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New organic dye-sensitized solar cells based on the D-A- π -A structure for efficient DSSCs: DFT/TD-DFT investigations

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Global energy consumption has increased due to population growth and economic development. Solar energy is one of the most important renewable energy sources for human consumption. In this research, four novel organic dyes (D2–D5) of the D–A– π –A structure based on triphenylamine (TPA) were studied theoretically using DFT and TD-DFT techniques for future usage as dye-sensitized solar cells (DSSCs). The effects of modifying the π -spacer of the reference molecule D1 on the structural, electronic, photovoltaic, and optical characteristics of the D2–D5 dyes were studied in detail. D2–D5 exhibited band gaps ($E_{\rm gap}$) in the range from 1.89 to 2.10 eV with $\lambda_{\rm abs}$ in the range of 508 to 563 nm. The results obtained show that modifying the π -spacer of the dye D1 increased its hole injection and reinforced the intramolecular charge-transfer (ICT) impact, which resulted in a red-shifted ICT absorption with a greater molar extinction coefficient. The theoretically calculated open-circuit voltage ($V_{\rm oc}$) values ranged from 0.69 to 1.06 eV, while the light-harvesting efficiency (LHE) values varied from 0.95 to 0.99. Indeed, this theoretical research could guide chemists to synthesize effective dyes for DSSCs.

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Introduction

Global energy consumption has expanded rapidly due to economic and population development but has been accompanied by the increasing consumption of non-renewable energy sources and increased environmental pollution. Therefore, resorting to renewable and low-cost energy sources that can help reduce environmental pollution emissions has become the most appropriate option for future energy needs.1-3 Recently, dye-sensitized solar cells (DSSCs) (or Grätzel cells) have been widely used as a new alternative to meet the growing energy demand for safe environmentally friendly, natural, and economical resources, 4-6 allowing the opportunity for converting solar energy into electricity with an efficiency of more than 13%.7 The engineering design of DSSCs is based on using photosensitive materials in solar cells; generally, a DSSC includes four main components: the photoanode,8 the cathode,9 the electrolyte, and the sensitizer.10 The photoanode of a DSSC consists of a thin layer of nanocrystalline metal oxide

⁽e.g. titanium dioxide (TiO₂) or zinc oxide (ZnO)) deposited on a glass substrate.11 Natural or artificial sensitizers are critical components of DSSCs and play a fundamental role in extending their absorption spectra. 12 In this sense, different organic, organometallic, and natural dyes can be utilized as sensitizers.13 The electrolyte of a DSSC consists of a redox couple designated as I⁻/I₃⁻ (iodide/triiodide), closed by a counter-electrode typically composed of platinum (cathode).14 In general, most used organic dyes for DSSCs are artificial sensitizers that possess the donor- π -spacer-acceptor (D- π -A) molecular architecture, which is intended to provide intramolecular charge transport (ICT) from the donor unit (D) to the acceptor group (A) via the π spacer. 15-17 Other potential structures of artificial sensitizers, including D-A- π -A, ¹⁸⁻²⁰ D-D- π -A, ^{21,22} (D- π -A)₂, ^{23,24} D- π -A- $A_{25,26}$ and $A-\pi-D-\pi-A_{27,28}$ have also been established and gained significant importance due to their high efficacy,29 but the most common structure is $(D-\pi-A)$ because it facilitates electron movement, which improves the efficiency of dyes and allows them to be used in DSSCs.30-32 Since the efficiency of a DSSC cell is highly dependent on the photosensitive dye used in the cell, the following points can be used when designing the cell to improve and raise the efficiency of DSSCs: (a) it must include the most appropriate π -spacers between the electron donor and the acceptor; (b) it is important to have a broad and strong absorption band capable of covering practically all the visible and near-infrared spectral region (NIR); (c) appropriate redox and oxidation potentials of the compound should be

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these sensitizers, PT2 (reference molecule), which we named in investigated to ensure regeneration; (d) a high electron this article as D1, demonstrated a good performance of 6.11% connection is necessary between the molecules and conduction $(V_{\rm oc} = 0.668 \text{ V}, J_{\rm sc} = 12.61 \text{ mA cm}^{-2}, \text{ and FF} = 0.74), \text{ using TPA}$ band of the semiconductor (TiO2) to accelerate the transfer of electrons from the dye to TiO2.33-35 It is essential to study the as the donor, [1,2,5]-thiadiazolo[3,4-c]-pyridine (PT) as the electronic and optoelectronic characteristics of sensitizers for supplementary acceptor fragment, benzene as the π -bridge, and DSSC devices.³⁶ Also, the highest occupied molecular orbital cyanoacetic acid as the electron-acceptor moiety (Fig. 1). The π -(HOMO) level of the dye being examined should be less than the bridge plays an essential role in the rapid transfer of electrons redox couple energy level (iodide/triiodide I⁻/I₃⁻) for the dye to for the DSSC charge; due to this, it should have the following be successfully regenerated.37 In addition to allowing the elecfeatures: easy charging, prolonged π -spacer that diminishes tron to be injected into the anode quickly, the lowest occupied a large part of the light-absorption spectrum, and it should facilitate the absorption capacity. 70,71 Hence, choosing a suitable molecular orbital (LUMO) should be above the conduction band (CB) of TiO2.38 Recent research introduced an electron auxiliary π -bridge is one of the most important steps in building highacceptor into the conjugated system between the electron-donor performance photovoltaic cells. Several π -bridges have been moiety and π -linker and used this in the design of a new implemented experimentally and theoretically to fabricate architecture known as the D-A- π -A system, ³⁹⁻⁴² where sensinovel sensitizers.72 tizers are coupled with auxiliary acceptor units, such as quinoxaline, 43-46 benzothiadiazole, 47-49 thieno[3,4-c]-pyrrole-4,6-

