Journal of Materials Chemistry C



View Article Online

View Journal | View Issue

PAPER

Check for updates

Cite this: J. Mater. Chem. C, 2024, **12**, 903

Received 13th October 2023, Accepted 7th December 2023

DOI: 10.1039/d3tc03740a

rsc.li/materials-c

1. Introduction

Solar energy stands out as one of the most accessible and costeffective renewable energy sources. Consequently, a wide range of photovoltaic technologies have emerged to efficiently convert sunlight into electrical current. While the majority of photovoltaic cells in current use rely on silicon,¹ efficient alternatives,

Charge transfer properties of novel linear carbon chain-based dyes[†]

Giuseppe Consiglio, 🝺^a Adam Gorcyński, 🝺^b Salvatore Petralia 🝺^c and Giuseppe Forte 🕩 *^c

In this study, we designed a series of dyes based on the D- π -A scheme, featuring the novel 2-amino pyrrole group, D1, as the donor unit, combined with three different acceptor groups: benzothiadiazole (A2), cyanoacrilic acid, (A3), and a unique pyrimidin-pyridinium salt derivative, A1, not previously employed in DSSC applications. A linear carbon chain (LCC) of varying lengths was used as the π -bridge. We compared the performance of D1 dyes with the well-established electron donor phenothiazine, D2. Spectroscopic, electronic and photochemical properties of these designed compounds were investigated through density functional theory (DFT) and time-dependent density functional theory (TDDFT) simulations. The calculated light-harvesting efficiency (LHE) values showed that A1 yielded the best results, and generally, D1 donors outperformed the reference D2. Additionally, longer LCCs contributed positively to LHE. The study of electron density difference (EDD) maps showed clear charge transfer from the donor unit to the acceptor unit in all cases. Dyes with D1 exhibited lower intramolecular reorganization energy (λ_i) values, indicating faster charge transfer compared to those with D2. Negative free energy injection values ($\Delta G^{(n)}$) were calculated for all sensitizers, signifying efficient electron injection, especially in D2containing dyes. However, these compounds exhibited positive free energy regeneration values (ΔG^{reg}), implying thermodynamically unfavorable regeneration processes. Conversely, D1-containing compounds demonstrated favorable values for regeneration. Increasing the length of the LCC positively affected ΔG^{reg} but had a less favorable impact on ΔG^{inj} . The calculated charge transfer (q_{CT}) values revealed that the most favorable results occurred when pairing the donor with A1, indicating that this acceptor group promoted charge separation. Simulations of dye@(TiO2)14 anatase complexes indicated that the binding of the designed dyes to the semiconductor improved their transport properties. In summary, all the calculated parameters indicated that D1 possesses superior properties for use in DSSC applications when coupled to A1 and, generally, dyes featuring D1 were predicted to perform well in photovoltaic applications. It is worth noting that ΔG^{reg} values may hinder regeneration in D2-containing dyes, despite their excellent values in other calculated parameters. Our study offers valuable insights for refining dye design in order to boost the performance of dye-sensitized solar cells (DSSCs).

such as perovskite solar cells and dye-sensitized solar cells (DSSCs), have come into prominence.² Since their introduction by Grätzel and O'Regan,³ DSSCs have emerged as a promising technology for efficiently harnessing solar energy to generate electrical power. They have attracted growing interest due to their cost-effective manufacturing, distinctive optical and mechanical properties, and high device efficiency.^{4,5} An essential component of DSSCs is the photosensitizer, which plays a crucial role in absorbing photons and initiating the charge separation process.⁶ Over time, extensive research efforts have been devoted to optimizing dye design with the goal of improving DSSC performance. These investigations have explored a wide range of photosensitizers, primarily grouped into three main classes: ruthenium(II) complexes,^{7–11} zinc(II) porphyrin derivatives^{12–18} and metal-free organic dyes. Among these classes, free-metal

^a Department of Chemical Science, University of Catania, Via S. Sofia 64, 95125, Italy

^b Faculty of Chemistry, Adam Mickiewicz University, Universytetu Poznańskiego 8, 61-614 Poznań, Poland

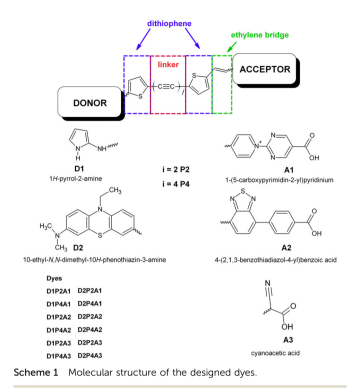
^c Department of Drug Science and Health University of Catania, Via S. Sofia 64, 95125, Italy. E-mail: gforte@unict.it

[†] Electronic supplementary information (ESI) available. See DOI: https://doi.org/ 10.1039/d3tc03740a

Paper

organic dyes have garnered particular attention due to their lowcost fabrication processes and favorable environmental impact. This category typically comprises three distinct regions, each serving a different purpose. First, an electron acceptor moiety (A) anchors the dye to the semiconducting electrode, ensuring an efficient electronic coupling between the sensitizers and the substrate. Second, a linker section (π) , consisting of a conjugated system, facilitates electron transfer. Finally, an electron donor moiety (D), which is located at the free end of the molecule. As a result, these dyes typically adopt a push-pull structure, denoted as D- π -A. Various strategies have been explored to optimize both the electron acceptor and electron donor moieties in order to achieve higher efficiency and improved performance in photovoltaic applications. In order to enhance the efficiency of charge transfer, the careful selection of appropriate donor-acceptor pairs is essential. Within this context, the role of the π -linker holds significant importance. A multitude of innovative metalfree DSSCs have been developed, utilizing a range of compounds including triphenylamine, dialkylaniline, coumarin, oligoene, indoline, perylene, phenothiazine, tetrahydroquinoline, and carbazole.¹⁹⁻²⁸ Among the assortment of novel metal-free DSSCs, dyes based on phenothiazine (PTZ) have demonstrated remarkable performance in DSSC applications. This group of sensitizers exhibiting strong electron-donating properties, superior to triphenylamine, is arranged in a non-planar butterfly conformation in the ground state, thus obstructing the molecular aggregation.²⁹ In particular, some reports indicate that DSSCs based on PTZ dyes display power conversion efficiency (PCE) superior to the commercial N719 sensitizer under the same fabricating and testing conditions.^{30,31} In combination with the cyanoacrilic group as an electron-attracting component, the PTZ-based dye has shown significantly increased photocurrent and PCE when codeposited with deoxycholic acid.³² Xie and coworkers synthesized two innovative organic sensitizers utilizing PTZ and oligothiophene vinylene as conjugation spacers, which yielded a PCE value of 5.4%.33 Additionally, co-sensitizing a PTZ-containing porphyrin sensitizer with a metal-free dye resulted in a co-sensitized device, achieving an efficiency of 11.5%.³⁴ Buene et al. synthesized several PTZ-based dyes to investigate the impact of auxiliary donors, the role of the π -spacer, and anchoring units. They found that the π -spacer is most effective when linked to the 3-position of PTZ, and thiophene outperformed the phenyl spacer.³⁵ In another study, the same group explored various sensitizers, featuring different auxiliary donor groups and determined that pyrene was the most effective, achieving the highest cell efficiency at 5.36%.³⁶ Han et al. introduced auxiliary acceptors (quinoxaline, benzothiadiazole, and BTZ) into PTZ-based dyes to enhance LHE.³⁷ Hua and coworkers developed two novel PTZ-based dyes with the encapsulated insulated molecular wire as an auxiliary donor, effectively reducing dye aggregation and improving photovoltaic cell performance.³⁸ Two new sensitizers with a D/A– π –D– π – A framework were introduced by Slodek and coworkers. The D- π -D- π -A structure exhibited higher PCE, and both dyes achieved higher PCE compared to the N719-based device.³⁹ A comprehensive review of recent articles concerning the use of PTZ derivatives in single, double and multi anchoring configurations for DSSC

