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Effect of tailoring π -linkers with extended conjugation on the SJ-IC molecule for achieving high $V_{\rm OC}$ and improved charge mobility towards enhanced photovoltaic applications†

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The problem of low efficiency of organic solar cells can be solved by improving the charge mobility and open circuit voltage of these cells. The current research aims to present the role of π -linkers, having extended conjugation, between the donor and acceptor moieties of indacenodithiophene core-based A- π -D- π -A type SJ-IC molecule to improve the photovoltaic performance of pre-existing SJ-IC. Several crucial photovoltaic parameters of SJ-IC and seven newly proposed molecules were studied using density functional theory. Surprisingly, this theoretical framework manifested that the tailoring of SJ-IC by replacing its π -linker with linkers having extended π -conjugation gives a redshift in maximum absorption coefficient in the range of 731.69-1112.86 nm in a solvent. In addition, newly designed molecules exhibited significantly narrower bandgaps (ranging from 1.33 eV to 1.93 eV) than SJ-IC having a bandgap of 2.01 eV. Similarly, newly designed molecules show significantly less excitation energy in gaseous and solvent phases than SJ-IC. Furthermore, the reorganization energies of DL1-DL7 are much lower than that of SJ-IC, indicating high charge mobility in these molecules. DL6 and DL7 have shown considerably improved open circuit voltage ($V_{\rm OC}$), reaching 1.49 eV and 1.48 eV, respectively. Thus, the modification strategy employed herein has been fruitful with productive effects, including better tuning of the energy levels, lower bandgaps, broader absorption, improved charge mobility, and increased $V_{\rm OC}$ Based on these results, it can be suggested that these newly presented molecules can be considered for practical applications in the future.

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1. Introduction

The impressive photoactivity of bulk heterojunction (BHJ) organic solar cells (OSCs) is linked to non-fullerene acceptors (NFAs) that possess easily tunable frontier energy levels. This characteristic is crucial in reducing the band gap and improving the absorption properties of photoactive materials.¹⁻³ Fused-

ring NFAs have upgraded the photovoltaic activity of OSCs. The photovoltaic performance of OSCs was limited due to fixed frontier energy levels and poor absorption of fullerene acceptors in the visible and near-infrared spectral ranges. Significant improvements in the performance of OSCs have been attained because of continuous efforts from both computational and synthetic approaches. Although noticeable advancements have improved fullerene-free OSCs, their efficiency is still insufficient for commercial applications.

In order to bring about an improvement in the overall efficiency of fullerene-free OSCs, it is essential to focus on important factors such as narrowing the band gap, increasing the range of light absorption of chromophores, reducing excitation energy and reorganization energy, and increasing the open circuit voltage ($V_{\rm OC}$). Narrowing the band gap is vital as it increases the number of photons that can produce photovoltaic current. By expanding the light absorption range of chromophores, OSCs can capture a broader spectrum of sunlight, resulting in higher efficiency. Moreover, reducing excitation and reorganization energy can minimize energy loss during the

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charge transfer process and increase charge mobility. Increasing $V_{\rm OC}$ is also essential as it is directly related to the maximum power output of a solar cell. By improving any of these crucial factors, researchers and scientists can work towards achieving greater efficiency in fullerene-free OSCs. 11-14

Different computational and synthetic techniques have been used to improve the efficiency of fullerene-free OSCs. Among these techniques, increasing the conjugation length in molecules is an effective strategy that can be achieved by inserting π bridges for further enhancement of the ability of the molecules to harvest light and, in turn, boost V_{OC} of OSCs. Such a method also forms resultant channels for the transport of electrons between the donor core and acceptor units.15 Electrons are delocalized over the entire molecule surface in conjugated systems with extended π -electron systems. This results in many closely spaced energy levels, which can lead to light absorption at longer wavelengths, i.e., in the near-infrared region.16 Furthermore, π -bridges function as "conformational locks"; thus, they can improve planarity. 17 Recently, Hai and coworkers investigated the effect of incorporating vinylene π -linkers in a central ladder-type fused core and developed two new molecules named BTP-2V-2F and BTP-1V-2F. BTP-2V-2F showed a power conversion efficiency (PCE) of 11.22% and a high shortcircuit current of 26.50 mA cm⁻², while the latter exhibited a remarkable PCE of 14.24%.18

In 2017, Li *et al.* used the strategy of π -linker insertion. They developed a novel compound named SJ-IC, having a ladder-type core of pentacyclic arene 4,9-dihydro-*s*-indaceno[1,2-*b*:5,6-*b*'] dithiophene (IDT) with four phenyl units as side chains to enhance its electron-donating character. In SJ-IC, the IDT donor part is separated from the IC acceptor *via* a vinyl group as a π -linker. Owing to the proficient structural fragments, SJ-IC exhibits strong absorption in chloroform solvent in the 560–800 nm wavelength range with a peak prominent at 720 nm. SJ-IC acting as acceptor along with J61 donor in a solar cell shows a high PCE of about 9.27%, notable $J_{\rm SC}$ of approximately 16.99 mA cm⁻², $V_{\rm OC}$ of about 0.83 V, film factor with a value of 0.6595 and having an electron mobility of 5.73 \times 10⁻⁴ cm² V⁻¹ s⁻¹.19

The current research further explores the effect of π -bridging in SJ-IC to enhance its already remarkable optoelectronic properties to design novel molecules for efficient OSCs. Until now, 19% PCE has been achieved in non-fullerene OSCs. But SJ-IC has significantly lagged in showing promising efficiency. SJ-IC (R) is structurally tailored to improve IDT core-based molecules by replacing vinyl spacers with other π -linkers with enhanced conjugation. Thus, seven new molecules have been designed. Herein, we derived all the investigated molecules from SJ-IC by replacing the simple π -bridge with methoxythiophene in DL1, 2-thia-4,6,9-triazacyclopenta [b]naphthalene in DL2, 2-oxa-4,6,9-triazacyclopenta [b]naphthalene in DL3, [3,4-b]quinoxaline in DL4, methylthieno[3,4-b]thiophene-2-carboxylate in DL5, 3,4-dihydro-2*H*-thieno[3,4-*b*] [1,4]dioxepine in DL6 and 2,3-dihydrothieno[3,4-b][1,4]dioxine in DL7 as shown in Fig. 1. These π -bridges have the potential to enhance crucial photovoltaic parameters such as increasing the light absorption ability, narrowing the bandgap, reducing reorganization and excitation energy, and enhancing $V_{\rm OC}$ of these molecules.

2. Computational methodology

The 3D molecular frameworks of all the molecules involved in the current research work were designed with the help of GaussView 6.0 to study the various optical and electronic parameters. For simplicity, long alkyl substituents attached to the phenyls on the core of the reference (SJ-IC) molecule were replaced by methyl to save computational time and cost. The quantum simulations were progressed by Gaussian 09 software20 using DFT (density functional theory)21 and TD-DFT (time-dependent DFT)22 approaches, which are considered reliable for calculating ground state and excited state properties of OSC molecules. The reference (SJ-IC) molecule was optimized in the ground state with a restricted electronic spin by using four different functionals of the DFT, i.e., B3LYP,23 CAM-B3LYP,²⁴ MPW1PW91,²⁵ and WB97XD²⁶ in conjunction with the 6-31G(d,p) basis set.27 After achieving geometry optimization with different functionals, the calculations in the solvent medium were performed by employing chloroform solvent via the IEFPCM²⁸ model of the TD-DFT approach. Chloroform was used as a solvent because it was used initially for SJ-IC in the literature¹⁹ to check out the effect of solvent on the light absorption capability of molecules in the excited state.

To determine the best molecular computational method, the obtained results with these different functionals were compared to the experimental results of the SJ-IC molecule. It was observed that experimental optoelectronic attributes of SJ-IC molecules, such as band gap and the maximum absorption (λ_{max}) , strongly agreed with the results obtained using the B3LYP/6-31G(d,p) level of theory. The λ_{max}^{cal} with B3LYP functional had a value of 731.69 nm, while the value of λ_{max}^{cal} was 546.11 nm with CAM-B3LYP functional, 691.11 nm with MPW1PW91, and 525.12 nm with WB97XD. This shows that use of B3LYP functional leads to more concordance with the experimental maximum spectral absorption (720 nm). For the convenience of comparison, Fig. S1 (ESI†) presents a graphical comparison of the results obtained using various functionals. Therefore, all the designed molecules were processed using the B3LYP functional of DFT to get all the results under similar conditions for parallel comparison.

Spectral illustrations for the absorption maxima (λ_{max}) in gaseous and solvent phases were realized by Origin 6.0 software. Multiwfn 3.8 software was used to reconstruct TDM (transition density matrix) data into actual graphs to inspect exciton dynamics and their interactions. The same software was used for developing non-covalent interaction (NCI) graphs and isosurfaces and the VMD program was used to visualize isosurfaces. DOS (density of states) graphs were generated using the PyMOlyze1.1 (ref. 22) program to reveal the extent of each molecule's involvement of acceptors, linkers, and donor parts.

