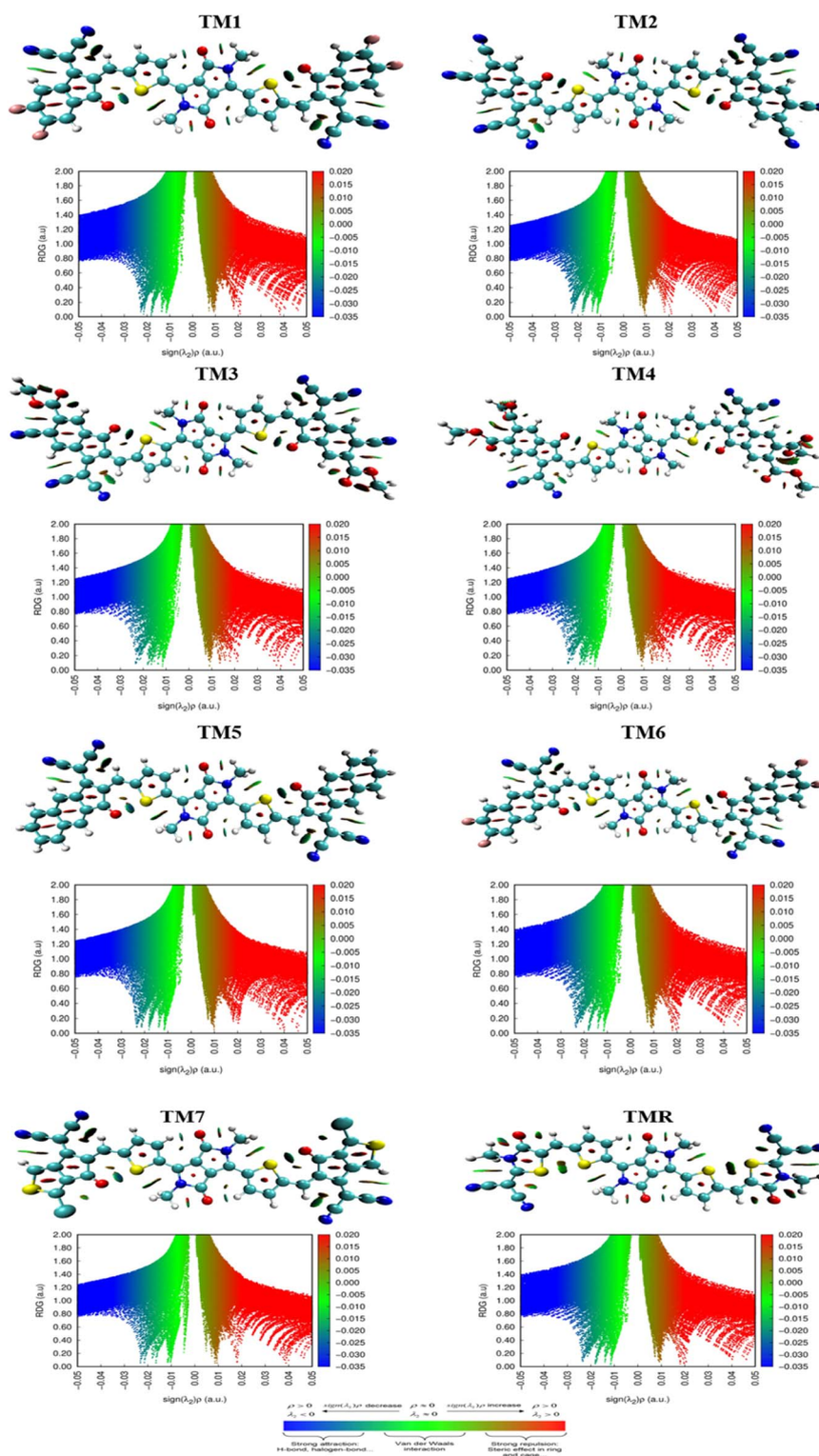


Fig. 10 NCIs and iso-surfaces of TMR and TM1–TM7 molecules.