PT2 was used to investigate the efficiency of D–A– π –A structural molecules of thiophene derivatives as the π -bridge (Fig. 1) by using theoretical DFT and TD-DFT methods to explain the influences of the π -spacer compounds D2–D5. By finding the critical theoretical elements, a high current charge density ($J_{\rm sc}$), open-circuit voltage ($V_{\rm oc}$), light-harvesting efficiency (LHE), reorganization energy ($\lambda_{\rm total}$), electronic injection driving force ($\Delta G^{\rm inject}$), and driving force of regeneration ($\Delta G_{\rm reg}$) were determined. Furthermore, other parameters, such as the energy levels, frontier molecular orbitals (FMOs), natural bond orbital (NBO), nonlinear optical properties (NLO), density of states (DOS), chemical indices, and absorption spectra were also considered. These theoretical parameters help to elucidate the relationship between the molecular structure and performance, which aids selecting better dyes.

Computational section

Several computational parameters can influence the functioning of DSSCs, including the power conversion efficiency (η), and this parameter depends on several variables, including the values of the V_{oc} , J_{sc} , fill factor (FF), and the incident solar power (P_{inc}):⁷³

Fig. 1 Chemical structures of the proposed molecules in this work, D_i (i = 1-5).

dione, 50,51 benzotriazole, 52-54 and diketopyrrolopyrrole. 55,56

Indeed, the D-A- π -A system has shown multiple advantages,

including reducing electron recombination, minimizing

aggregation, adjusting the molecular energy levels, causing

a red-shift of the absorption band, and increasing the photovoltaic efficiency and stability;^{57–59} whereas several organic dyes

of triphenylamine-based compounds have been used as elec-

tron donors due to the excellent light-harvesting ability of tri-

phenylamine (TPA)-based molecules, their capacity to control

the energy levels of molecules, promote charge generation, and

facilitate the formation of intermolecular interactions. 60-64 For

DSSCs, several electron-acceptor materials have been developed

and synthesized.65-67 However, cyanoacrylic acid is often

employed as an electron-acceptor material due to its high electron-attracting capacity and solid binding to the TiO₂

surface, which facilitates electron injection.⁶⁸ The selection of the π -spacer is critical when designing D- π -A dyes because it

plays a crucial function in regulating the HOMO-LUMO energy

levels and expanding the absorption range of organic sensi-

tizers.69 Ying et al. synthesized and elaborated a series of

materials with the D-A- π -A system as sensitizers.⁵⁸ Among

 $\eta = FF \frac{V_{\rm oc} J_{\rm sc}}{P_{\rm inc}} \tag{1}$

The parameter V_{oc} can be evaluated by the analytical relation according to eqn (2):⁷⁴

$$V_{\rm oc} = E_{\rm LUMO} - E_{\rm CB} \tag{2}$$

where $E_{\rm LUMO}$ indicates the dye's energy level, and $E_{\rm CB}$ designates the energy of the CB of TiO₂ ($E_{\rm CB}=-4.0$ eV).⁷⁵ In a DSSC, $J_{\rm sc}$ can also be determined by the following formula:⁷⁶

$$J_{\rm sc} = \int LHE(\lambda) \ \phi_{\rm inject} \eta_{\rm collect} d\lambda \tag{3}$$

where LHE(λ) signifies the light-harvesting efficiency at a specific wavelength, $\Phi_{\rm inject}$ represents the electron injection efficiency, and η indicates the charge-collection efficiency.

It is vital to note that the LHE of DSSCs is a significant element that affects their performance. In general, a large LHE generates the most significant amount of photocurrent. To compute the LHE, the following equation is used:⁷⁷

$$LHE = 1 - 10^{-f} (4)$$

where f denotes the strength of the oscillator proportional to the $\lambda_{\rm max}$ of the dye. Indeed, this strength has a direct impact on the LHE. Eqn (5) ref. 78 may be used to compute the injection driving force into the CB of the ${\rm TiO_2}$.

$$\Delta G^{\text{inject}} = E^{\text{dye}*} - E_{\text{CB}} \tag{5}$$

where E^{dye^*} signifies the oxidation potential energy of the molecule in the excited state and E_{CB} symbolizes the reduction potential of the conduction band of TiO₂. The value of E^{dye^*} may be computed in the following manner:⁷⁹

$$E^{\text{dye}^*} = E^{\text{dye}} - E^{00} \tag{6}$$

where $E^{\rm dye}$ defines the oxidation potential of the molecule in the neutral state ($E^{\rm dye}=-E_{\rm HOMO}$), and $E^{\rm 00}$ denotes the vertical transition energy corresponding to the $\lambda_{\rm max}$.⁸⁰

The driving force for dye regeneration denoted by the symbol $\Delta G_{\rm reg}$ can be expressed as follows:⁸¹

$$\Delta G_{\text{reg}} = E^{\text{dye}} - E_{\text{Redox}}^{\text{Electrolyt}} \tag{7}$$

where $E_{\rm Redox}^{\rm Electrolyt}$ is represented as the redox potential of the electrolyte I^/I_3^ (-4.80 eV).⁷⁵

The DFT approach can be used to optimize dyes' positive and negative states to calculate their total energy of reorganization. ⁸² It is known that a significant value of $J_{\rm sc}$ requires a low total energy of reorganization; this can be confirmed by combining the hole energy of reorganization with the electron one, ⁸³ as represented by the following formula (8):⁸⁴

$$\lambda_i = (E_0^{\pm} - E_+^{\pm}) + (E_+^0 - E_0) \tag{8}$$

wherhe energy of the catie E_0^{\pm} represents ton (anion) calculated with the optimized structure of the neutral molecule, E_{\pm}^{\pm} is the energy of the cation (anion) calculated with the optimized cation (anion) structure, E_{\pm}^0 denotes the energy of the neutral molecule calculated at the cationic (anionic) state, and E_0 is the energy of the neutral molecule at the ground state.