It is important to note that the reorganization energy influences the efficiency of a solar cell. It gives information about the charge transfer capability of OSC molecules, specifically the mobility of holes and electrons during excitation. This essential

ig. 1 Pictorial representation of the molecular structures for the reference and the newly designed molecules with inserted linkers.

parameter can be calculated using the Marcus equation. Using the pre-selected level of theory, *i.e.*, B3LYP/6-31G(d,p), all the studied molecules were optimized in cationic and anionic states, followed by simulations for energy calculations in cationic and anionic states. Eqn (1) and (2) (ref. 31) were applied to calculate the reorganization energy of electrons (λ_e) and holes λ_h of each molecule:

$$\lambda_{\rm h} = [E_0^+ - E_+] + [E_+^0 - E_0] \tag{1}$$

$$\lambda_{e} = [E_{0}^{-} - E_{-}] + [E_{-}^{0} - E_{0}] \tag{2}$$

in the above expressions, E represents the heat of formation of molecules (eV). The negative sign (-) depicts the optimized anionic state, the positive sign (+) depicts the optimized

cationic state, and the optimized neutral state is depicted by the zero (0) symbol in subscripts. In contrast, the superscripts represent the charge state in which the energy of optimized cation, anion, and neutral molecules was calculated.³²

Results and discussion

3.1 Geometry analysis

The SJ-IC molecule has three parts: a ladder-shaped donor or "push" part (IDT), C–C double bonds as π -bridges, and an electron-deficient acceptor, IC, "pull" fragment. As portrayed in Fig. 2, it can be observed that the SJ-IC molecule is optimized in one plane. The four phenyl moieties are arranged in another plane in order to minimize steric hindrance at low potential energy with 113° torsion angles. These phenyls lessen the

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Front view Side view R DL1 DL2 DL3 DL5 DL6 DL7

Fig. 2 Optimized structures of R and designed molecules (DL1–DL7) with side views at right.

intermolecular interactions due to steric hindrance and thus prevent self-aggregation.³³ Lengths of the bonds connecting donor-linker and linker-acceptor parts of the SJ-IC molecule

have values ranging from 1.42680 to 1.42034 Å. The dihedral angle between the donor and spacer is represented by θ_1 while θ_2 represents the dihedral angle between the spacer and acceptor. The dihedral angles between these connecting fragments have values of 0.00129° and 0.00476°, respectively. These values are very close to zero. The acquired outcomes indicate a higher degree of planarity and conjugation, both essential for charge transfer.

Planarity is one of the vital factors responsible for charge transfer. Planarity is highly affected by the dihedral angle between the molecule's constituents. Planarity will be more prominent in molecules with smaller dihedral angles, increasing the possibility of charge transfer. The dihedral angle values for all our proposed molecules range from 0.00603° to 5.99593°, as summarized in Table 1. It is predicted that all the torsion angle values are considerably smaller for two reasons. One reason is the rotation restriction over single bonds due to the insertion of π -linkers with extended conjugation. The second reason for the smaller torsion angle can be the presence of conformation locks such as noncovalent interactions (S···O, S···H, O···H, and S···N) among subunits of each molecule which hold aromatic rings in the same plane.34 DL5 and DL6 have shifted slightly from the central plane due to lone pairs on the oxygen atoms of linkers.

The bond lengths between atoms that connect donor-spacer (l_1) and spacer-acceptor (l_2) were also calculated. Their values were observed to be in the 1.39471–1.43451 Å range. These data indicate that the bond lengths between molecule fragments are somewhere between those for C–C single (1.54 Å) and C=C double (1.34 Å) bonds. This shows the presence of conjugation at these points. As these bonds connect the molecules' fragments, it can be inferred that charges can transfer from the donor to the acceptor portion in these molecules due to conjugation.

3.1.1 Role of non-covalent interactions and isosurfaces. To understand the strength and nature of NCI present in SJ-IC and the designed structures, NCI graphs based on a reduced density gradient (RDG) were plotted using Multiwfn 3.8 software, as shown in Fig. 3. NCI graphs visualize repulsive (steric hindrance) and attractive interactions (like H-bonding, S···O, S···N, and van der Waals forces) among various parts of the molecule based on electron density (ρ) and RDG. NCI plots show

Table 1 Computed values of bond angles and bond lengths for SJ-IC and DL1–DL7

	Dihedral ar	ngle (θ°)	Bond length (L_{C-C})	
Molecule	$\theta_{1}^{^{\circ}}$	$\theta_2^{^\circ}$	l_1	l_2
R	0.00129	0.00476	1.42680	1.42034
DL1	0.37208	0.00603	1.43025	1.41556
DL2	0.10320	0.02090	1.42059	1.40571
DL3	0.36913	0.43839	1.40986	1.39471
DL4	0.05060	0.00880	1.42106	1.40350
DL5	5.91159	0.12303	1.43216	1.41254
DL6	5.99593	0.65967	1.43451	1.40838
DL7	2.32446	0.19666	1.43203	1.40981

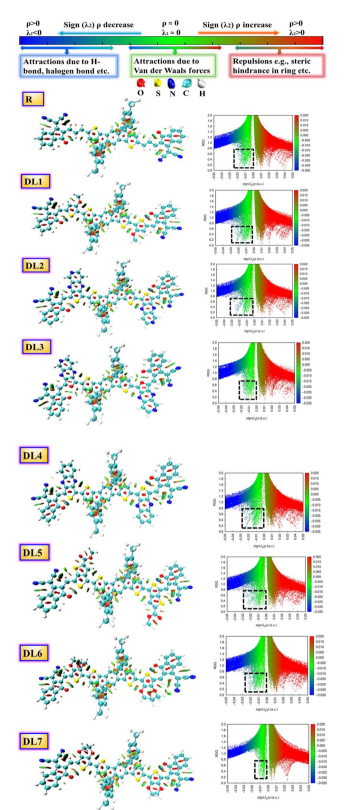


Fig. 3 NCI graphs and isosurfaces for the researched molecules. Black dotted squares highlight the presence of non-covalent interactions for R and DI 1–DI 7

RDG on the *y*-axis and the product of energy and the second eigenvalue $(\text{sign}(\lambda_2)\rho)$ on the *x*-axis. The regions where the electron density is non-interacting have high RDG but low ρ ,

whereas high ρ values and RDG indicate regions of covalent bonds. However, if both RDG and ρ are low, this indicates noncovalent interactions.

In this research, we focused on the appearance of peaks on the left y-axis of the graphs. These peaks have a negative $\operatorname{sign}(\lambda_2)$ and correspond to various non-covalent interactions. When the electronegativity of atoms within a molecule increases, the peak spread of $\operatorname{sign}(\lambda_2)\rho$ is shifted further towards negative values. This indicates that stronger hydrogen bonding is present in the molecule.

The presence of attractive non-covalent interactions in the newly designed molecules is evident from NCI graphs in Fig. 3. It can be seen that the green-colored peaks are spread toward the left. At the same time, the negative values of the effective density " $(\text{sign}(\lambda_2)\rho)$ " in all of our conceptualized molecules reveal the regions of electron depletion. Vastly reduced density gradients indicate strong interactions, whereas less reduced density gradients show weaker interactions.

It can be seen from the graphs that van der Waals forces of attraction are highly dominant in DL2 and DL6 molecules as compared to the other molecules. These weak forces are very much responsible for enhancing the stability of molecules. Furthermore, the blue line (on the left side) is above the red band (on the right side) for all the molecules, which signifies that attractive forces are more dominant than repulsive ones. The domination of attractive forces makes these molecules more stable.

Different colors of isosurfaces symbolize different interactions according to electron density. The labeled scale in Fig. 3 has the blue color showing strong, attractive interactions like strong H-bonding. The green color corresponds to mediumweak interactions, whereas red shows repulsive interactions. In DL1–DL7, disc-like blue isosurfaces depict the presence of various interactions, including S···O, S···H, S···N, O···H, *etc.* While remaining irregular patches symbolize some interactions, such as London dispersion and van der Waals forces. The small red patches inside the five-membered rings of the central donor core and linkers show repulsive interactions.

Similarly, among the phenyl substituents, large isosurface patches of red and green colors show the presence of repulsive and attractive forces. Repulsions push them out of the plane while the attractive interactions try to keep them in the plane with the rest of the molecule. It can be observed that green patches are present across each molecule in higher quantity, which confirms the dominance of attractive interactions in all these molecules leading towards enhanced stability.