modified compounds **TM1–TM7** can be measured with the application of eqn (9):⁷⁶

$$FF = \frac{\frac{eV_{OC}}{K_B T} - \ln\left(\frac{eV_{OC}}{K_B T} + 0.72\right)}{\frac{eV_{OC}}{K_B T} + 1} \quad (9)$$

In this equation, open circuit voltage is represented by V_{OC} , e denotes the charge of each compound, T characterizes the temperature of the system that was 300 K, Boltzmann constant is symbolized by K_B and the normalized V_{OC} is represented by $\frac{eV_{OC}}{K_B T}$. The calculated fill factor values for the reference molecule and the designed chromophores (**TM1–TM7**) are summarized in Table 9, these values being 0.89, 0.86, 0.74, 0.83, 0.87, 0.88, 0.87 and 0.86, respectively. The measured results displayed that the molecules **TMR**, **TM4**, **TM5** and **TM6** have almost the similar value of fill factor, which is why they exhibit admirable photovoltaic properties compared with **TM1**, **TM2**, **TM3**, and **TM7** having lower FF value. The descending pattern of FF of all compounds follows the sequence: **TMR** > **TM5** > **TM4** = **TM6** > **TM1** = **TM7** > **TM3** > **TM2**.

J_{sc} (short circuit current), V_{OC} , power of incident light and FF contribute collectively to impart a positive effect on the PCE of NFA-based OSCs as represented by eqn (10):^{77,78}

$$PCE = \frac{J_{sc} V_{OC} FF}{P_{in}} \quad (10)$$

From the mentioned equation, it is clear that V_{OC} , J_{sc} , and FF directly affect the PCE of OSC materials: by increasing the values of these parameters PCE will also be increased.⁷⁹ On the other hand, power of incident light is inversely related to PCE. By knowing the numerical values of these parameters, we can judge effectively the value for PCE.

3.12. Iso-surface and reduced density gradient (RDG) analysis

With an emphasis on non-covalent interactions between molecules, the non-covalent analysis is a 2D RDG graph that can discriminate among weak van der Waals forces, hydrogen bonds and repulsive steric forces. Molecular scatter plots are developed by the Multiwfn software tool. λ_2 values are taken into account while assessing the type of NCIs. Graphical illustration of iso-surfaces and their respective NCIs for all derived molecules (**TM1–TM7**) along with **TMR** are shown in Fig. 10. Blue, green and red colours in graphs indicate strong interactions, van der Waal forces and weak interactions respectively as indicted in the coloured bar in Fig. 10.

4. Conclusion

The effects of end-capped acceptors on the charge transfer ability as well as on structural and photochemical activities of **TMR** have been investigated using state-of-the-art quantum chemical and quantum mechanical methods. Seven different

acceptor molecules (**TM1–TM7**) were formulated *via* the end-cap engineering of the reference **TMR**. These compounds were designed with the purpose of investigating their opto-electronic characteristics for use in OSCs with enhanced efficiency. Light absorption, electron–hole RE, nucleophilic as well as electrophilic sections inside the molecule (MEP), band gap between FMOs, involvement of acceptor and donor in FMOs (DOS), and TDM were all investigated. All seven developed structures (**TM1–TM7**) when compared to the published reference **TMR** molecule exhibited notable improvement regarding all the above mentioned parameters, *i.e.* a reduced band gap in the range of 1.87–1.78 eV, a dipole moment that is larger than that of **TMR** in both the examined phases (gas and solvent), lower hole as well as electron REs, and maximum photon absorption (799 nm in gas phase and 878 nm in chlorobenzene solvent), which were very impressive, thanks to modifications by end-capped acceptor moieties. Therefore, it is suggested that all formulated candidates are promising for use in OSCs in order to obtain high PCE in the near future, since almost all of the unique acceptor molecules **TM1–TM7** have better optical and electrical characteristics compared to the reference **TMR**.

Data availability

Data will be made available on request.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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