In addition, excellent nonlinear optical (NLO) characteristics may affect the short-circuit current density. ⁸⁵ According to eqn (9) and (10), the polarizability (α) and hyperpolarizability (β_{tot}) are determined as:

$$\alpha = \frac{1}{3} \left(\alpha_{xx} + \alpha_{yy} + \alpha_{zz} \right) \tag{9}$$

$$\beta_{\text{tot}} = \sqrt{\beta_x^2 + \beta_y^2 + \beta_z^2} \tag{10}$$

where xx, yy, and zz represent the tensor coefficients of polarizability, and $\beta_i = (i = x, y, z)$ combines the different quantities:

$$eta_i = \left(rac{1}{3}
ight) \sum_{j=x,y,z} \left(eta_{ijj} + eta_{jij} + eta_{jji}
ight)$$

Indeed, different reactivity parameters have been studied to explain and predict various physicochemical phenomena that provide details about the electrical structure of the chemistry of target compounds, ⁸⁶ *e.g.* employing E_{HOMO} and E_{LUMO} can give the chemical potential (μ) , ⁸⁷ chemical hardness (η) , ^{88,89} softness (S), and electronegativity (χ) by using the following formulas:

Chemical potential
$$\mu = \frac{E_{\text{LUMO}} + E_{\text{HOMO}}}{2}$$
 (11)

Electronegativity
$$\chi = -\mu = \frac{-(E_{\text{LUMO}} + E_{\text{HOMO}})}{2}$$
 (12)

Chemical hardness
$$\eta = \frac{E_{\text{LUMO}} - E_{\text{HOMO}}}{2}$$
 (13)

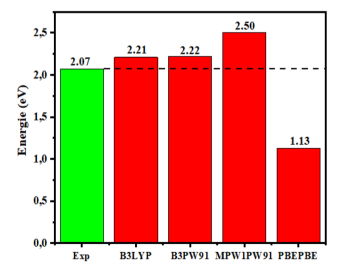


Fig. 2 $E_{\rm gap}$ of D1 determined using DFT with different functionals and comparison with the experimental values.

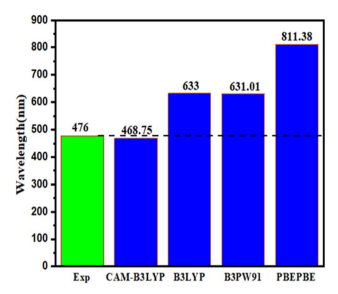


Fig. 3 Examining the functional used in the TD-DFT calculation of the maximum absorption wavelengths (λ_{max}) for D1.

All the computations in this research were performed using the Gaussian 09 software package, ⁹¹ and GaussView 6.0.16 for the simulated spectra and orbital density.

Selecting the test method

To examine the impact of different functionals on the electronic characteristics: the E_g of the dye D1 was studied theoretically

using the PBEPBE,⁹² MPW1PW91,⁹³ B3PW91,⁹⁴ and B3LYP⁹⁵ functionals combined with the 6-31G** basis set, and the data obtained are presented in Fig. 2, which shows the bandgap energies determined from B3LYP/6-31G**. The results were close to the experimental results with a reliable and acceptable error variance, while the error margin given by other functionals was higher. Therefore, the B3LYP/6-31G** functional was considered to be much more suitable for determining our developed molecules' structural and electronic characteristics.

The choice of the TD-DFT theoretical function to describe the vertical transition energy of a potential sensitizer depends on the reference molecule's absorption wavelengths (λ_{max}). Fig. 3 shows the absorption wavelengths (λ_{max}) of the reference molecule examined using TD-DFT with various functionals, such as PBEPBE, ⁹² CAM-B3LYP, ⁹⁶ B3LYP, ⁹⁵ and B3PW91 (ref. 97) combined with the 6-31G** basis set. The results indicated that the CAM-B3LYP with 6-31G** offered a maximum value of λ equal to 468.75 nm, similar to the experimental value (476 nm). ⁵⁸ Based on the findings, CAM-B3LYP with the 6-31G** level was chosen to compute the optical characteristics of the five proposed molecules (D1, D2, D3, D4, and D5) (Fig. 4).

Results and discussion

Molecular geometry

Table 1 gives the geometric characteristics of the potential sensitizer structures, where d explains the bond distance and θ represents the dihedral angle (Scheme 1). As illustrated in Table

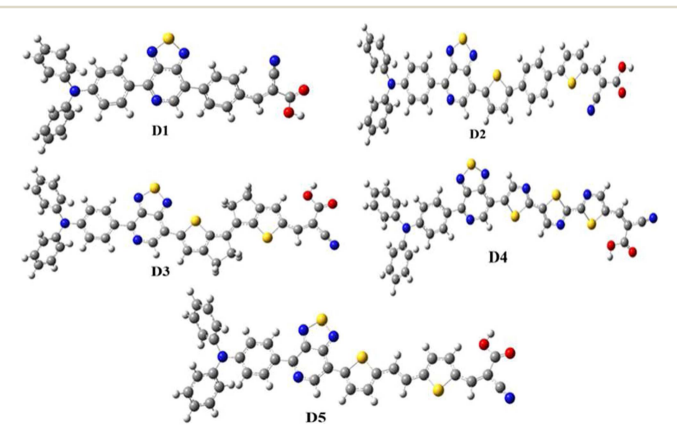


Fig. 4 Structural optimization of the studied molecules (D1-D5).