3.2 Frontier molecular orbitals and absorption properties

Frontier molecular orbital (FMO) analysis is mandatory to study the excited state features of the computationally designed molecules. The efficiency of a solar cell is related directly to charge transfer.³⁷ So, to gain insight into the electronic, opto-electrical, and conductive properties of the designed structures, we performed an FMO analysis.³⁸ Band theory explains the push–pull mechanism of electrons by taking HOMO as an electron donor (valence band) and LUMO as an electron

acceptor (conduction band).39,40 The difference between energy levels of HOMO and LUMO is called a band gap which is calculated using eqn (3):

$$E_{\rm g} = E_{\rm LUMO} - E_{\rm HOMO} \tag{3}$$

The lesser the distance between HOMO and LUMO, the smaller will be the band gap and, hence, the better will be charge transfer. Improved charge transfer boosts a device's overall efficiency.41 For narrower band gaps, the HOMO level of a molecule should lie at a higher energy state. The LUMO should lie at a low energy state, i.e., close to the HOMO. Stable LUMOs can hold maximum electrons in the acceptor vicinity for extended periods, allowing more photon absorption and lifting device performance.37

To reveal the impact of π -spacers on designed molecules DL1-DL7, we studied HOMO and LUMO energy levels along with their respective band gaps at B3LYP/6-31G(d,p) level of DFT, and results are summarized in Table 2. For SJ-IC, the HOMO and LUMO levels are present at -5.43 eV and -3.42 eV, respectively, resulting in a band gap of 2.01 eV. The HOMO levels for DL1-DL7 showed values of -5.02, -5.29, -5.34, -5.14, -5.30, -5.04, and -5.07 eV. On the other hand, energies of their LUMO appear to be at energy onset of -3.26, -3.95, -3.99, -3.73, -3.45, -3.11, and -3.12 eV, respectively. Meanwhile, the HOMO and LUMO of SJ-IC are present at energy levels of -5.43 and -3.42 eV, respectively. These data show that the

LUMO of DL2, DL3, DL4, and DL5 reside at higher energy levels than the LUMO of the SI-IC molecule. This indicates that these molecules have a higher probability of charge transfer from HOMO to LUMO than the reference SJ-IC molecule.

The ideality of the designed molecules is further confirmed by examining each studied molecule's energy difference among FMOs. The newly developed molecules (DL1-DL7) showed bandgap values of 1.76, 1.33, 1.35, 1.40, 1.85, 1.93, and 1.95 eV, respectively. These results show that energy gaps for all investigated molecules are smaller than that for reference (SJ-IC), having a bandgap of 2.01 eV.

DL2 seemed to have the narrowest band gap among all these investigated structures. This confirms that the spacer subunit with conjugated rings containing electronegative nitrogen and sulfur atoms performs better than the other spacers. Furthermore, DL3 and DL4 also show remarkable narrowing of the bandgap due to the presence of the extended conjugated network in spacer subunits. However, all the newly developed molecules (DL1-DL7) exhibited smaller bandgaps than the reference (SI-IC). Hence, these molecules have marked charge transfer, confirming that inserting π -linkers with extended conjugation effectively modulates the molecules to boost their efficiency.

Further confirmation of the above observations was carried out by obtaining diagrams of FMOs from the relaxed geometry of each molecule. It visually represents intramolecular charge

Table 2 Computed values for energies of HOMO and LUMO of investigated structures alongside energy gaps between them and FWHM

Molecule	π -linker	$E_{\mathrm{HOMO}}\left(\mathrm{eV}\right)$	$E_{ m LUMO}$ (eV)	E_{g} (eV)	FWHM (nm)
R		-5.43	-3.42	2.01	306
DL1	S	-5.02	-3.26	1.76	393
DL2	N N N N N N N N N N N N N N N N N N N	-5.29	-3.95	1.33	840
DL3	N N O	-5.34	-3.99	1.35	768
DL4		-5.14	-3.73	1.40	675
DL5		-5.30	-3.45	1.85	395
DL6	Co	-5.04	-3.11	1.93	389
DL7	Co Co	-5.07	-3.12	1.95	383

transfer⁴¹ and the relation between structure and energy levels.³⁵ Dense colors have been used in FMO pictograms (Fig. 4) to express the flow pattern of the charges (electrons and holes)

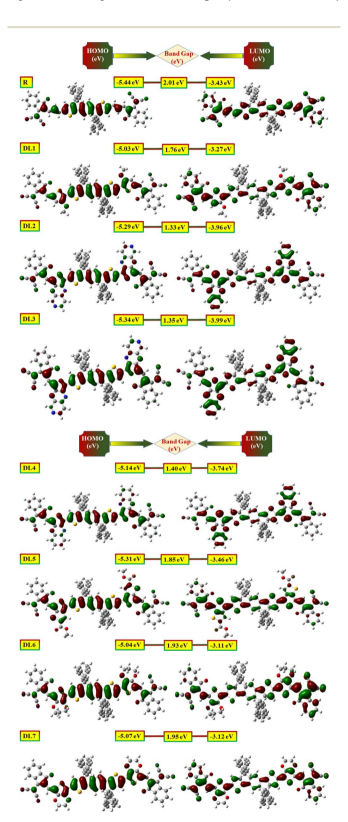


Fig. 4 Visual display of highest occupied molecular orbital (HOMO) (left) along with lowest unoccupied molecular orbital (LUMO) (right) of reference and DL1–DL7 molecules.

whose conduction provokes electric current. In R, HOMO is located on the donor part, representing that charge density is located on the donor in the ground state. In contrast, the LUMO part is mainly on the acceptor and π -bridges. It shows that the charges are attracted towards acceptor moieties of the molecule through π -bridges upon excitation. In all designed molecules, the pictures of their FMOs reveal that the donor units hold HOMO density more efficiently than the acceptors.

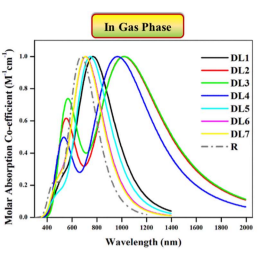
Likewise, LUMO density is mainly positioned on acceptor units (especially in DL1, DL6, and DL7). The charge density on π -bridges depicts their contribution to charge flow from HOMO to LUMO. These bridges are highly involved in pulling the electric charge density from the central donor core and passing these charges toward acceptor moieties. The above discussion clarifies the importance of our work, as electronic parameters show that these proposed molecules are appropriate candidates to be used for the construction of improved OSCs.

Optical properties play a pivotal role in estimating the overall performance of a device.⁴² Range of photon absorption, full width at half maximum (FWHM), excitation energy, oscillating strength, and involvement of orbitals contribute to the optical properties of the active layer of OSCs.⁴³ All the studied molecules (R and DL1–DL7) were investigated with the B3LYP/6-31G(d,p) functional of DFT approach in gas and solvent phase. This study was performed to record spectra for the absorption in the UV-visible region and then translated into graphs using the Origin 6.0 program. These graphs are illustrated in Fig. 5 to visualize these molecules' maximum absorption of photons.

Experimental absorption coefficient (λ_{max}^{exp}) for R is reported at 720 nm, which is found to be in good agreement with DFT-based calculated absorption maximum (λ_{max}^{cal}) located at 731.69 nm. The literature shows that a molecule's absorption spectrum strongly depends on the nature of its substituents.³⁵ As in all our planned molecules, building blocks differ regarding their π -linkers. The study of the optical properties helps to decide whether the insertion of investigated π -linkers is fruitful. In the gas phase, all the investigated molecules gave λ_{max}^{cal} in the range of 709–1023 nm. All the investigated molecules show redshifts ranging from 34 to 348 nm compared to R as evident from their λ_{max}^{cal} provided in Table 3.

We used chloroform for solvent-phase calculations to determine these molecules' working potential in a solvent medium. The parameters calculated for absorption in the solvent phase are illustrated in Table 3. It can be observed that $\lambda_{\rm max}^{\rm cal}$ of each molecule is higher in the solvent phase than in the gas phase. The highest λ_{max}^{cal} values are exhibited by DL2 and DL3, i.e., 1112 nm and 1094 nm, respectively. UV-visible regions are important spectral regions that are responsible for high light-to-current transformation. Interestingly, λ_{max}^{cal} for DL2, DL3, and DL4 extend to the NIR region while covering the UVvisible range. DL2 exhibited a redshift of 381 nm in λ_{max}^{cal} compared to R. This can be due to the mutual electron withdrawal effect of nitrogen atoms in bridges along with CN groups of the acceptor. Similarly, in all designed molecules, the sulfur, nitrogen, and oxygen in linkers having an enormous electronegative difference provide good electron delocalization

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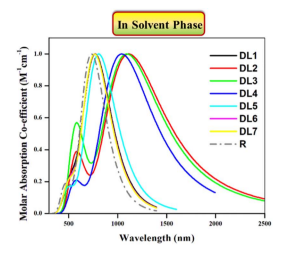


Fig. 5 Absorption profiles of investigated molecules in UV-visible region in the gas phase (left) and in chloroform solvent (right) obtained through Origin 6.0 software at B3LYP functional.

compared to a carbon atom in R, leading to higher absorption maxima. High values of $\lambda_{\rm max}^{\rm cal}$ in the solvent phase also reveal that all these tailored molecules can be effectively used for solution processing of OSCs. Moreover, as all these molecules show light absorption in a broad range, they can produce more photoelectric current by absorbing radiation in a wide range which is beneficial for improving efficiency.