applications is offered by Devadiga and coworkers.⁴⁰ The acceptor-anchoring groups are mainly represented by cyanoacrylic moieties which are present in the vast majority of DSSCs with best performance (see all references reported above). Several research groups have explored the employment of a benzothiadiazole group (BTD) with promising outcomes. Yella et al. introduced a novel porphyrin sensitizer employing a push-pull system that incorporated this strong electron-withdrawing group. The resulting device achieved an impressive efficiency of approximately 13% when employing a cobalt-based electrolyte.41 Wang and coworkers linked the BTZ-benzoic acid moiety to a rigidified N-annulated benzoindenopentaphene through an ethynyl group. This led to an exceptional efficiency of 12.6% for the DSSC using this dye, representing one of the highest efficiencies observed for a DSSC sensitized with a single metal-free dye.42 Recently, Ding et al. synthesized two new PTZ-based dyes that incorporated the BTDbenzoic acid unit with a distinguishing ethynyl linkage between BTD and benzoic acid. The results indicated that the insertion of an ethynyl group resulted in a more planar structure, leading to increased dye loading and redshifted intramolecular charge transfer (ICT) absorption with a higher molar extinction coefficient obtaining improved photovoltaic performance.43 To investigate the impact of various acceptors with varying amounts of electron withdrawing groups on free-metal dyes, a series of organic analogues with alternative acceptors were computationally examined. The authors observed that introducing the BTD unit into dyes enhanced π -conjugacy, resulting in a redshift in the UV-VIS spectra and longer excited-state lifetimes. Further investigations demonstrated that the increased presence of BTD groups in the dyes had an adverse effect on properties such as ICT, interface electronic transfer, regeneration driving force and electron-injection efficiency.44 The solvent also plays a critical role in influencing the DSSC properties.45,46 Its primary function is to dissolve the dye and establish contact with the semiconductor surface, thereby initiating the adsorption process. Both aprotic and protic solvents, including acetonitrile, methoxypropionitrile, dimethylformamide, dimethylsulfoxide, ethanol and water, have been thoroughly investigated in numerous experimental and theoretical studies.45-50 Several research studies have shown that aprotic solvents with high polarity, such as acetonitrile, are the preferred choice for DSSCs. This preference stems from their ability to facilitate both dye adsorption and electron injection.^{45,46,50} Our research group recently investigated the role of linear carbon chains (LCCs) in free metal dyes utilizing the donor-acceptor pair formed by the 2-amine pyrrole as the electron-donating group and the cyanoacrylic group as the electron-acceptor. We thoroughly assessed the optoelectronic characteristics of these dyes and evaluated their potential performance in DSSC applications.^{51,52} In this study, we investigate the charge transfer efficiency of 2-amine pyrrole, hereafter referred as D1, in combination with three different electron-acceptor groups, as illustrated in Scheme 1. As discussed above, two of these electron-acceptor groups are widely employed in DSSCs, the BTD derivative, referred to as A2, and the cyanoacrilic group, referred to as A3. The third group, labeled as A1, is based on a pyrimidin-pyridinium salt derivative, and, to the best of our



knowledge, has not been employed in DSSC applications before. The π -linker between the donor and the acceptor groups is represented by a LCC of varying lengths (C4, C8). It is worth noting that LCCs exhibit electrical conduction and possess unusual optoelectronic properties due to their extensive π -conjugated system.53-59 Additionally, they can be easily synthesized and exhibit long-term stability.⁶⁰ In this study, the LCC is flanked by thiophene groups at both ends, as recent literature suggests that good efficiency has been achieved using dithiophene derivatives in the π -bridge.^{61,62} Furthermore, an ethylenic bridge is introduced to link the thiophene ring with the acceptor group, thereby increasing the planarity of the structure. We assess the charge transfer efficiency of the D1 group in comparison to that of the PTZ derivative, hereafter referred to as D2 (see Scheme 1). This choice is based on the demonstrated excellent results of this electron-donating group in DSSC applications. Finally, simulations of dye@(TiO₂)₁₄ anatase nanocluster systems were carried out to elucidate the electronic structure at the interface and to analyze the impact of bonding with semiconductors on electronic transition and injection. The insight gained from this study can serve as a valuable reference for the synthesis and optimization of a novel and more efficient class of metal free dyes for DSSC applications.

2. Methods

Molecular simulations of the investigated dyes were carried out using the Gaussian 16 program package.⁶³ The geometries of all compounds were fully optimized using the B3LYP functional employing the 6-311+G(2d,p) basis set. For the computation of UV/VIS spectra, we employed the TD-DFT approach, along with

the same basis set, considering the lowest 10 singlet-singlet transitions. It is worth highlighting that as demonstrated previously, the long range corrected functional, CAM-B3LYP, has consistently yielded results that align closely with experimental data for compounds with a similar structural framework.^{64,65} However, it should be noted that the molecules under investigation are not currently available for experimental analysis, consequently, there is a lack of experimental data available for comparative assessment. In our earlier investigations, we employed the solvated model based on density (SMD) to account the solvation effects. When comparing this approach with the conductor-like polarizable continuum model (CPCM) and IEFPCM methods, we observed minimal variations in both UV-Vis absorption values and absorption intensities.⁵¹ As a result, we have consistently adopted this methodology. In order to model the semiconductor, we selected a $(TiO_2)_{14}$ cluster, which was carved out from the crystallographic structure of TiO_2 anatase by cleaving the $(1 \ 0 \ 1)$ surface. The dyes were anchored to the TiO₂ surface through bonding between the two oxygen atoms of the cyanoacrylate group and two titanium atoms. This choice is based on previous studies that consistently demonstrate the bidentate mode as the most stable binding geometry.^{66–68} It's worth mentioning that the $(TiO_2)_9$ cluster has been proven to be the smallest acceptable model capable of accurately reproducing the physical parameters of a real TiO₂ semiconductor. As such, the size of the cluster proposed here can be considered a representative model of TiO2.69-71 The optimization of dye@(TiO2)14 geometries was carried out employing the effective core potential LANL2DZ and its accompanying basis set for the Ti atom, whereas the double- ξ basis set 6-31G(d) together with the B3LYP-D3 functional was selected for the other atoms. To consider the solvation effect, we utilized the solvated SMD model. The natural transition orbitals (NTOs) were visualized using Chimera software, and the Multiwfn 3.8 program⁷² was chosen to evaluate the transferred charge, $q_{\rm CT}$, and the charge-transfer distance, $D_{\rm CT}$.