Table 1 Distances of d_i (Å) and the dihedral angles θ_i (°) of the studied dyes

Dye	$d_1(\mathring{\mathrm{A}})$	$d_2(\mathring{ m A})$	d_3 (Å)	θ_1 (°)	θ_2 (°)	θ_3 (°)
D1 D2 D3	1.46 1.46 1.46	1.46 1.45 1.45	1.45 1.43 1.43	-176.28 -175.45 -176.69	-150.61 -170.76 -175.60	-179.06 -179.09 -179.58
D4 D5	1.46 1.46	1.45 1.45	1.43 1.43	-178.05 -177.28	-179.08 -179.64	-179.99 -179.36

Scheme 1 Studied geometric parameters.

1, all the parameters were determined at the DFT/B3LYP/6-31G** level. It is important to note that the short bond lengths facilitate the ICT in the D- π -A architecture. ⁹⁸ The optimized structures of all the studied dyes as sensitizers showed identical

conformations, as confirmed by the bond length values all lying in the range of 1.43–1.46 Å. These bond lengths revealed a C=C character in all the dyes, which facilitated ICT between the donor fragment and the -spacer and the -bridge and the acceptor fragment. According to Table 1, the values of θ_1, θ_2 , and θ_3 for all the potential proposed dyes ranged between -175° and -179° , which revealed that all the compounds possessed a planar geometry with the dihedral angle θ_2 of the D1 dye. This was slightly twisted due to the steric interactions between the π -spacer and acceptor fragments. 99 Thus, altering the π -bridge influenced the planarity of these designed molecules, hence increasing their light-collecting effectiveness.

Frontier molecular orbitals

Frontier molecular orbitals (HOMO and LUMO) are quantum mechanical descriptors that give essential information about the optoelectronic properties of dyes. These orbitals assist in examining the distribution and stabilization of the charge density throughout the dyes. According to the literature, electron transfer ($S_0 \rightarrow S_1$) must occur in each noticed dye from the ground state (HOMO) to the excited state (LUMO). HOMO-LUMO band gap (E_g) indicates the decrease and increase in reactivity due to various π -spacers. Charge transfer is facilitated when the band gap is small and *vice versa*. Theoretical determination of the energy levels for the HOMO and LUMO for any dye is vital for understanding the

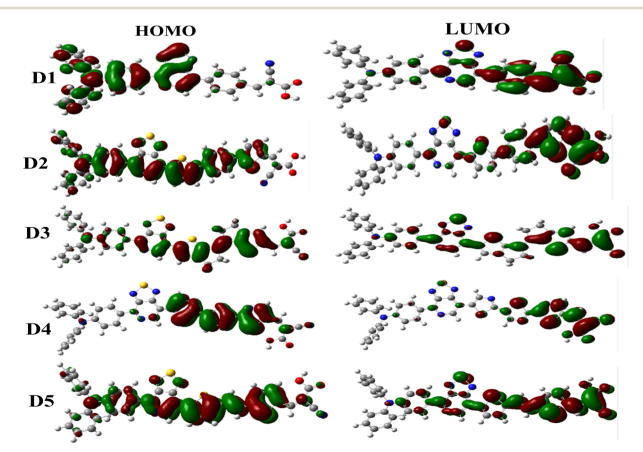


Fig. 5 Contour plots of the FMOs of all dyes examined.

construction of the electronic features in dye systems.¹⁰⁴ This helps to determine the suitability of suggested dyes to fit the fundamental conditions for DSSC photosensitizer materials.¹⁰⁵ The E_{HOMO} , E_{LUMO} , and the various E_{gap} values of some currently investigated research molecules are illustrated in Fig. 6, and the FMO schematic is shown in Fig. 5.

In this work, the proposed dyes (D1–D5) comprised triphenylamine as electron-donor unit [1,2,5]-thiadiazolo[3,4-c]-pyridine as the auxiliary acceptor group, different π -linkers, and acceptor parts (cyanoacrylic acid). The analysis of the frontier orbitals demonstrated that the HOMO of dye D1 was concentrated on the donor triphenylamine and [1,2,5]-thiadiazolo[3,4-c]-pyridine. At the same time, the LUMO was distributed on the π -spacers and the acceptor regions (cyanoacrylic acid). For the dyes D2, D3, and D5, the HOMO electrons were widely distributed on these dyes with a maximum density on different π -spacers. In contrast, the LUMOs were principally situated on cyanoacrylic acid, with a minimum density on the π -spacers groups. Finally, the D4 dye showed a high HOMO on the π -spacers group and a strong LUMO on the cyanoacrylic acid group.

Based on DSSC devices' operating principle, photosensitizer materials should have appropriate energy levels to allow for electron injection and material regeneration. The energy levels of the D1 to D5 molecules were estimated at the B3LYP/6-31G** level, and their energy diagrams are shown in Fig. 6, showing that the HOMO levels of all the molecules were below the redox potential of I⁻/I₃⁻. Therefore, the oxidized molecules (photosensitizers) could be reduced by electrons from the electrolyte (I⁻/I₃⁻). Moreover, the molecules had LUMO values higher than the CB of TiO₂ (-4.00 eV), which implied that the electrons could be injected quickly into the TiO2 semiconductor during the excitation of the molecules. As previously stated, the proposed dyes (D1-D5) are good photosensitizer selections and are suitable for DSSC applications. The HOMO energy values of the proposed dyes (D1-D5) followed the order: D3 (-4.979 eV) > D2 (-5.045 eV) > D5 (-5.118 eV) > D1 (-5.221 eV) > D4 (-5.255 eV), and the values implied a regeneration capacity and good

performance. Hence, the proposed molecules D2, D3, and D5 had greater regeneration capacities than D1. Meanwhile, the LUMO levels of the molecules D2–D5 and D1 were -2.942, -3.086, -3.316, -3.159, and -3.011 eV, respectively. The results showed no significant difference except for D2.