If we examine the graphs for $\lambda_{\rm max}^{\rm cal}$ in the gas and solvent phases, it can be seen that two peaks characterize each molecule. One peak of smaller height located between 400 and 700 nm corresponds to π - π^* transitions present in the respective molecule. In contrast, the heightened peak depicts absorption maxima for each molecule. The efficiency of introduced π -linkers is evident from the graphs via NIR region absorption peaks. Compared to R, the smaller peaks for the designed molecules are also increased due to the high number of π - π^* transitions. The descending order of $\lambda_{\rm max}^{\rm cal}$ in solvent phase calculated for reference and designed molecules (DL1–DL7) is: DL2 > DL3 > DL4 > DL5 > DL1 > DL6 > DL7 > R. This also shows that the decreased band gaps lead to enhanced absorption maxima as a lower band gap facilitates excitation of electrons from HOMO (ground state) to LUMO (excited state).

The photovoltaic properties of molecules of the photochemically active layer also depend on excitation energy.⁴³ It is the amount of energy required for the transition of charge from a relaxed ground state to an excited state.44 For current generation, charge excitation is necessary as photocurrent is produced due to excited-state charges. The lower the excitation energy of charges, the higher the excitation of charges upon relaxation even for low energy radiations. 43,44 So lower excitation energy causes accelerated transitions between HOMO and LUMO, leading to more charge transfer, guaranteeing efficient photovoltaic properties. 44,45 The E_x values for the reference and DL1-DL7 are provided in Table 3. These calculated values show that the excitation energies range between 1.21 and 1.74 eV in the gas phase and 1.11 and 1.61 eV in the solvent phase. All the tailored molecules have smaller excitation energy values than R. which means these molecules have reasonable charge excitation potential. Furthermore, lower E_x in solvent phase indicates their capability of being solution-processable.

It can be noticed from Fig. 5 that molecules show significant absorption up to 1600 nm in a gaseous and solvent medium. The molecules possess multiple electronic transitions with different energies contributing to their absorption profile. The absorption at shorter wavelengths could be associated with

 Table 3
 B3LYP/6-31G(d,p) computed values for photovoltaic attributes of investigated structures in the gas phase and solvent phase

Molecule	$\lambda_{\max}^{exp}\left(nm\right)$	$\lambda_{\max}^{\operatorname{cal}}{}^{a}\left(\operatorname{nm}\right)$	$\lambda_{\max}^{\operatorname{cal}\ b}\left(\operatorname{nm}\right)$	$E_{\rm x}^{\ a}$ (eV)	E_{x}^{b} (eV)	$f_{ m os}{}^a$	f_{os}^{b}
R	720	675.24	731.69	1.8362	1.6945	2.9570	3.2324
DL1	_	772.38	772.38	1.6052	1.6052	2.6922	2.6922
DL2	_	1018.32	1112.86	1.2175	1.1141	1.9092	2.2414
DL3	_	1023.52	1094.78	1.2114	1.1325	1.5462	1.8899
DL4	_	966.91	1040.86	1.2823	1.1912	2.0930	2.4747
DL5	_	748.59	806.48	1.6562	1.5373	2.7547	3.0468
DL6	_	715.02	768.28	1.7340	1.6138	2.583	2.8693
DL7	_	709.70	765.72	1.7470	1.6192	2.7440	3.0080

^a Gas phase. ^b Solvent phase.

higher energy transitions. While the absorption at longer wavelengths can be related to lower energy transitions. 46,47 This broader absorption profile would allow the molecules to absorb light over various wavelengths.

In specific conjugated organic chromophores, excitonic effects can occur. Excitons are bound electron-hole pairs due to the coulombic interaction between an excited electron and a positively charged hole. These excitons can have lower energy than the calculated HOMO-LUMO gap and can contribute to the absorption profile at longer wavelengths.^{48,49}

Furthermore, the presence of multiple excited states can also explain the broadening of the absorption profile.⁵⁰ As these studies were carried out at multiple excited states, the absorption of light up to 1600 nm can be due to lower excited states.

Oscillator strength $(f_{\rm OS})$ measures the intensity of electromagnetic emission resulting from electronic transitions. Organic chromophores with higher oscillation strength show an enhanced probability for photon absorption resulting in higher light harvesting efficiency. For R and designed molecules, $f_{\rm OS}$ values in gas and solvent phases are provided in Table 3. Although R exhibited higher $f_{\rm OS}$ in both phases, $f_{\rm OS}$ for the tailored molecules is also sufficient for successful transitions. Among all the newly designed molecules, DL5 and DL7 show the highest oscillator strength with values of 3.0468 and 3.0080, respectively, giving them superiority over other molecules.

3.3 Natural transition orbitals (NTOs)

For systems with extended conjugation, the consideration of NTOs is required in addition to simple FMO analysis. NTOs provide a more reliable description of inter- or intramolecular charge transfers and quantitatively explain electronic transitions. The study of NTOs can be considered an alternative method to simplify the complex descriptions of transitions obtained through the transition density matrix. Thus, it also helps to visualize the distribution pattern of the electrons and holes in a molecule's excited state. In the case of NTOs, the occupied orbital is named a "hole" as an electron is ejected from it. In contrast, the virtual orbital is said to be an "electron" as an ejected electron from the hole is occupied by the virtual orbital.^{51,52}

For all our studied molecules, we selected only those transition states with high oscillation strength and were found to be the first transition states. Fig. S3 (ESI†) shows that the hole area is predominantly spread on the donor regions in all designed molecules. In contrast, the electron area is localized on the bridge and acceptor fragments of the molecules. Thus, upon excitation, the charges from the orbital of the donor are transferred to unoccupied orbitals located on the acceptor or π -linker. This process ensures that a well-defined ICT is present among various fragments of each molecule.

We also noted the highest eigenvalue of transitions for each molecule. This represents the electron transport contributions (ETC) and is essential for understanding the extent of transitions. The calculated values of ETC as percentages are illustrated in Table 5. DL1 and DL4 show the most efficient electronic transitions, with the highest ETC values of 99.9%.

The rest of the molecules also have better transport of electrons from HOMO to LUMO, as evidenced by their high value of transition characters, above 97% for all designed molecules. These data of electronic transitions from donor to acceptor regions confirm the high success rate of charge transfer in all these molecules, which is better for improving net efficiency.

3.4 Density of states

Results obtained by examining FMOs have been verified by DOS analysis. It was performed at B3LYP/6-31G(d,p) for the reference and structurally tailored (DL1–DL7) molecules. DOS depicts the contribution of various molecule fragments in forming the FMOs. 45,53 The DOS determines the total number of states that electrons can occupy at a specific energy level. The higher the number of states that are occupied by electrons, the higher the value for DOS; hence, the higher will be the electron density. 54 Thus, through DOS, we get insight into the contribution of the molecule's building blocks in constructing borderline molecular orbitals (FMOs). The DOS graphs for the reference and all designed molecules (DL1–DL7) are provided in Fig. 6.

The DOS graphs label the *y*-axis with the available states at specific energy levels, while the *x*-axis is labelled with three central values. Negative ones on the left side depict HOMO energy levels, and positive ones towards the right side represent LUMO. At the same time, the central region corresponds to the band gap between HOMO and LUMO.⁵⁵ The zigzag lined curves represent HOMO–LUMO charge density, and the area under these curves corresponds to possible excitation states for charge density. DOS calculations have the advantage that in addition to total DOS, they also give partial density of states (PDOS) for each molecule fragment.

While inspecting PDOS, the individual participation of each fragment of a molecule in FMOs can be observed.⁴⁴ On the other hand, the total DOS gives the total charge flow density of the whole molecule.⁴⁰ For PDOS, the reference and our newly developed molecules were portioned into three parts: acceptor (shown with black), donor (represented by the red line), and bridge (highlighted with green color in graphs). All these fragments collectively constructed the total DOS, represented with the blue dotted line. The fragment that strongly contributes to forming HOMO or LUMO will have greater electron density. Such a fragment gives a peak with a higher crest in the DOS graph in the HOMO or LUMO region.⁵⁶

The calculated values of DOS for each molecule involved in the current study are summarized in Table 4. If we examine the DOS data for the reference, it can be seen that the significant contribution for HOMO is from the donor part, *i.e.*, 60.1%. In comparison, the acceptor has a minor (25.7%) contribution to the formation of HOMO. The bridge has the lowest contribution (14.3%) in the elevation of HOMO. In raising LUMO, the acceptor and bridge participate effectively in LUMO, contributing 52.6% and 17.5%, respectively.