The light-harvesting efficiency (LHE) helps in calculating the dye's efficiency and is determined using the equation:

$$LHE(\lambda) = 1 - 10^{-\varepsilon(\lambda)\Gamma}$$
(1)

where $\varepsilon(\lambda)$ is the molar absorption coefficient whose value depends on wavelength and Γ is the dye loading amount corresponding to the product of the dye concentration *c* with TiO₂ film thickness *b*, ($\Gamma = c \times b$). The short circuit photocurrent, *J*_{sc}, a crucial performance indicator for DSSC devices, is directly proportional to the light harvesting efficiency value and can be determined by the following equation:⁷³

$$J_{\rm SC} = q \int LHE(\lambda) \Phi_{\rm inj} \eta_{\rm coll} \vartheta_{\rm ph.AM1.5G}(\lambda) d\lambda$$
 (2)

where q is the elementary charge of an electron, $\vartheta_{\rm ph.AM1.5G}$ stands the photon flux corresponding to the AM 1.5G solar radiation spectrum. $\Phi_{\rm inj}$ denotes the electron injection efficiency which depends on the injection time and is defined as follows:⁷⁴

$$\Phi_{\rm inj} = \frac{1}{\left(1 + \frac{\tau_{\rm inj}}{\tau_{\rm relax}}\right)} \tag{3}$$

 τ_{inj} is the injection time and τ_{relax} is the relaxation lifetime of the dye's excited states. On the other hand, η_{coll} represents the electron collection efficiency and its value can be calculated as:

$$\eta_{\rm coll} = \frac{1}{\left(1 + \frac{\tau_{\rm trans}}{\tau_{\rm rec}}\right)} \tag{4}$$

where $\tau_{\rm rec}$ denotes the recombination time, whereas $\tau_{\rm trans}$ is the electron-transport time from the CB of the semiconductor to the electrode. Both $\Phi_{\rm inj}$ and $\eta_{\rm coll}$ are related to free energy injection and free energy recombination because of the fact that $\tau_{\rm inj}$ and $\tau_{\rm rec}$ are the inverses of $k_{\rm rec}$ and $k_{\rm inj}$, which, according to Marcus theory,⁷⁵ are defined as:

$$k_{\rm inj/rec} = A \sqrt{\frac{\pi}{\hbar^2 \lambda k_{\rm B} T}} \exp\left(-\beta r\right) \exp\left(\frac{-\left(\Delta G^{\rm inj/rec} + \lambda\right)^2}{4\lambda k_{\rm B} T}\right) \quad (5)$$

where A is a constant, β is the attenuation factor, r is the electron transfer distance and denotes the reorganization energy comprising both intramolecular (λ_i) and intermolecular (λ_o) reorganization energies. The intramolecular reorganization energy quantifies the change in energy associated with the molecule's structural reorganization during the charge transfer process, while the intermolecular reorganization energy characterizes the relaxation of the surrounding medium in which the charge transfer occurs.⁷⁶ Due to its significantly smaller values compared to λ_i , λ_o is usually neglected.⁷⁷ λ_i can be further divided in hole and electron reorganization energy ($\lambda_h + \lambda_e$) which can be calculated using the following formulas:⁷⁶

$$\lambda_{\rm e} = [E^{-}(\mathbf{M}) - E^{-}(\mathbf{M}^{-})] + [E(\mathbf{M}^{-}) - E(\mathbf{M})]$$
(6)

$$\lambda_{\rm h} = [E^+(\mathbf{M}) - E^+(\mathbf{M}^+)] + [E(\mathbf{M}^+) - E(\mathbf{M})]$$
(7)

$$\lambda_{\rm i} = (\lambda_{\rm h} + \lambda_{\rm e}) \tag{8}$$

where $E(\mathbf{M}^+)/E(\mathbf{M}^-)$ represents the neutral energy levels of the optimized cationic/anionic geometries, while $E^+(\mathbf{M})/E^-(\mathbf{M})$ denotes the cationic/anionic energies of the optimized neutral structures.

Finally, $k_{\rm CT}$ is a critical parameter that is related to λ and, in the absence of any barrier, can be calculated as:

$$k_{\rm CT} = \sqrt{\frac{\pi}{\lambda k_{\rm B}T}} \cdot \frac{|V_{12}|^2}{\hbar} \cdot \exp\left(-\frac{\lambda}{4k_{\rm B}T}\right)$$
(9)

where V_{12} represents the coupling constant between the potential curves of the reagent and the product, and it can be determined using the generalized Mulliken–Hush formalism as⁷⁸

$$V_{12} = \frac{|\mu_{12}| \cdot \Delta E_{12}}{\sqrt{(\mu_1 - \mu_2)^2 + 4(\mu_{12})^2}}$$
(10)

where μ_{12} represents the transition dipole moment, ΔE_{12} is the energy difference, μ_1 and μ_{12} are the adiabatic electronic dipoles.

3. Results and discussion

3.1 Frontiers molecular orbital analysis and absorption spectra

An essential prerequisite for the potential utilization of dve sensitizers in DSSCs is the appropriate energy level of the LUMO, enabling efficient electron injection into the TiO₂ conduction band (CB). Moreover, an optimal HOMO energy level plays a significant role in facilitating efficient electron collection. Fig. 1 displays the energy diagram of the selected molecular orbitals for the investigated dyes. This diagram clearly illustrates that the compounds featuring the amino-pyrrolic donor group, D1, have HOMO energy levels positioned below -4.8 eV, which correspond to the redox potential of the electrolyte I^{-}/I_{3}^{-} with respect to the vacuum level,⁷⁹ thus ensuring efficient electron recovery. Conversely, dyes featuring the PTZ group as the electron-donor, D2, show the HOMO energy level slightly above -4.8 eV, suggesting that, in these instances, the recombination process may be hindered. On the other hand, in all considered cases, the LUMO energy levels are significantly higher than the TiO_2 CB, which is at -4.0 eV,⁸⁰ enabling the rapid injection of excited dyes' electrons into the semiconductor.