The energy gap has a considerable effect on the activity of molecules; whereby a sensitizer with a small bandgap energy is more favourable for electron excitation, and has enhanced absorption properties. For that reason, sensitizers may absorb more photons in the visible region, increasing the $J_{\rm sc}$ and PCE.^{106,107} The findings in Fig. 6 show that the bandgap energy of the proposed molecules could be modulated by inserting various aromatic spacers. D3 offered the lowest $E_{\rm g}$ of 1.89 eV, followed by D4 (1.93 eV), D5 (1.95 eV), D2 (2.10 eV), and D1 (2.21 eV). The proposed molecules (D2–D5) had smaller $E_{\rm g}$ values than D1, implying that D2–D5 would have a better short-circuit current density ($J_{\rm sc}$) than D1. Due to a smaller energy gap, the UV-vis spectra of D2–D5 may be red-shifted relative to D1.

Density of states (DOS)

Numerous studies have shown that examination of the density of states can explain the charge-transfer characteristics of organic semiconductors.108 Indeed, it also provides the contribution of the MOs of each molecule's different components to the overall system. The GaussSum can be used to determine the density of states (DOS) function. 109 Fig. 7 shows the DOS results for predicting the alpha and beta free electrons for the valence and conduction bands, respectively.110 The energy of the band gap increased in parallel with the DOS of the proposed molecules. In DSSCs, the voltage is associated with the energy gap between the Fermi level (next to the conduction-band level) of the TiO₂ semiconductor electrode and the redox potential in the electrolyte. Evidently, when $E_{\rm F}$ (Fermi energy) is near the midgap, the charge is transported from/to the metal electrode and organic material. Hence, the potential energy profile stays flat whenever $E_{\rm F}$ approaches either the HOMO or LUMO level, while charge transfer is significant.33

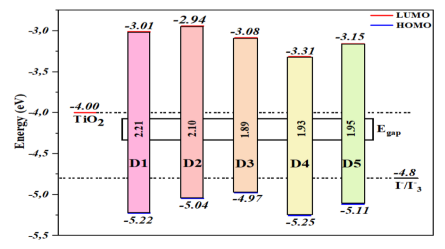


Fig. 6 Illustration of the energy levels of all molecules studied, TiO_2 , and the electrolyte (I^-/I_3^-).

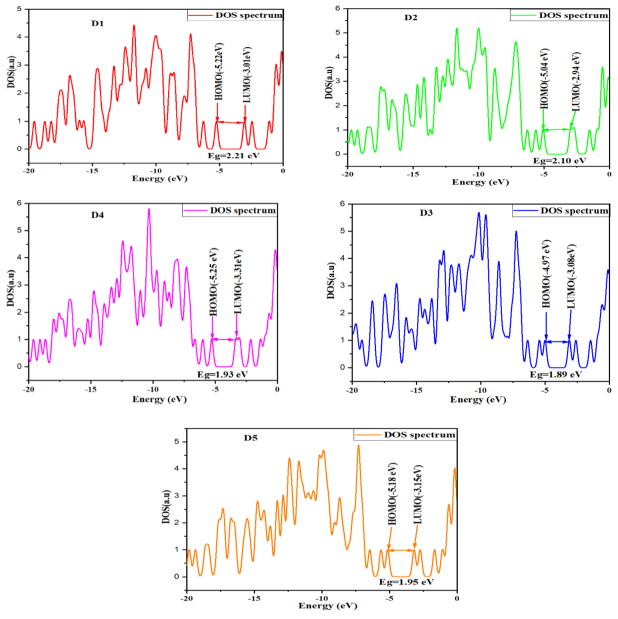


Fig. 7 Density of states for the examined molecules D1–D5.

Quantum chemical reactivity indices

The chemical potential (μ), electronegativity (χ), and chemical hardness (η) of the proposed molecules were obtained from eqn

 $\begin{tabular}{ll} \textbf{Table 2} & \textbf{Calculated values for the electronic and chemical parameters} \\ \textbf{of the five dyes studied} \\ \end{tabular}$

Dye	E_{HOMO} (eV)	E_{LUMO} (eV)	$E_{\rm gap}$ (eV)	μ (eV)	η (eV)	χ (eV)
D1	-5.223	-3.012	2.21	-4.12	1.10	4.12
D2	-5.045	-2.942	2.10	-3.99	1.05	3.99
D3	-4.979	-3.086	1.89	-4.06	0.94	4.06
D4	-5.255	-3.316	1.93	-4.28	0.97	4.28
D5	-5.118	-3.159	1.95	-4.14	0.98	4.14
TiO_2	_	-4.00	_	-4.66	3.78	4.66

(11)–(13) using the B3LYP method with the 631G** basis set, and the findings are tabulated in Table 2. According to the data in Table 2, the TiO₂ compound had the lowest chemical potential (–4.66 eV)¹¹¹ compared with the potential chemical values calculated for the proposed molecules D1–D5. This suggests the electron transfer from the dyes as electron donors to the TiO₂ semiconductors as electron acceptors would be relatively straightforward.¹¹² Furthermore, the chemical hardness (η) of all the studied molecules was lower than that of TiO₂ ($\eta=3.78$ eV), indicating that all the dyes could release one or more electrons when compared to the TiO₂ semiconductor. Additionally, the electronegativity of TiO₂ (–4.66 eV) was higher than for the dyes, implying that TiO₂ may readily accept electrons from these dyes.