Similarly, for all our designed molecules, the HOMO levels have been raised mainly due to donor parts. In contrast, the rise in LUMO levels can be seen due to the higher percentage contribution of acceptors and bridges. All the π -bridges in our planned molecules seem to be better charge distributors than

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R DL1 Fermi level = -4.43 Fermi level = -4.14 DL2 DL3 Fermi level = -4.62 Fermi level = -4.67 DL4 DL5 Fermi level = -4.44 Fermi level = -4.38 DL7 DL6 Fermi level = -4.08 Fermi level = -4.10

Fig. 6 DOS graphs reveal the involvement of three fragments (donor, spacer, and acceptor) of the planned molecules (DL1–DL7) in raising HOMO and LUMO energy levels.

a double-bond-containing bridge present in R. These outcomes are supported through their tabulated percentage participation obtained from Mulliken's calculations shown in Table 4. Interestingly, in the case of DL2 and DL3, it can be seen that the contribution of spacer subunits is up to 60.6% in forming the total LUMO. This means these highly conjugated spacers efficiently collect charges from the central donor portion.

The conductive property of any part corresponds to its location. If a part is located near LUMO, its involvement in conductivity will be higher due to the many available states. As inserted π -spacers have more conjugation and more π - π * transition possibilities, these spacers and acceptors gave higher

Table 4 Percentage involvement of three fragments of the reference and planned molecules in the density of FMOs

	НОМО			LUMO		
Molecule	Acceptor (%)	Donor (%)	Linker (%)	Acceptor (%)	Donor (%)	Linker (%)
R	25.7	60.1	14.3	52.6	29.8	17.5
DL1	20.0	23.3	56.8	56.7	18.6	24.7
DL2	25.8	40.8	30.4	20.3	19.1	60.6
DL3	33.7	35.8	30.5	19.4	20.0	60.6
DL4	30.1	46.1	30.6	40.7	27.8	31.5
DL5	23.3	46.1	30.1	40.7	27.8	31.8
DL6	19.6	57.9	22.5	55.5	19.6	24.9
DL7	20.2	57.3	22.5	53.1	21.4	25.5

output than SJ-IC (R). As a result, LUMO peaks showed suitable intensities, confirming that replacing the double bond in R with the above-mentioned π -spacers is a fruitful strategy.

3.5 Dipole moment

The dipole moment vector provides information regarding a molecule's planarity and solubility in a desired solvent.¹⁹ The dipole moments for R and all designed molecules in the gas and solvent phases are provided in Table 5. These values were computed through the B3LYP/6-31G(d,p) level of DFT. Our compounds are mostly planar except for the phenyl ring substituents attached to the core. Their dipole moments are lower due to their linear geometry attained *via* optimization. Due to symmetrical geometry, opposite charges come in front of each other in opposite directions. Thus, they cancel each other's effect to some extent. So, as revealed for all our planned structures, low dipole moments, higher planarity and more conjugation result in higher absorption maxima.

It is a fact that a higher dipole moment is necessary for improved movement of charges.³⁷ The dipole moment values calculated in the solvent are slightly higher than those of the gas state. Interestingly, DL1 has a similar dipole moment in both gas and solvent phases, indicating that the solvent does not affect the dipole moment of this molecule. Such dipole moment vectors indicate that DL1-DL7 have positioned themselves to lower the charge recombination. Among all these molecules, DL6 shows the highest dipole moment in gas and solvent phases, with values of 2.17295 D and 2.72379 D, respectively. This shows that the DL6 molecule can have better charge mobility based on dipole moment than the other molecules.

The dipole moment also influences intramolecular charge transfer by affecting molecular fabrication. Two molecules with different dipole moments arrange such that their dipole moments are antiparallel. In this way, they create long tracks for the charges to move because their opposite faces attract each other due to the antiparallel alignment of their dipole moments. ^{57,58} So, a higher dipole moment of a molecule enhances its solubility along with improved crystallinity leading toward closer molecular assemblages. ⁴⁰ The notable thing about

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Table 5 Computed dipole moment (μ), binding energy (E_b), and transition character for R and DL1-DL7 in gas and chloroform solvent

Molecule	$\mu_{ m g}$ (D)	μ_{s} (D)	$E_{\mathbf{b}}^{b}$ (eV)	Interaction coefficient ^a	Interaction coefficient ^b	ETC^{a} (%)
R	0.04380	0.02930	0.316	0.7058	0.7025	99.6
DL1	0.04160	0.04160	0.158	0.7069	0.7069	99.9
DL2	0.09180	0.08630	0.224	0.7062	0.7036	99.7
DL3	0.02460	0.13240	0.218	0.7031	0.7016	98.9
DL4	0.01071	0.04260	0.215	0.7069	0.7035	99.9
DL5	0.03478	0.05622	0.318	0.7045	0.7004	99.3
DL6	2.17295	2.72379	0.325	0.7001	0.6973	98.0
DL7	0.56570	0.67680	0.337	0.7058	0.6999	99.1

^a Gas. ^b Chloroform solvent.

our planned molecules is that they all have improved dipole moments compared to R. Thus, they are well compatible with chloroform solvent.

3.6 Exploration of physiochemical properties through an MEP map

To reveal the physiochemical properties of molecules such as three-dimensional electron density, reactive sites, and nucleophilic and electrophilic points,59 their molecular electrostatic surface potentials (MESPs) are portrayed. This study gives information about charge transfer across a molecule in the excited state.35 MESP surfaces appear like a cloud with three primary colors. Each color code corresponds to the specific strength of the electrostatic potential of the molecule.

Red shades represent negative electrostatic potential. The red portion of the molecule is electron-rich, a possible attack site for electrophiles, and attractive points for protons due to aggregated electron density at these regions. 60 In comparison, cyan and blue shades represent positive electrostatic potentials. It symbolizes that these areas correspond to electron-deficient sites, susceptible to nucleophilic attack, and repulsive areas for protons due to the atomic nuclei of molecules at these sites. In comparison, sites shaded with green represent areas with neutral electrostatic potential.

A color scale is also formed along with each MEP cloud with deep red color at one end, moving towards deep blue color at the other end.37 MEP plots for R and the structurally engineered molecules are displayed in Fig. S4 (ESI†). Fascinatingly, it can be seen that more red color exists at the acceptor regions of DL2, DL3, and DL5 molecules than the other molecules. This shows enhanced charge separation in these molecules depicting that these newly planned molecules can be efficiently designed for OSCs.

MEPs for the reference and designed molecules show cyan colors on the sides of the core, methyl groups of linkers, and phenyl rings of acceptors. Thus, these areas are susceptible to nucleophilic attack. In contrast, the negative potential represented by the red color is present on the acceptor and π -linkers having N, O, and S atoms. It shows that these regions provide a robust and attractive force for electrons and are responsible for better charge delocalization. But if we look at those N, O, and S atoms of various bridges facing towards the core, they do not show negative potential there, although they are still

electronegative. The reason is the canceling effect produced by the positive potential of hydrogens and carbons of the core located in their vicinity. There is no high prevalence of blue color (positive electrostatic potential), while various areas of negative potential exist. So, it can also be inferred from obtained MESP surfaces that each of our planned molecules has the potential to behave as an acceptor molecule in the active layer of OSCs.

3.7 Chemical reactivity

There are five main parameters whose values give an idea about the reactivity of acceptor and donor molecules. These parameters also specify the charge transfer direction from one molecule to another. Namely, these are chemical potential (μ) , chemical softness (S), chemical hardness (η), electrophilicity index (ω) , and electronegativity (χ) . We calculated these values using eqn (4)-(8) to get an insight into chemical reactivity. 61-63

For an acceptor, the chemical potential (μ) is the tendency to acquire a negative charge. For a donor, it is the electrondonating capacity of the molecule. The chemical potentials calculated for DL1-DL7 through eqn (4) are provided in Table 6.

$$\mu = \frac{(E_{\text{HOMO}} + E_{\text{LUMO}})}{2} \tag{4}$$

PTB1, PCPDTBT, and PTB7-Th are selected as donors for our structurally engineered molecules and have a chemical

Table 6 Computed chemical potential (μ) , hardness (η) , softness (S), electronegativity (χ), and electrophilicity (ω) obtained through B3LYP/ 6-31G(d.p) level

Molecule	μ (eV)	η (eV)	$S\left(\mathrm{eV}^{-1}\right)$	χ (eV)	ω (eV)
R	-4.43	1.01	0.99	4.43	9.79
DL1	-4.14	0.88	1.13	4.14	9.75
DL2	-4.62	0.67	1.49	4.62	16.01
DL3	-4.66	0.67	1.48	4.67	16.14
DL4	-4.43	0.70	1.42	4.44	14.01
DL5	-4.38	0.93	1.08	4.38	10.38
DL6	-4.07	0.97	1.03	4.08	8.64
DL7	-4.09	0.98	1.02	4.09	8.58
PTB1	-4.05	0.85	1.17	4.05	9.64
PCPDTBT	-4.42	0.72	1.38	4.42	661
PTB7-Th	-4.40	0.80	1.25	4.40	12.10

potential of -4.05 eV, -4.42 eV, and -4.40 eV, respectively. PTB1 is selected for the reference, DL1, DL6, and DL7. PCPDTBT is selected for DL2, DL3, and DL4. PTB7-Th is selected for DL5 due to better matching of HOMO and LUMO energy levels. Except for DL5, all the other newly designed molecules have better chemical potential values than the donor used for them. DL2 and DL3 have significantly higher chemical potential than the corresponding donors, meaning that DL2 and DL3 are promising acceptor molecules.