The absorption properties have been calculated for ten excited states and the dominant electronic transitions are summarized in Table 1, which include information on vertical electronic absorption energies, oscillator strengths, and maximum short circuit photocurrent, Jsc-max. Fig. 2 displays the UV/ VIS absorption spectra of the investigated molecules, revealing a prominent absorption peak in the visible region with considerable intensity. The main contribution to this peak is due to the HOMO \rightarrow LUMO transition and is indicative of intermolecular charge transfer. Additionally, a secondary, albeit less intense, absorption band is observed in the UV region across nearly all cases. Fig. 3 depicts the natural transition orbitals (NTOs) of the lowest energy excited state for the dyes in question. This illustration, which indicates rearrangement after the excitation, demonstrates the favorable inductive and electron-drawing characteristics of both the donor and acceptor groups.

To assess the impact on photovoltaic properties, it is essential to consider both absorption intensity and the optical

D1P2A1 D1P4A1 -1 D1P242 D1P4A2 -2 D1P2A3 D1P4A3 D2P2A1 -3 D2P4A1 D2P2A2 D2P4A2 Š D2P2A3 D2P4A3 -5 17/. - TiO -6 -7

Fig. 1 Selected MOs energy levels for the designed dyes at the B3LYP/6-311+G(2d,p)/SMD level together with TiO₂ CB and I^-/I_3^- redox potential.

Table 1 Absorption wavelengths, oscillator strength, J_{sc} -max (mA cm⁻²) under ideal conditions where $\Phi_{inj} = \eta_{coll} = 1$, $\Gamma = 30$ nmol cm⁻² is considered for all dyes, and main contributions (f > 0.2) to the transitions of dyes in acetonitrile at the CAM-B3LYP/6-311+G(2d,p)/SMD level

	λ				Main contribution
Compound	nm	eV	f	$J_{\rm sc}$ -max	to the transition
D1P2A1	549.30	2.26	2.47	21.08	$H \rightarrow L (64\%)$
	389.14	3.19	0.24		$H-1 \rightarrow L(42\%)$
	363.28	3.41	0.25		$H \rightarrow L+1 (34\%)$
	309.52	4.01	0.32		$H-2 \rightarrow L(21\%)$
D1P4A1	547.36	2.27	2.40	23.02	$H \rightarrow L(58\%)$
	536.54	2.31	0.40		$H \rightarrow L+2 (60\%)$
	426.43	2.91	0.39		$H-1 \rightarrow L(30\%)$
	358.53	3.46	1.18		$H-1 \rightarrow L+2 (15\%)$
	355.61	3.49	0.38		$H-1 \rightarrow L+2$ (33%)
	297.46	4.17	0.34		$H-2 \rightarrow L(31\%)$
D1P2A2	497.49	2.49	2.44	16.07	$H \rightarrow L(51\%)$
	404.05	3.07	0.31		$H \rightarrow L+1 (47\%)$
	299.51	4.14	0.30		$H-2 \rightarrow L(21\%)$
D1P4A2	517.24	2.40	3.03	20.16	$H \rightarrow L (40\%)$
	345.93	3.58	1.71		$H-3 \rightarrow L+2$ (30%)
D1P2A3	519.24	2.39	1.85	17.63	$H \rightarrow L (76\%)$
	294.64	4.21	0.30		$H-2 \rightarrow L(25\%)$
	281.58	4.40	0.24		$H-6 \rightarrow L(68\%)$
D1P4A3	538.99	2.30	1.75	21.06	$H \rightarrow L (50\%)$
	366.70	3.38	0.86		$H-2 \rightarrow L+1 (37\%)$
	343.14	3.61	1.09		$H \rightarrow L+2 (30\%)$
	287.05	4.32	0.33		$H-3 \rightarrow L (38\%)$
D2P2A1	528.35	2.35	2.55	20.01	$H \rightarrow L (39\%)$
	431.47	2.87	0.31	20101	$H-1 \rightarrow L (40\%)$
	395.92	3.13	0.39		$H \rightarrow L+1 (30\%)$
D2P4A1	521.95	2.38	2.30	20.18	$H \rightarrow L (49\%)$
D21 III	482.11	2.50 2.57	0.74	20.10	$H^{-1} \rightarrow L^{+2} (30\%)$
	453.11	2.74	0.60		$H-2 \rightarrow L(28\%)$
	411.30	3.01	0.48		$H \rightarrow L+1 (39\%)$
	337.22	3.68	0.55		$H \rightarrow L+2 (27\%)$
D2P2A2	485.52	2.55	2.58	16.07	$H \rightarrow L (56\%)$
D21 2112	444.18	2.79	0.28	10.07	$H^{-1} \rightarrow L(22\%)$
	410.19	3.02	0.20		$H^{-1} \rightarrow L^{+1} (27\%)$
	364.59	3.40	0.52		$H \rightarrow L (23\%)$
	329.57	3.76	1.06		$H \rightarrow L(23\%)$ $H \rightarrow L+8(20\%)$
D2P4A2	512.65	2.42	2.95	19.87	$H \rightarrow L (50\%)$
D2F4A2	444.18	2.42 2.79	0.28	19.07	$H \rightarrow L (30\%)$ $H-1 \rightarrow L (22\%)$
	444.18	3.02	0.28		$H-1 \rightarrow L(22\%)$ $H-1 \rightarrow L+1 (27\%)$
	364.59	3.40	0.29		$H \rightarrow L (23\%)$
	364.59 329.57	3.40 3.76	0.52 1.06		$H \rightarrow L (23\%)$ $H \rightarrow L+8 (20\%)$
D2P2A3				12 60	
DZPZAS	498.75	2.49	2.02	13.62	$H \rightarrow L (45\%)$ $H \rightarrow L + 1 (24\%)$
D0D442	328.25	3.78	0.24	10.00	$H-1 \rightarrow L+1 (24\%)$
D2P4A3	523.42	2.37	2.28	19.69	$H \rightarrow L (34\%)$
	336.76	3.68	1.03		$H-4 \rightarrow L+1 (21\%)$
	333.77	3.71	1.14		$H-4 \rightarrow L+1 (21\%)$

absorption range. These combined data can be summarized by the light harvesting efficiency value, LHE (λ), whose curves along with the AM 1.5G solar spectrum are reported in the ESI,† Fig. S1. Table 1 reports the summation of the products of the LHE (λ) with the AM 1.5G (λ) solar spectrum within the wavelength range of 280 to 830 nm. Here we made the assumption that under optimal conditions, where $\Phi_{inj} = \eta_{coll} = 1$, this sum corresponds to J_{sc} -max. This information highlights that the most favorable LHE (λ) performance is achieved when employing the acceptor group A1; furthermore, D1 yields the most promising results as an electron donor. Although the PTZ donor group results in a high absorption intensity (see Fig. 2), it exhibits a narrower optical absorption range, and best outcomes are observed when it is paired with the electron acceptor A1, whereas

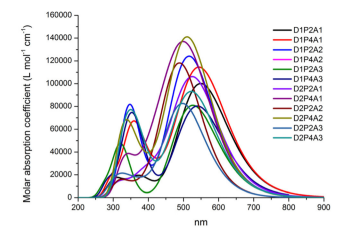


Fig. 2 Absorption spectra of dyes in acetonitrile at the TD-CAM-B3LYP/ 6-311+G(2d,d)/SMD level. The spectra are Gaussian-broadened with a half width half maximum of 0.35 eV.

results are more modest when combined with A3 and A2 acceptor groups. In all studied cases, increasing the length of LCC leads to a higher LHE (λ).