Table 3 NBO analysis for all the proposed molecules (D1-D5)

Dyes	D	A	π	A	$\Delta q(ext{D-A})$
D1	0.124	-0.107	0.085	-0.102	0.226
D2	0.111	-0.133	0.167	-0.145	0.256
D3	0.117	-0.141	0.222	-0.198	0.315
D4	0.136	-0.128	0.102	-0.110	0.246
D5	0.134	-0.139	0.178	-0.173	0.307

Natural bond orbital (NBO) analysis

NBO analysis is a key factor for studying the charge transfer or conjugative interactions of molecular systems. It offers a good platform for examining intramolecular and intermolecular bonds and bond interactions. Therefore, NBO analysis was carried out here at the DFT/B3LYP/6-31G** level to determine charge populations for the proposed molecules, and the data are reported in Table 3. The donor and π -linker groups of all the molecules exhibited a positive NBO charge, indicating that they were efficient electron-donating groups. On the other hand, the

negative NBO value of the supplementary acceptor and main acceptor groups suggested they were good electron-withdrawing groups. During photoexcitation, the NBO charges on the donor groups of each molecule were arranged as follows: D4 (0.136e) > D5 (00.134e) > D1 (0.124e) > D3 (0.117e) > D2 (0.111e), suggesting that D4 and D5 had a better electron-donating capacity than the reference molecule D1. For the acceptor, the NBO charges for D1–D5 were $-0.102,\,-0.145,\,-0.198,\,-0.110,$ and -0.173e, respectively, indicating that the insertion of thiophene derivatives as π -spacers could augment the electron density in the acceptor. In addition, $\Delta q(\text{D-A})$ indicates the difference between the electron donor and acceptor. According to Table 3, D3 had a maximum q(D-A) (0.315e), followed by D5 (0.307e), D2 (0.256e), D4 (0.246e), and D1 (0.226e), signifying that all the dyes had a greater charge separation than D1.

Electrostatic potential (ESP)

The electron density is an essential parameter for predicting both the reactivity of electrophilic and nucleophilic sites (*i.e.* an

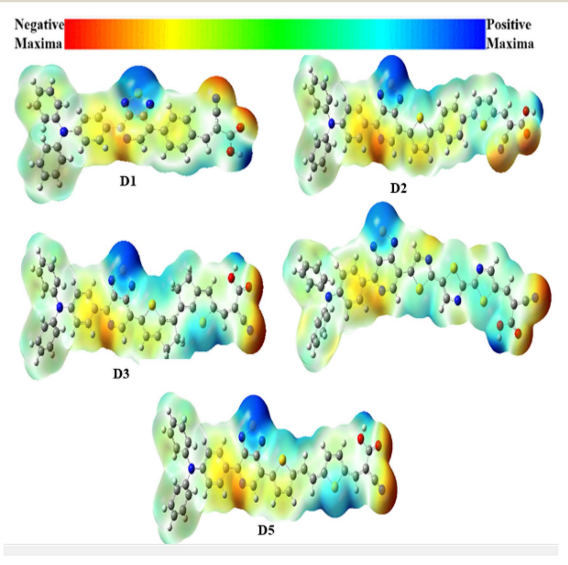


Fig. 8 Calculated electrostatic potentials on the molecular surfaces of the studied dyes.

Table 4 Calculated average isotropic polarizability $\langle \alpha \rangle$ and the first hyperpolarizability (β_{tot}) for D1–D5 and their constituents applying the B3LYP/6-31G** set

Parameter	D1	D2	D3	D4	D5
α_{xx}	1127.90	1826.55	2200.52	1883.56	2119.14
α_{xy}	24.07	6.94	24.00	-11.03	-6.22
α_{yy}	437.46	521.44	550.56	555.79	487.52
α_{xz}	-0.16	10.55	-4.12	-11.68	0.92
α_{yz}	-12.34	-30.75	-9.01	2.33	-16.86
α_{zz}	204.27	284.85	260.63	250.67	235.99
$\langle \alpha \rangle$ (a.u.)	589.88	877.61	1003.90	896.67	947.55
$\langle \alpha \rangle (10^{-24} \text{ esu})$	87.42	130.06	148.77	132.88	140.42
β_{xxx}	83 624.40	-127466.00	-108879.00	-177392.00	-155029.00
β_{xxy}	4543.67	-9817.01	-9986.1	15 466.20	-10344.30
β_{xyy}	-113.06	1108.60	1254.72	-5037.69	994.63
β_{yyy}	-63.76	-69.38	-170.44	810.49	-142.33
β_{xxz}	1854.24	1092.00	2854.64	1835.06	2672.25
β_{xyz}	417.07	407.57	127.05	-1096.19	153.20
β_{yyz}	35.34	22.73	8.679	268.18	35.59
β_{xzz}	48.42	-275.22	137.68	25.50	149.13
eta_{yzz}	22.90	-98.71	-10.98	18.76	-19.46
$\hat{eta_{zzz}}$	17.75	16.73	6.76	-5.30	10.85
β_{tot} (a.u.)	83 702.73	127 030.72	108 004.56	183 142.65	154 267.40
$\beta_{\text{tot}} \left(10^{-30} \text{ esu}\right)$	723.133	1097.457	933.084	1582.225	1332.624

explanation of chemical reactions) and the interactions through hydrogen bonding.¹¹⁴ The molecular electrostatic potentials (MEPs) for the D1–D5 dyes are shown in Fig. 8, where the red area corresponds to the negative potentials (the electron-rich field), which characterizes the reactive zone for an electrophilic attack. In contrast, the blue area relates to the positive potentials (electron-poor fields), which determine the reactive zone for nucleophilic attack. Thus, a specific colour defined each MEP value in the following order: red < orange < yellow < blue. As illustrated in Fig. 8, the MEP revealed that all the dyes'

maximum positive area (blue colour) was centred on the carboxyl hydrogen atom and the sulfur atoms of the auxiliary acceptor, mainly for D1, D4, and D5. In contrast, the negative area (red colour) was primarily centred around the nitrogen atoms of the acceptor groups (–CN) in the D1–D5 dyes, which were the most favourable sites for the electrolyte.