Chemical potential further depends on the chemical softness (S) and hardness (η), which are inversely interlinked. Table 6 illustrates these parameters, which we obtained through eqn (5) and (6):

$$\eta = \frac{(E_{\text{LUMO}} - E_{\text{HOMO}})}{2} \tag{5}$$

$$S = \frac{1}{\eta} \tag{6}$$

Higher softness is responsible for smooth electron transfer due to low resistance. As evident from softness values in Table 6, all our newly developed molecules (DL1–DL7) show higher softness than the reference. DL2, DL3, and DL4 show a remarkable improvement in softness with values of 1.49 eV, 1.48 eV, and 1.43 eV, respectively. This shows the probability of enhanced charge transfer rate in these designed molecules making our strategy of inserting conjugated π -linkers successful. All other newly designed molecules also show improvement in softness by lowering the hardness. This might be because these π -linkers have enhanced interactions of various molecular parts, resulting in an increase in ICT.

Electrophilicity (ω) and electronegativity (χ) explain excessive electron flow. The direction of electron flow depends on the values obtained from eqn (7) and (8):

$$\chi = \frac{(E_{\text{HOMO}} + E_{\text{LUMO}})}{2} \tag{7}$$

$$\omega = \frac{\chi^2}{2(\eta)} \tag{8}$$

Table 6 illustrates the higher tendency of DL2, DL3, and DL4 for electrons with higher ω values of 16.0, 16.1, and 14.0 eV as compared to the other molecules. Similarly, the high electronegativity of the same molecules represents their better electron-accepting ability from donor molecules. These aspects of the mentioned molecules make them a superior choice for the development of efficient OSCs in the future.

3.8 Examination of charge transfer

3.8.1 Binding energy. The large binding energy between excitons is a bottleneck for developing efficient organic photovoltaic devices. When a photon impinges on an organic photoactive acceptor material, an electron-hole pair (exciton) is generated instead of forming separate electrons and holes confined in space. Dissociation of an exciton requires

overcoming the coulombic attraction between the electron and hole. For this purpose, an exciton diffuses into a donor/acceptor interface where the difference in electron affinity of the donor and acceptor provides a driving force for the conversion of the exciton into free charges. These free charges approach their respective electrodes by moving along the acceptor and donor and generating electricity. ⁶⁴ The minor coulombic barrier between excitons results in lower energy required for successful exciton dissociation. If E_g is the band gap between permitted (HOMO–LUMO) bands and E_x is the optical energy gap, then the exciton binding energy that stabilizes this exciton is given by eqn (9):

$$E_{\rm b} = E_{\rm g} - E_{\rm x} \tag{9}$$

Table 5 illustrates the values of the binding energy of excitons for the proposed molecules with values of 0.158 eV, 0.224 eV, 0.218 eV, 0.215 eV, 0.318 eV, 0.325 eV, and 0.337 eV for DL1–DL7 respectively. On the other hand, the binding energy of R is 0.316 eV. DL1 shows the lowest value of exciton binding energy. DL2, DL3, and DL4 also show lower exciton binding energy than the reference molecule. This means that these molecules have improved potential for the dissociation of excitons. Thus, based on the results of exciton $E_{\rm b}$, it can be proposed that these molecules might have the potential for significant charge mobility.

3.8.2 TDM. To explore the electronic behavior of a compound, TDM analysis is carried out because, through TDM maps, we can visualize the interactions between building blocks of molecules. The TDM also explains the locations of electrons and holes and elaborates on electron excitations in the excited state. All electronic transitions of the proposed molecules were analyzed by computing their TDM by applying the B3LYP level of theory with a 6-31G(d,p) basis set. Then this information was converted into TDM maps through multiwavefunction (Multiwfn 3.8) software. These TDM plots were portrayed while neglecting H atoms because they show minute contributions in electronic transitions. For the convenient study of transitions, the TDM graph for each molecule was sectioned into three parts (A = acceptor, π -L = linker, D = donor) as shown in Fig. 7.

Atom numbers are displayed on the bottom x-axis and left yaxis. The electron density coefficient in the colored scale is also displayed for each molecule on the right y-axis. The bright blue-green areas against dark blue background symbolize charge transfer. From TDM plots for all of our proposed molecules, it can be observed that the electrons are distributed throughout the molecules due to extended conjugation. It can be observed that there is coherence in the regions of TDM maps which are specific for atoms of π -bridges and acceptor, separately. This confirms that π -linkers further facilitated charge transfer from one edge of the molecule to the other edge without charge trapping. Here π - π * transitions are represented through the coherence of electrons along the diagonal pathway. Like R, some core regions lack electronic transitions in all designed structures. Moreover, similar charge distribution behavior was revealed for R and the

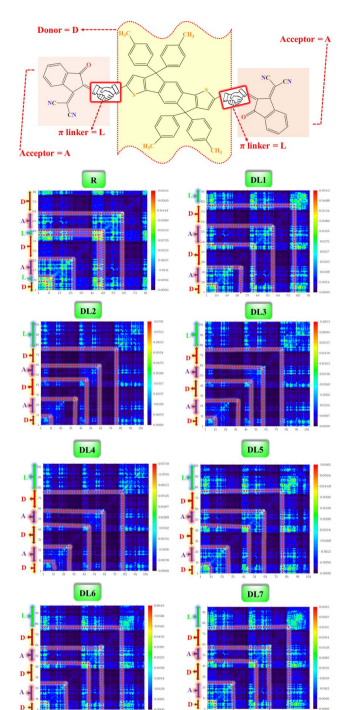


Fig. 7 $\,$ TDM heat maps for R and DL1–DL7 with red dotted lines partitioning the fragments of molecules.

designed molecules, which shows the effectiveness of inserting π -linkers. From in-depth observations, it can be seen that there exist more electron interactions in the acceptor and spacer region of DL1 and DL5 molecules, indicated by more bright spots in the maps of these molecules.

Dissociation of excitons and capability of charges to migrate show inverse relation with interaction coefficient. A low interaction coefficient leads to fewer interactions among charges. Hence, molecules with low interaction coefficients have a high potential for the mobility of charges in various parts of the molecule. The interaction coefficient of all the studied molecules varies in chloroform solvent in increasing order as: DL6 < DL7 < DL5 < DL3 < R < DL4 < DL2 < DL1.

DL6 and DL7 have small interaction coefficients; hence, the easiest charge transfer can occur in them in contrast to the others. However, it can be observed that a tiny difference exists in the values of all molecules' interaction coefficients. So, all these molecules have the probability of only a few interactions among charges which leads to a higher charge transfer rate in all these molecules. The cyanide groups present in the acceptor of each molecule improve the electron-withdrawing effect. These acceptor subunits possess electronegative nitrogen, sulfur, and oxygen atoms sp-hybridized in π -linkers. In conjunction with conjugated π -linkers, the cyanide groups further increase the conjugation of tailored structures. Hence, interactions among excitons are lowered, and excitons are easily separated, as evident from small values for the interaction coefficient of all designed molecules.

3.8.3 Ionization potential (IP) and electron affinity (EA). Energy for dissociation of charge transfer into free charges is one of the fundamental factors determining a device's performance in OSCs. It can be calculated by taking the difference between the acceptor's EA and the IP of the donor (IP $_{\rm D}$ – EA $_{\rm A}$). ^{65,66} Thus, IP and EA are essential parameters to estimate the charge transport ability. ⁶⁷

High EA is concerned with better electrophilicity and improved electron transport and injection. Molecules with high EA are used as efficient electron acceptors because they can take up electrons easily. However, donor materials should have low IP to donate electrons easily. Electron-donating moieties attached to a molecule cause a decrease in IP because HOMO levels are destabilized by facilitated electron transfer by these electron-donating groups. However, molecules having electron-withdrawing groups attached to them exhibit elevated IP because of the stabilization of HOMO, and electron removal becomes challenging.