To visualize the electron density during the $S_0 \rightarrow S_1$ transition, an analysis of the dyes' electronic structure was conducted. The resulting electron density difference (EDD) maps are presented in Fig. 4 for D1P2A1 and D1P4A1, while those for the remaining dyes can be found in the ESI,† Fig. S2. In these maps, the blue region indicates an increase in electron density, while the red region signifies a decrease in electron density due to the excitation of electrons during the transition. Both figures clearly show that the acceptor unit exhibits the highest electron density, whereas the donor unit displays the lowest electron density. As a consequence, we can conclude that during the transition, there is a charge transfer from the donor unit to the acceptor unit in all investigated dyes.

3.2 Ionization potential, electron affinity and reorganization energy

The ionization potential (IP) refers to the energy required to extract an electron from a molecule's neutral state. Conversely, electron affinity (EA) is the energy gap between the neutral molecule and its anionic form, both in their lowest energy charge-transfer states. The values of IP and EA play a crucial role in determining the suitability of a dye as a photosensitizer for DSSCs. These parameters were computed using the following equations:

$$IP = E^{+}(M^{+}) - E(M)$$
(11)

$$EA = E(M) - E^{-}(M^{-})$$
 (12)

where $E(\mathbf{M})$, $E^+(\mathbf{M}^+)$ and $E^-(\mathbf{M}^-)$ are the energies of the dyes related to the neutral, cationic and anionic optimized structures. The computed values of ionization potentials, electron affinities and intramolecular reorganization energies, for the investigated dyes are presented in Table 2. A lower value of λ_i is indicative of slower recombination processes.

This article is licensed under a Creative Commons Attribution-NonCommercial 3.0 Unported Licence.

Open Access Article. Published on 08 2023. Downloaded on 14-10-24 20.05.05.

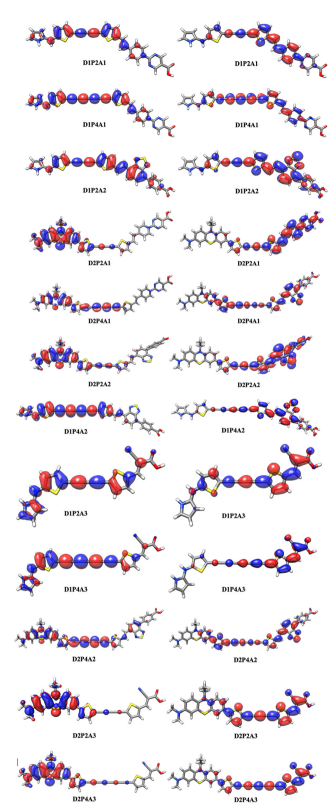


Fig. 3 Diagram of natural transition orbitals (NTOs, $S_0 \rightarrow S_1$) of the dyes in acetonitrile at the TD-CAM-B3LYP/6-311+G(2d,p)/SMD level. The surfaces are generated with an isovalue at 0.02.

The IP values vary within a range of 4.63 to 5.11 eV. Notably, D1P4A1 stands out with the highest IP value, suggesting



Fig. 4 Electron density difference (EDD) maps of D1P2A1 and D1P4A1 dyes simulated by Gaussview (the red and blue colors indicate a decrease and increase of charge densities, respectively). The density differences are 0.0001 a.u.

reduced ability for electron injection into the TiO₂ semiconductor. Consistently, it displays one of the higher EA values, specifically 3.78 eV, signifying a more favorable regeneration process. In general, dyes featuring D1 exhibit the higher IP values, whereas the A1 acceptor, characterized by a positive charge on the pyridine nitrogen atom, induces higher EA values to the dyes. Both λ_h and λ_e values are positive, signifying favorable hole and electron transfer characteristics for the designed sensitizers. However, those with the D2 donor group exhibit smaller λ_e values compared to λ_h . This implies that for these dyes, electron mobility exceeds hole mobility due to the inverse relationship between the charge transfer rate, $k_{\rm CT}$, and the reorganization energy.^{81–84}

In the case of the D1 donor group, λ_e values are lower than λ_h only when the LCC length is shorter (n = 2). As the chain length increases, the mobility of negative charges decreases relative to that of holes. The discrepancy between λ_h and λ_e suggests a more balanced transport in dyes employing D1. From the data presented in Table 2, it results that λ_i is lower for compounds featuring D1, thus indicating that D1-containing dyes exhibit superior charge transfer abilities, a trait that is advantageous for efficient photocurrent generation and, as a result, indicates improved performance in solar devices.

3.3 Electron injection and electron regeneration driving forces ($\Delta G^{\text{inj}}, \Delta G^{\text{reg}}$)

The fundamental importance of aligning the frontier orbitals of both the dye and the semiconductor in order to achieve efficient electron injection and dye regeneration is widely recognized.^{85–88} To delve deeper into the charge transfer

Table 2 Calculated values of charge transfer rate (k_{CT} in s⁻¹), reorganization energies λ_i , λ_e , λ_h (eV), ionization potential (IP in eV), electron affinity (EA in eV) and V_{12} (eV) for all the dyes