Nonlinear optical properties (NLO)

NLO characteristics were measured *via* the intra-molecular charge delocalization in conjugated molecular systems. Hence,

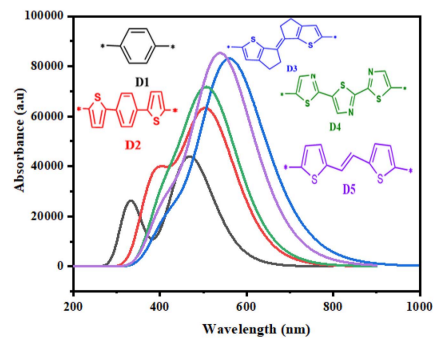


Fig. 9 Absorption spectra of the studied dyes.

Table 5 Absorption spectral data calculated at the TD-DFT/CAM-B3LYP/6-31G** level

Dyes	λ_{max} (nm)	$E_{\mathrm{ex}}\left(\mathrm{eV}\right)$	f	MO/character
D1	468.749	2.64	1.08	HOMO → LUMO (83%)
	338.338	3.66	0.20	HOMO-1 → $LUMO$ (77%
	329.920	3.75	0.44	$HOMO \rightarrow LUMO+1 (53\%)$
D2	508.840	2.43	1.52	HOMO → LUMO (81%)
	395.307	3.13	0.78	HOMO-1 → $LUMO$ (31%)
	368.605	3.36	0.16	$HOMO-1 \rightarrow LUMO (49\%)$
D3	563.103	2.20	2.01	$HOMO \rightarrow LUMO (89\%)$
	438.804	2.82	0.37	$HOMO \rightarrow LUMO+1 (65\%)$
	402.245	3.08	0.17	$HOMO-1 \rightarrow LUMO (65\%)$
D4	514.051	2.41	1.67	$HOMO \rightarrow LUMO (49\%)$
	413.363	2.99	0.69	HOMO-1 → $LUMO$ (38%)
	367.851	3.37	0.01	$HOMO-1 \rightarrow LUMO$ (27%)
D5	542.007	2.28	2.06	HOMO → LUMO (83%)
	421.542	2.94	0.55	$HOMO \rightarrow LUMO+1 (49\%)$
	384.554	3.22	0.10	$HOMO-1 \rightarrow LUMO (52\%)$

these properties might be utilized to assess the efficiency of the hole-electron transfer of the sensitizer, which can also affect the $V_{\rm oc}$ and the solar cell performance. It has been shown that conjugated molecular systems possess excellent NLO properties. Therefore, it often offers excellent photoelectric conversion characteristics. The α and β_{tot} of the molecules are computed employing the B3LYP approach and 6-31G** basis set, and the calculation results are listed in Table 4. Based on the values in Table 4, the polarizability (α) of all dyes is arranged as follows: D3 (148.77 \times 10⁻²⁴ esu) > D5 (140.42 \times 10⁻²⁴ esu) > D4 (132.88 \times 10⁻²⁴ esu) > D2 (130.06 \times 10⁻²⁴ esu) > D1 (87.42 \times 10⁻²⁴ esu). These results indicate that D2-D3 performs better in electron charge transfer than D1. For the hyperpolarizability (β_{tot}) , D2–D5 have greater first hyperpolarizability than D1, and values are in the following order: D4 > D5 > D2 > D3 > D1. It indicates that the proposed D2-D5 dyes will present a more efficient photocurrent response and perform better than D1 due to their excellent NLO characteristics.

Optical properties

Photovoltaic characteristics are strongly related to the absorption of photons by dyes. An excellent organic solar cell must possess a broad and strong visible absorption spectrum.104 To examine the effects of various π -spacers of the proposed dyes (D2-D5) based on the molecule (D1) and to get a clearer idea of the possible electronic transitions of the five studied dyes, the proposed dyes' electronic transitions and absorption characteristics were calculated using TD-DFT/CAM-B3LYP/6-31G**, 118 in the gas phase. The absorption spectra of the reference molecule (D1) and the investigated molecules (D2-D5) are shown in Fig. 9, and the results are tabulated in Table 5. From the results in Fig. 9 and Table 5, the maximum absorption values obtained for the molecule (D1) and its derivatives D2, D3, D4, and D5 were 468.749, 508.840, 563.103, 514.051, and 542.007 nm, respectively. The proposed dyes showed high absorption between 400 and 800 nm in the visible zone. These values correlated with the lowest energy and were ascribed to an

Table 6 Calculated electronic properties of the studied dyes (in eV)

Dyes	$E_{ m HOMO}$	$E_{ m LUMO}$	E^{00}	E^{dyes}	E ^{dyes*}	$\Delta G^{ m inject}$	ΔG_{reg}	LHE	$V_{ m oc}$
D1	-5.223	-3.012	2.64	5.22	2.58	-1.42	0.42	0.91	0.99
D2	-5.045	-2.942	2.43	5.04	2.61	-1.39	0.24	0.96	1.06
D3	-4.979	-3.086	2.20	4.97	2.78	-1.22	0.17	0.99	0.92
D4	-5.255	-3.316	2.41	5.25	2.84	-1.16	0.45	0.98	0.69
D5	-5.118	-3.159	2.28	5.11	2.83	-1.17	0.31	0.99	0.85

electronic transition from the ground state S_0 to the excited state S_1 . In addition, the maximum absorption wavelengths of these dyes were 542.007 nm and 563.103 nm for D5 and D3, respectively. Indeed, a dominant transition with low energy (2.20-2.64 eV) and a higher value of oscillation force (1.08-2.06) accompanied the absorption spectra.