We calculated IP and EA according to eqn (10) and (11), respectively,⁵⁵ and summarize the results in Table 7:

$$IP = [E_0^+ - E_0] \tag{10}$$

Table 7 B3LYP/6-31G(d,p) computed values for reorganization energies of hole (λ_h) and electron (λ_e) alongside calculated electron affinity (EA) and ionization potential (IP)

Molecule	$\lambda_{\rm h}$ (eV)	λ_{e} (eV)	EA (eV)	IP (eV)
R	0.185	0.209	2.656	6,211
DL1	0.188	0.129	2.536	5.766
DL2	0.058	0.156	3.249	5.962
DL3	0.139	0.154	3.274	6.009
DL4	0.139	0.160	3.036	5.807
DL5	0.161	0.182	2.753	6.013
DL6	0.195	0.154	2.379	5.788
DL7	0.186	0.147	2.396	5.814

$$EA = [E_0 - E_0^-] \tag{11}$$

All newly developed molecules (DL1–DL7) showed lower IP because their HOMO levels are at higher energy onsets than that of the reference. The least value of IP was shown by DL1 having an IP of 5.766 eV. Similarly, DL2 (3.249 eV), DL3 (3.274 eV), DL4 (3.036 eV), and DL5 (2.753 eV) showed significantly improved EA compared to R (2.656 eV). This is because the inserted $\pi\text{-linkers}$ played a role in stabilizing the LUMOs, making electron injection easier. Hence, these are expected to have smooth electron flow.

3.8.4 Reorganization energy. Non-radiative recombination is an important parameter that affects energy losses in organic photovoltaic cells. Non-radiative recombination is associated with reorganization energy. So, in our research work, reorganization energy is one of the necessary calculations which pave a path to obtaining an efficient OSC. The Marcus theory of charge transfer can calculate the rate of charge transfer. The critical element of the Marcus model is reorganization energy $(\lambda_{h/e})$ which represents the energy required for geometry relaxation during charge transfer. The reorganization energy is divided into two forms. One is internal reorganization energy related to electronic states in relaxed and charged geometries. The other is outer-sphere reorganization energy which is concerned with the effect of solvent, polarization, temperature, and other external factors. Since we used similar conditions in our calculations and are mainly concerned with the intramolecular transfer of charge carriers, we only focused on the internal RE (λ_{int}) . The external RE (λ_{ext}) is not calculated as it relates to the outer environment and polarization effects produced during charge transfer and is negligibly related to charge transmission.³¹ The λ_{in} is a combination of two reorganization energies, namely hole reorganization (λ_h or λ_+) and electron reorganization (λ_e or λ_-) energy (*i.e.*, $\lambda^{in} = \lambda_h + \lambda_e$).⁶⁹ Eqn (1) and (2) are used to calculate the hole transfer λ_h and electron transfer λ_e rates, respectively, and results are summarized in Table 7.

The sum of ground- and excited-state reorganization energies (i.e., $\lambda_1 + \lambda_2$) gives λ_h as depicted in Fig. S5 (ESI†). λ_1 (reorganization energy in ground state) equals the difference of energies of optimized neutral state and cationic geometry. On the other hand, λ_2 represents the excited-state reorganization energy which is calculated as the energy difference of cationic energy of optimized molecule in charged state and neutral state geometry.

Charge mobility is inversely dependent on reorganization energy. Those molecules with lower reorganization energy have higher charge mobilities because they need a small driving force for exciton dissociation. A high value of reorganization energy results in non-radiative recombination. Our calculations revealed a decrease in reorganization energies for most of our structurally engineered molecules compared to the reference. These results are fruitful for enhancing exciton lifetime, improving charge transfer, and suppressing non-radiative recombination of charges. DL1 showed the lowest $\lambda_{\rm e}$ (0.129 eV), while DL2 showed minimum $\lambda_{\rm h}$ (0.058 eV). The $\lambda_{\rm h}$ of all the

molecules increases in the order: DL2 (0.058 eV) < DL3 = DL4 (0.139 eV) < DL5 (0.161 eV) < R (0.185 eV) < DL7 (0.186 eV) < DL1 (1.88 eV) < DL6 (1.95 eV). On the other hand, all newly designed molecules show lower λ_e than the reference molecule. This means all these newly developed molecules have significantly improved electron transfer potential compared to R. The added π -linkers successfully reduced the energy barrier by conjugation, and as a result, charge mobility increased.

3.9 Light harvesting efficiency

Light harvesting efficiency (LHE) is another significant parameter that shows the photocurrent response of a molecule by absorption of radiation. When the LHE of a molecule is more significant, it will produce a high quantity short circuit current and intensify the molecule's photocurrent response. LHE is determined through oscillator strength (f_{OS}) which measures the emission/absorption of radiation during the transition of electrons from one energy level to another. LHE of R and DL1–DL7 is presented in Table 8. It is calculated through eqn (12):⁷⁰

$$LHE = 1 - 10^{-f} \tag{12}$$

Eqn (12) shows the direct dependence of LHE on oscillator strength $f_{\rm OS}$. A higher value of oscillator frequency of a molecule intensifies the absorption peak of that molecule. ⁵⁴ All our modelled molecules have comparable oscillator frequencies, and hence due to the direct relation of $f_{\rm OS}$ with LHE, their LHEs also lie in a similar range. Among all the newly designed molecules, DL5 shows a maximum LHE value (0.991 eV), which is slightly better than those of the other designed molecules. These comparable values of LHE suggest that all these molecules can perform efficiently in OSC devices.

3.10 Open circuit voltage

Among the technical parameters which are most challenging to improve, open circuit voltage $(V_{\rm OC})$ is a crucial one. When the current flow is zero, the voltage value attained across a photovoltaic device is called open-circuit voltage. A higher value of a system's $V_{\rm OC}$ leads to higher photocurrent conversion efficiency. To theoretically estimate the $V_{\rm OC}$ of any photovoltaic system, Scharber's equation is used, which is given as:

Table 8 Calculated open-circuit voltage ($V_{\rm OC}$), normalized $V_{\rm OC}$, fill factor (FF), and light harvesting efficiency of all molecules

Molecule	ΔE (eV)	$V_{\rm OC}$ (eV)	Normalized $V_{\rm OC}$	FF	LHE
R	1.48	1.18	61.8357	0.9183	0.9994
DL1	1.64	1.34	68.0193	0.9242	0.9971
DL2	1.20	0.9	41.3527	0.8881	0.9943
DL3	1.16	0.86	39.8068	0.8848	0.9871
DL4	1.42	1.12	49.8551	0.9032	0.9967
DL5	1.75	1.45	60.6763	0.9170	0.9991
DL6	1.79	1.49	73.8164	0.9290	0.9987
DL7	1.78	1.48	73.4300	0.9287	0.9990

$$V_{\rm OC} = \frac{1}{e} \left[E_{\rm HOMO}^{\rm donor} - E_{\rm LUMO}^{\rm acceptor} \right] - 0.3 \tag{13}$$

where $E_{\rm HOMO}^{\rm donor}$ is the energy of the HOMO level of the donor, $E_{\rm LUMO}^{\rm acceptor}$ is the energy of the LUMO level of the acceptor, and e is an elementary charge value of a point molecule which is usually taken as constant '1'. Whereas 0.3 is an empirical factor compensating for the usual losses occurring in a BHJ. The 0.3 factor is kept constant for all the molecules to get a comparison of their efficiency under similar conditions. In this research work, we selected different donors for designed acceptors due to the compatibility of HOMO–LUMO energy levels. The $V_{\rm OC}$ of the reference molecule and DL1, DL6, and DL7 acceptors was calculated by combining them with PTB1 donor due to good matching of their HOMO–LUMO energy levels. Furthermore, DL2, DL3, and DL4 were combined with the PCPDTBT donor molecule, and DL5 with the PTB7-Th donor for theoretical calculations of $V_{\rm OC}$.

Hence, we calculated the $V_{\rm OC}$ by computing the difference between the HOMO of the donor and the LUMO of designed acceptors (DL1–DL7), and the obtained results are portrayed in Fig. 8. The HOMO–LUMO of PTB1, PCPDTBT, and PTB7-Th lie at 4.90/3.20 eV, 5.15/3.70 eV, and 3.60/5.20 eV, respectively, as reported in the literature.^{72–74} The difference between the energy

of HOMO and LUMO of donor and acceptor material is known as the electrochemical potential difference (ΔE), which signifies the interlayer bandgap.^{72,73} The tabulated values of ΔE of all molecules are also mentioned in Table 8.