Compound	$k_{ m CT} imes 10^{13}$	$\lambda_{\mathbf{h}}$	λ_{e}	λ_{i}	IP	EA	V_{12}
D1P2A1	4.99	0.249	0.200	0.449	5.07	3.71	0.392
D1P4A1	2.54	0.255	0.261	0.515	5.11	3.78	0.399
D1P2A2	5.73	0.176	0.240	0.416	4.96	3.14	0.352
D1P4A2	3.49	0.214	0.263	0.478	5.08	3.22	0.382
D1P2A3	4.51	0.258	0.200	0.458	5.03	3.26	0.391
D1P4A3	2.41	0.244	0.278	0.522	5.06	3.33	0.402
D2P2A1	0.99	0.381	0.233	0.614	4.75	3.73	0.420
D2P4A1	1.14	0.367	0.235	0.601	4.75	3.81	0.422
D2P2A2	0.84	0.373	0.242	0.615	4.63	3.16	0.387
D2P4A2	1.58	0.321	0.235	0.556	4.65	3.25	0.392
D2P2A3	0.61	0.369	0.260	0.629	4.70	3.38	0.355
D2P4A3	0.32	0.371	0.312	0.683	4.72	3.53	0.343

process, driving force for electron injection, ΔG^{inj} , and regeneration, ΔG^{reg} , were determined. ΔG^{inj} quantifies the free energy associated with the electron injection process, represents the difference between the excited electron of the sensitizer and the injected electron onto the TiO₂ surface and is given by:

$$\Delta G^{\rm inj} = E^{\rm ox-dye^*} - E_{\rm CB} \tag{13}$$

where E_{CB} is the reduction potential of the TiO₂ CB (E_{CB} = 4.0 eV), $E^{\text{ox-dye}*}$ is the oxidation potential of the dye in its excited state and can be computed as:

$$E^{\text{ox-dye}^{*}} = E^{\text{ox-dye}} - E_{0-0}$$
(14)

where $E^{\text{ox-dye}}$ refers to the oxidation potential of the ground state, while E_{0-0} is the vertical electronic transition energy corresponding to λ_{max} . Dye regeneration is the process wherein the oxidized dye is restored by the redox electrolyte after electron injection. Several studies have shown that this parameter significantly impacts the photovoltaic performance of DSSCs. The driving force for the regeneration can be determined from $E^{\text{ox-dye}}$ as follows:

$$\Delta G^{\text{reg}} = E^{\text{redox-electrolyte}} - E^{\text{ox-dye}}$$
(15)

where the redox $E^{\text{redox-electrolyte}}$ redox potential is -4.8 eV. The values of ΔG^{inj} and ΔG^{reg} are presented in Table 3.

The $E^{\text{ox-dye}*}$ values reveal that dyes featuring D2 undergo oxidation more easily than those with the D1 donor group. Additionally, the ΔG^{inj} values consistently show a negative trend, highlighting that electron injection is a thermodynamically favored process. Notably, these values decrease as we move from the series characterized by D1 to the one featuring D2. In general, lower ΔG^{inj} values correspond to enhanced efficiency in electron injection, consequently, the data from Table 2 suggest that electron injection is most effective in D2containing compounds. However, it should be noted that k_{inj} , which represents the inverse of injection time, also depends on the reorganization energy, as mentioned earlier, therefore a more detailed investigation into the adsorption of the sensitizer on TiO₂ is necessary for more accurate predictions.

Efficient dye regeneration necessitates ΔG^{reg} values to be negative and within the range of 0.2–0.3 eV.⁷³ Our calculations reveal that dyes with D2 exhibit positive ΔG^{reg} values, indicating

Table 3 Calculated $E^{\text{ox-dye}}$, $E^{\text{ox-dye}*}$, ΔG^{inj} , ΔG^{reg} (eV), q_{CT} (a.u.) and D_{CT} in Å for all the dyes

Compound	E ^{ox-dye}	$E^{\text{ox-dye}_{\star}}$	ΔG^{inj}	ΔG^{reg}	$q_{\rm CT}$	$D_{\rm CT}$
D1P2A1	5.09	2.83	-1.17	-0.29	0.880	10.450
D1P4A1	5.12	2.86	-1.14	-0.32	0.940	13.137
D1P2A2	4.93	2.44	-1.16	-0.13	0.842	10.840
D1P4A2	5.03	2.64	-1.14	-0.23	0.852	12.564
D1P2A3	5.05	2.66	-1.34	-0.25	0.752	8.932
D1P4A3	5.07	2.77	-1.23	-0.27	0.772	10.131
D2P2A1	4.76	2.42	-1.58	0.04	0.934	13.707
D2P4A1	4.78	2.40	-1.60	0.03	0.987	16.671
D2P2A2	4.73	2.18	-1.82	0.07	0.864	11.240
D2P4A2	4.75	2.33	-1.67	0.05	0.878	13.786
D2P2A3	4.75	2.27	-1.73	0.05	0.875	14.173
D2P4A3	4.76	2.39	-1.61	0.04	0.893	15.334

that the regeneration process is thermodynamically unfavorable for these sensitizers. In contrast, compounds featuring D1 yield conducive values toward regeneration. To evaluate how charge transfer occurs within the dyes during electron transition, we employed Multiwfn 3.8 software to assess parameters such as transferred charge (q_{CT}) and transferred charge distance $(D_{\rm CT})$.⁷² Upon examining the numerical values of $q_{\rm CT}$ shown in Table 2, it becomes evident that sensitizers featuring D2 tend to show slightly higher $q_{\rm CT}$ values in comparison to those with D1. In general, the most favorable results are observed when the donor is paired with A1, indicating that this acceptor group facilitates the formation of a charge-separated state. Simulated charge transfer distance corroborates this notion, as a longer distance is conducive to charge separated state of dyes. Considering all the parameters outlined in Tables 2 and 3, it can be inferred that D1 generally confers superior properties for use in DSSC applications when coupled to A1. In a broader context, sensitizers featuring D1 are predicted to perform well in photovoltaic applications. On the other hand, it is worth noting that in the specific configuration we studied, $\Delta G^{\rm reg}$ values may hinder regeneration in dyes featuring D2, despite they show excellent values for the other calculated parameters.

3.4 TiO₂-dye adsorption

To effectively predict the performance of a solar cell device, it is essential to carry out prediction studies focused on the interface between the dye and the semiconductor. In pursuit of this objective, we have employed a $(TiO_2)_{14}$ cluster as a representative model for the TiO₂ semiconductor surface. As mentioned earlier, bidentate carboxylate groups have been utilized as anchoring agents due to the stability they offer, as supported by several studies.^{66–68} In Table 4 are reported the dipole moment, μ of dye@TiO₂, in comparison with that of free dyes, and the interaction energies, E_{int} , calculated as:

$$E_{\text{int}} = E_{\text{dye}(\underline{a})\text{TiO}_2} - (E_{\text{dye}} + E_{\text{TiO}_2})$$

Calculated Ti-O distances fall within the range of 1.96-2.10 Å, aligning closely with the theoretically reported Ti-O bond lengths (ranging from 2.03 to 2.24 Å) for different dye-TiO₂ clusters.73 These results are indicative of the favorable interaction between the dyes and the semiconductor surface, as supported by the calculated E_{int} values for dye@(TiO₂)₁₄. Additionally, Table 4 reveals that the calculated dipole moments of the dye@TiO₂ clusters are larger than those of the free dyes. These findings imply that the binding to the semiconductor enhances the transport properties. From Fig. 5 and Fig. S3 in the ESI,† it become evident that the HOMO levels of dyes@(- TiO_2 ₁₄ are localized to the sensitizers, preserving the HOMO characteristics found in isolated dyes. These HOMO values show an increase when compared to the free dyes. On the other hand, the LUMO levels, along with most of the LUMO+n levels, are primarily centered on the substrate, displaying characteristics similar to TiO₂.