Major parameters affecting the photovoltaic performance of DSSCs

The influence of multiple π -linkers on the effectiveness of DSSCs was determined by investigating some essential parameters, such as $\Delta G^{\rm inject}$, $\Delta G^{\rm reg}$, LHE, and the excited-state lifetime (τ) , which were used to examine and assess the $J_{\rm sc}$ and $V_{\rm oc}$. $\Delta G^{\rm inject}$ and LHE are essential parameters influencing the $J_{\rm sc}$ obtained from eqn (6). Suitable dyes must have a significant LHE that reflects the absorption ability of most photons in the UV-visible region and subsequently injects photoexcited electrons into the CB of TiO₂. Table 6 summarizes the LHE, $\Delta G^{\rm inject}$, $\Delta G_{\rm reg}$, and $V_{\rm oc}$ values, where the LHE value for D3 and D5 exhibited the largest value (0.99), followed by D2 (0.96), D4 (0.98), and D1 (0.9), suggesting that the developed dyes D2–D5 showed higher light-absorption potentials compared to D1. Hence, D2–D5 will present higher $J_{\rm sc}$ values because of their superior LHE.

The negative $\Delta G^{\rm inject}$ values demonstrated that electrons could be injected easily into TiO₂. The $\Delta G^{\rm inject}$ values in Table 6 show that D1 and D2 had greater values than D3, D4, and D5, indicating that electron injection would be easier into TiO₂ for them. To get fast electron transfer, $\Delta G_{\rm reg}$ must be as low as possible; as shown in Table 6, except for D4, the $\Delta G_{\rm reg}$ values of all the developed compounds were lower than that of the reference molecule D1, indicating that those dyes had a greater power conversion efficiency.

The charge-transfer rate (k) also influences a DSSC device's efficiency, which may be defined as follows:²⁶

Table 7 Reorganization energies, λ_{total} values (eV), and excited-state lifetime τ (ns) of the molecules (D1–D5)

$\lambda_{e}\;(eV)$	$\lambda_h \; (eV)$	$\lambda_{\rm total} \ (eV)$	τ (ns)
0.1	0.20	0.30	0.19
0.09	0.17	0.26	0.16
0.20	0.32	0.52	0.15
0.06	0.13	0.19	0.15
0.08	0.15	0.23	0.14
	0.1 0.09 0.20 0.06	0.1 0.20 0.09 0.17 0.20 0.32 0.06 0.13	0.1 0.20 0.30 0.09 0.17 0.26 0.20 0.32 0.52 0.06 0.13 0.19

$$k = \left(\frac{\pi}{\lambda_{\text{total}} k_b T}\right)^{\frac{1}{2}} \frac{V^2}{\hbar} \exp\left(-\frac{\lambda_{\text{total}}}{4k_b T}\right) \tag{14}$$

where V is the charge-transfer coupling constant, T is the temperature, $k_{\rm b}$ is the Boltzmann constant, and the $\lambda_{\rm total}$ is the reorganization energy. Except for the $\lambda_{\rm total}$, all the parameters in eqn (14) are believed to be constants; while a lower $\lambda_{\rm total}$ will expedite electron transport and boost the $J_{\rm sc}$. $\lambda_{\rm total}$ is the sum of the electron-reorganization energies ($\lambda_{\rm e}$) and the hole-reorganization energies ($\lambda_{\rm h}$); these energies were determined using eqn (8), and obtained values are listed in Table 7, where the computed $\lambda_{\rm total}$ for the proposed dyes (D1–D5) increased in the following order: D4 < D5 < D2 < D1 < D3. So, due to the low values of $\lambda_{\rm total}$, the molecules D2, D4, and D5 would have good electron-transfer efficiency compared to molecule D1 (the reference molecule).

Also, (τ) has an essential effect on the charge-transport characteristics of a material. Indeed, a longer lifetime of the excited state enables charge transport and reduces the loss of energy¹¹⁹ eqn (15) was used to determine the (τ) of the dyes:¹²⁰

$$\tau = \frac{1.499}{f \times E^2} \tag{15}$$

where *E* denotes the transition energy between various excited states, and *f* indicates the oscillator strength. Table 7 shows the data for (τ) of all the molecules. From Table 7, the values of the excited-state lifetimes were in the following order: D1 (0.19 ns) > D2 (0.16 ns) > D3 (0.15 ns) = 4 (0.15 ns) > D5 (0.14 ns).

Conclusion

In this research, four novel molecules (D2–D5) with different π -bridges were developed based on previous research-based experiment results of the dye D1. DFT and TD-DFT were applied as theoretical methods to test the critical parameters associated with the $J_{\rm sc}$ and $V_{\rm oc}$ values of the dyes. The results demonstrated that all the developed compounds (D2–D5) possessed lower energy gaps, thus resulting in their absorption spectra being red-shifted by a different degree than D1, particularly for D3 and D5. It is important to note that D2–D5 presented larger absorption bands, indicating a higher light-absorption possibility. The more outstanding $V_{\rm oc}$ and $J_{\rm sc}$ of the proposed dyes D2–D5 may be partly attributable to their lower $\lambda_{\rm total}$, $G_{\rm reg}$, η , $G^{\rm inject}$, and larger LHEs.

On the other hand, D2–D5 exhibited higher charge-transfer efficiencies and a reduced charge recombination protocol compared to D1. According to this investigation, the modifications of the π -spacer units of the dye reference affected the overall performance of the studied dyes. This study should give theoretical guidance for experiments to develop and test highly effective D–A– π –A type sensitizer materials.

Author contributions

A. Azaid: paper writing; T. Abram, R. Kacimi, M. Raftani, Y. Khaddam, and D. Nebbach: graph preparation, analyzed and interpreted the data; M. Bouachrine and M. Alaqarbeh:

analyzed the DFT calculations and supervision of the work; A. Sbai and T. Lakhlifi: computational part of the study and DFT calculations. All authors commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

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