It can be observed that there is a negative offset of the HOMO and LUMO levels between the donor and our acceptor molecules. A negative offset of HOMO means that the HOMO level of the acceptor is higher than that of the donor, *i.e.*, in the case of PCPDTBT:DL4 blend. In this case, charge transfer is governed mainly by a hole transfer mechanism. Facilitated hole transfer occurs when a hole in the donor material is first transferred to an intermediate state. Then the hole can be transferred from the intermediate state to the acceptor material. Once the hole is transferred to the acceptor material, it can move from the acceptor material towards the electrode, generating a photocurrent.^{74,75}

On the other hand, when the LUMO of the acceptor molecule is energetically lower than the LUMO of the donor molecule, a negative LUMO offset is created. The negative LUMO offset between the donor and acceptor molecules facilitates charge transfer by providing an energy gradient for electron movement. This offset ensures that the electron can quickly move from the donor to the acceptor, minimizing energy losses and enhancing the overall efficiency of the OSC. After absorbing

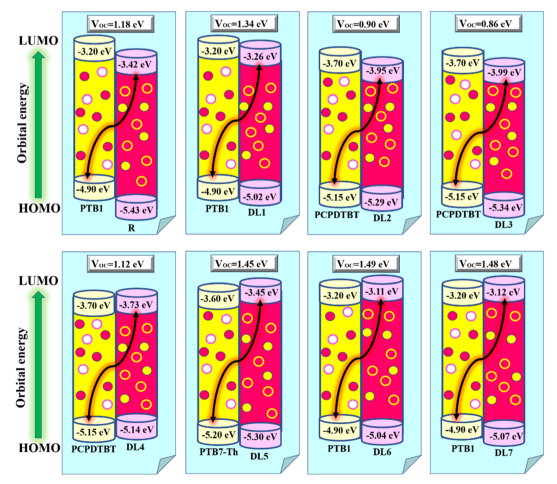


Fig. 8 $V_{\rm OC}$ of designed acceptors (DL1-DL7) against donor J61.

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light energy, electrons in the donor molecule are excited from the ground to a higher energy state, leaving behind a positively charged hole. The excited electron in the donor molecule absorbs enough energy to overcome the energy barrier created by the negative LUMO offset. It can then transfer from the donor molecule to the acceptor molecule. Once the electron is transferred to the acceptor molecule, it can move through the acceptor material, eventually reaching the electrode. Simultaneously, the positively charged hole left behind in the donor molecule can move in the opposite direction towards the other electrode. The movement of electrons and holes toward their respective electrodes creates an electric current. A negative offset of the LUMO energy level creates a separation of positive and negative charges, which is essential for the functioning of a solar cell.

Table 8 shows that our designed molecules DL1, DL5, DL6, and DL7 have higher $V_{\rm OC}$ with values of 1.34, 1.45, 1.49, and 1.48 eV, respectively, in comparison to the $V_{\rm OC}$ of R (1.18 eV). The remaining proposed acceptors also show significant $V_{\rm OC}$, enough for electron transfer from the respective donors to the acceptors (DL1–DL7). The reason for better $V_{\rm OC}$ is the low-lying HOMOs of donor materials while high-lying LUMOs of all planned NFAs. This increases the bandgap of the active layer, resulting in improved open circuit voltage. Lumber 12 to a be noticed that the $V_{\rm OC}$ of all these newly designed molecules is less than the active layer bandgap between the donor and acceptor molecules. This enhances the extent of their experimental applications. Due to this remarkable improvement in $V_{\rm OC}$, these newly designed molecules (especially DL6) should be considered for making advanced OSCs in the future.

3.11 Fill factor

The fill factor directly influences the light-to-electricity conversion efficiency of any photovoltaic device. It is mainly dependent on open circuit voltage. The higher the value of $V_{\rm OC}$, the higher will be the fill factor, and, consequently, the PCE of the device will also be higher. The ideal and perfect value for the fill factor is unity (*i.e.*, 100%).⁸³ We employed eqn (14) to calculate the fill factor of R and all optimized planned molecules:^{84,85}

$$FF = \frac{V_{OC} - \ln(V_{OC} + 0.72)}{V_{OC} + 1}$$
 (14)

in this equation, normalized open circuit voltage is represented by $\frac{V_{\rm OC}}{k_{\rm B}T}$. Normalized $V_{\rm OC}$ is used for the theoretical calculations of FF. $V_{\rm OC}$ represents open circuit voltage and $k_{\rm B}$ is Boltzmann's constant (8.61733 \times 10⁻⁵ eV K⁻¹).

All the proposed molecules give FF values that are close to the ideal value for FF with 0.9242 for DL1, 0.8881 for DL2, 0.8848 for DL3, 0.9032 for DL4, 0.9170 for DL5 while 0.9290 and 0.9287 for DL6 and DL7, respectively, as evident from Table 8. DL1, DL6, and DL7 show remarkable improvement in FF compared to the reference SJ-IC. There can be several reasons that can lead to improved fill factor in OSCs. One theory suggests that optimizing the charge transport and reducing the recombination losses cause a higher fill factor. ⁸⁶ All the newly

designed molecules have the potential for improved charge transport due to their lower reorganization energy than the reference. This can improve the fill factor in the newly designed molecules. Another scenario involves using low-bandgap polymers or small molecules, which can broaden the absorption spectrum and increase the open-circuit voltage, resulting in a higher fill factor. Enhanced light harvesting and increased light absorption in the active layer can improve the fill factor. As indicated from the absorption profiles of these studied molecules, newly designed molecules have broader absorption spectra which can enhance the fill factor of these molecules. Therefore, these molecules should be kept in focus for making efficient solar cells in the future as they emerge as potential candidates as a result of this study.

3.12 Short circuit current density

Short circuit current density $(J_{\rm SC})$ is an essential factor that influences the photovoltaic performance of solar cells. It measures the maximum current density that can be generated by a solar cell in the absence of any external resistance. Depending on several factors, $J_{\rm SC}$ is the highest current that can be drawn from the solar cell. It is mainly affected by the number of photons, the incident light spectrum, the absorption coefficient of the active layer material, the charge carrier mobility, and the electron and hole diffusion lengths. 89,90

A higher $J_{\rm SC}$ value generally results in a higher overall PCE of a solar cell. It is a direct measure of the quantity of current flowing through a circuit so that it can be accurately measured only by experimental methods. However, measuring other factors that affect $J_{\rm SC}$ makes it possible to predict the improvement or decrement of $J_{\rm SC}$ for the designed molecules compared to the reference molecule. It is noticed that higher absorption in the active layer leads to a higher $J_{\rm SC}$. Similarly, higher mobility allows for more efficient extraction of photogenerated charges, resulting in a higher $J_{\rm SC}$.

The $J_{\rm SC}$ value of the SJ-IC molecule is 13.70 mA cm $^{-2}$.¹⁹ It can be observed that our newly proposed molecules show a significant improvement in light absorption owing to a broader absorption profile. This can lead to improved short-circuit current density in these proposed molecules. Similarly, higher charge mobility in the newly proposed molecules due to low reorganization energy can also improve the $J_{\rm SC}$ of these molecules. It makes these molecules more suitable for developing efficient OSCs in the future.

4. Conclusion

In this quantum chemical approach, we studied the effect of inserting various π -linkers to reveal photovoltaic properties of SJ-IC. The conceptualized molecules outperformed the reference SJ-IC in the context of photovoltaic properties. All newly designed molecules exhibited significantly reduced band gap (E_g) between their respective FMOs in contrast to R. Furthermore, they show notably higher λ_{\max}^{cal} , *i.e.*, up to NIR in absorption spectra with broadened FWHM, thus having a better ability to harvest light. We studied various global descriptors,

including chemical potential (μ) , electronegativity (χ) , chemical hardness (η) , softness (S), along with electrophilicity (ω) . The improved photovoltaic properties of designed molecules make them promising for use in OSCs. Lower excitation energies shown by the tailored molecules lead to easier charge excitation with a considerably better mobility rate for charges. The presence of NCI played the role of conformational locks to retain planarity, which was confirmed through NCI graphs and isosurfaces. The main challenge for OSCs is to achieve higher open-circuit voltage. Our results reveal that DL1 and DL6 can stand as the best acceptors out of those studied. The $V_{\rm OC}$ values of these molecules range from 0.86 to 1.49 eV. The extent of improved charge mobility and photovoltaic efficiency shown by the newly designed molecules is remarkable. However, for conducive electron injection, these $V_{\rm OC}$ values are considered acceptable. Lastly, the study of the J61:DL6 complex verified the acceptor nature of our designed molecules and confirmed the efficient charge transfer from donor to acceptor. Overall our research suggests that all the newly tailored molecules are efficient acceptor materials in contrast to the reported reference SJ-IC.

Conflicts of interest

Authors have no conflict of interest.

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