Complexes	$E_{\rm int}$	μ
D1P2A1@(TiO ₂) ₁₄	-15.49	13.11 (10.97)
D1P4A1@(TiO ₂) ₁₄	-16.10	19.94 (15.34)
D1P2A2@(TiO ₂) ₁₄	-13.27	15.59 (11.24)
D1P4A2@(TiO ₂) ₁₄	-14.43	17.66 (12.46)
D1P2A3@(TiO ₂) ₁₄	-14.91	11.40 (9.53)
D1P4A3@(TiO ₂) ₁₄	-15.89	13.13 (11.55)
D2P2A1@(TiO ₂) ₁₄	-14.59	24.28 (19.56)
D2P4A1@(TiO ₂) ₁₄	-15.01	26.44 (23.94)
D2P2A2@(TiO ₂) ₁₄	-13.14	17.32 (14.79)
D2P4A2@(TiO ₂) ₁₄	-14.53	20.86 (18.22)
D2P2A3@(TiO ₂) ₁₄	-15.07	19.18 (15.51)
D2P4A3@(TiO ₂) ₁₄	-15.81	20.05 (16.74)

4. Conclusions

We conducted a theoretical investigation into the spectroscopic, electronic and photochemical properties of newly designed dyes which featured 2-amino pyrrole, D1, as the donor moiety, and were paired with three distinct withdrawing groups. Two of them, benzothiadiazole, A2, and cyanoacrylic acid, A3, are commonly used in DSSC applications, while a novel electron-attractor group, a derivative of pyridinium salt, was introduced for the first time. Additionally, a linear carbon chain (LCC) of varying lengths was used as the π -bridge. Analogs of these sensitizers were modeled using phenothiazine, D2, as the electron donor group, in place of D1, to compare the effectiveness of the 2-amino-pyrrole donor group. LHE (λ), as determined from the calculated absorption spectra, reveals that dyes containing D1 are expected to exhibit superior performance compared to their D2-containing counterparts. Furthermore, the introduction of the innovative electronattracting group A1 yields the most promising results. The EDD maps demonstrated charge transfer from the donor unit to the acceptor unit in all studied cases. D1-containing sensitizers exhibit lower reorganization energy (λ_i) values, resulting in faster charge transfer when compared to those with D2. Efficient electron injection is predicted for all sensitizers, particularly those featuring D2; however dyes with this donor

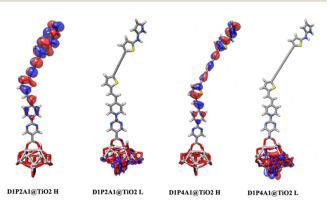


Fig. 5 Optimized geometries of the bidentate binding mode of D1P2A1 and D1P4A1 dyes on $(TiO_2)_{14}$. HOMOs (H) and LUMOs (L) are also depicted.

group show positive ΔG^{reg} values, implying an unfavorable regeneration process. Increasing the length of the LCC has a positive effect on the LHE (λ) and ΔG^{reg} but has a less favorable impact on ΔG^{inj} . The acceptor group A1 facilitates charge separation, as indicated by the calculated charge transfer values, q_{CT} . Simulation of dye@(TiO₂)₁₄ complexes suggests that the binding with the semiconductor enhanced the transport properties. In short, all calculated parameters indicate that the novel D1 and A1 groups are poised to deliver superior properties for DSSC applications. Consequently, we can conclude that most of the designed dyes may have the potential to serve as candidates for the fabrication of DSSCs.

Author contributions

Giuseppe Consiglio: conceptualization, investigation, writing, revision and editing. Adam Gorcyński: investigation, writing, revision and editing. Salvatore Petralia: investigation, writing, revision and editing. Giuseppe Forte: conceptualization, supervision, methodology, and writing original draft.

Conflicts of interest

The authors declare no conflicts of interest.

Acknowledgements

This work has been partially funded by the European Union (NextGeneration EU), through the MUR-PNRR project SAMO-THRACE (ECS00000022).

References

- 1 C. Battaglia, A. Cuevas and S. De Wolf, *Energy Environ. Sci.*, 2016, **9**, 1552–1576.
- 2 P. Y. Chen, J. Qi, M. T. Klug, X. Dang, P. T. Hammond and A. M. Belcher, *Energy Environ. Sci.*, 2014, 7, 3659–3665.
- 3 B. O'Regan and M. Grätzel, Nature, 1991, 353, 737-740.
- 4 M. L. Parisi, S. Maranghi and R. Basosi, *Renewable Sustable Energy Rev.*, 2014, **39**, 124–138.
- 5 X. Zhou, Y. Zhang, A. L. Porter, Y. Guo and D. Zhu, *Scientometrics*, 2014, **100**, 705–721.
- 6 S. Shalini, R. Balasundaraprabhu, T. Satish Kumar, N. Prabavathy, S. Senthilarasu and S. Prasanna, *Int. J. Energy Res.*, 2016, 40, 1303–1320.
- 7 M. K. Nazeeruddin, P. Péchy and M. Grätzel, *Chem. Commun.*, 1997, 1705–1706.
- 8 M. K. Nazeeruddin, P. Péchy, T. Renouard, S. M. Zakeeruddin, R. Humphry-Baker, P. Comte, P. Liska, L. Cevey, E. Costa, V. Shklover, L. Spiccia, G. D. Deacon, C. A. Bignozzi and M. Grätzel, *J. Am. Chem. Soc.*, 2001, 123, 1613–1624.
- 9 R. Buscaino, C. Baiocchi, C. Barolo, C. Medana, M. Grätzel, M. K. Nazeeruddin and G. Viscardi, *Inorg. Chim. Acta*, 2008, 361, 798–805.

- 10 Z. Z. Lu, J. Peng, W. De, C. H. Lin, C. G. Wu, K. C. Ho, Y. C. Lin and K. L. Lu, *Eur. J. Inorg. Chem.*, 2016, 1214–1224.
- 11 J. F. Huang, J. M. Liu, P. Y. Su, Y. F. Chen, Y. Shen, L. M. Xiao, D. B. Kuang and C. Y. Su, *Electrochim. Acta*, 2015, **174**, 494–501.
- 12 A. Yella, H. W. Lee, H. N. Tsao, C. Yi, A. K. Chandiran, M. K. Nazeeruddin, E. W. G. Diau, C. Y. Yeh, S. M. Zakeeruddin and M. Grätzel, *Science*, 2011, 334, 629–634.
- 13 S. Mathew, A. Yella, P. Gao, R. Humphry-Baker, B. F. Curchod, N. Ashari-Astani, I. Tavernelli, U. Rothlisberger, M. K. Nazeeruddin and M. Grätzel, *Nat. Chem.*, 2014, 6, 242–247.
- 14 G. Consiglio, S. Failla, C. G. Fortuna, L. D'Urso and G. Forte, *Comput. Theor. Chem.*, 2015, **1067**, 1–6.
- 15 I. P. Oliveri, G. Forte, G. Consiglio, S. Failla and S. Di Bella, *Inorg. Chem.*, 2017, 56, 14206–14213.
- 16 S. Aghazada, P. Gao, A. Yella, G. Marotta, T. Moehl, J. Teuscher, J. E. Moser, F. De Angelis, M. Grätzel and M. K. Nazeeruddin, *Inorg. Chem.*, 2016, 55, 6653–6659.
- N. V. Krishna, J. V. S. Krishna, M. Mrinalini,
 S. Prasanthkumar and L. Giribabu, *ChemSusChem*, 2017, 10, 4668–4689.
- 18 C. C. Chen, J. S. Chen, V. S. Nguyen, T. C. Wei and C. Y. Yeh, Angew. Chem., 2021, 60(9), 4886–4893.
- 19 M. Xu, S. Wenger, H. Bala, D. Shi, R. Li, Y. Zhou, S. M. Zakeeruddin, M. Grätzel and P. Wang, *J. Phys. Chem. C*, 2009, **113**, 2966–2973.
- 20 S. Ahmad, E. Guillén, L. Kavan, M. Grätzel and M. K. Nazeeruddin, *Energy Environ. Sci.*, 2013, **6**, 3439–3466.
- 21 C. Zhong, J. Gao, Y. Cui, T. Li and L. Han, *J. Power Sources*, 2015, **273**, 831–838.
- 22 A. Venkateswararao, P. Tyagi, K. R. J. Thomas, P. W. Chen and K. C. Ho, *Tetrahedron*, 2014, **70**, 6318–6327.
- 23 S. S. K. Raavi, P. Docampo, C. Wehrenfennig, M. J. P. Alcocer, G. Sadoughi, L. M. Herz, H. J. Snaith and A. Petrozza, *J. Phys. Chem. C*, 2014, **118**, 16825–16830.
- 24 M. Mao, X. L. Zhang, X. Q. Fang, G. H. Wu, S. Y. Dai, Q. H. Song and X. X. Zhang, *J. Power Sources*, 2014, 268, 965–976.
- 25 J. Liu, X. Sun, Z. Li, B. Jin, G. Lai, H. Li, C. Wang, Y. Shen and J. Hua, *J. Photochem. Photobiol.*, *A*, 2014, **294**, 54–61.
- 26 T. Kitamura, M. Ikeda, K. Shigaki, T. Inoue, N. A. Anderson, X. Ai, T. Lian and S. Yanagida, *Chem. Mater.*, 2004, **16**, 1806–1812.
- 27 S. L. Li, K. J. Jiang, K. F. Shao and L. M. Yang, *Chem. Commun.*, 2006, 2792–2794.
- 28 R. Chen, X. Yang, H. Tian, X. Wang, A. Hagfeldt and L. Sun, *Chem. Mater.*, 2007, **19**, 4007–4015.
- 29 X. Q. Zhu, Z. Dai, A. Yu, S. Wu and J. P. Cheng, *J. Phys. Chem. B*, 2008, **112**, 11694–11707.
- 30 R. Y. Y. Lin, F. L. Wu, C. T. Li, P. Y. Chen, K. C. Ho and J. T. Lin, *ChemSusChem*, 2015, 8, 2503–2513.
- 31 Y. Hua, S. Chang and D. Huang, *Chem. Mater.*, 2013, 25, 2146–2153.
- 32 M. Ling Jang, J. J. Wen, Z. M. Chen, W. H. Tsai, T. C. Lin, T. J. Chow and Y. J. Chang, *ChemSusChem*, 2019, 8, 3654–3665.
- 33 Z. B. Xie, A. Midya, K. P. Loh, S. Adams, D. J. Blackwood, J. Wang, X. J. Zhang and Z. K. Chen, *Prog. Photovoltaics*, 2010, **18**, 573-581.

- 34 Y. Xie, Y. Tang, W. Wu, Y. Wang, J. Liu and X. Li, J. Am. Chem. Soc., 2015, 137, 14055–14058.
- 35 A. F. Buene, A. Hagfeldt and B. H. Hoff, *Dyes Pigments.*, 2019, **169**, 66–72.
- 36 A. F. Buene, E. E. Ose, A. G. Zakariassen, A. Hagfeldt and B. H. Hoff, *J. Mater. Chem. A*, 2019, 7, 7581–7590.
- 37 M. L. Han, Y. Z. Zhu and S. Liu, J. Power Sources, 2018, 387, 117–125.
- 38 T. Hua, Z. S. Huang, K. Cai, L. Wang, H. Tang, H. Meier and D. Cao, *Electrochim. Acta*, 2019, **302**, 225–233.
- 39 A. Slodek, D. Zych, S. Golba, S. Zimosz and E. Schab-Balcerzak, J. Mater. Chem. C, 2019, 7, 5830–5840.
- 40 D. Devadiga, M. Selvakumar, P. Shetty, M. S. Santosh, R. S. Chandrabose and S. Karazhanov, *Int. J. Energy Res.*, 2021, 45, 6584–6643.
- 41 A. Yella, C. L. Mai, S. M. Zakeeruddin, S. N. Chang,
 C. H. Hsieh, C. Y. Yeh and M. Grätzel, *Angew. Chem.*, 2014, 53(11), 2973–2977.
- 42 J. Wang, H. Wu, L. Jin, J. Zhang, Y. Yuan and P. Wang, ChemSusChem, 2017, 10, 2962-2967.
- 43 Y. Ding, J. Li, S. Liu, Y. Tan, K. Shi, Y. Jiao, B. Mi, W. Xu and Z. Gao, *Dyes Pigm.*, 2021, **194**, 109664.
- 44 Z. Z. Sun, S. Feng and W. L. Ding, *Sol. Energy*, 2020, **195**, 491–498.
- 45 K. M. Lee, V. Suryanarayanan and K. C. Ho, *J. Power Sources*, 2009, **188**, 635–641.
- 46 J. Wu, Z. Lan, J. Lin, M. Huang and P. Li, J. Power Sources, 2007, 173, 585–591.
- 47 E. Mosconi, A. Selloni and F. De Angelis, *J. Phys. Chem. C*, 2012, **116**, 5932–5940.
- 48 H. Tian, X. Yang, R. Chen, R. Zhang, A. Hagfeldt and L. Sun, J. Phys. Chem. C, 2008, 112, 11023–11033.
- 49 T. Le Bahers, F. Labat, T. Pauportè and I. Ciofini, *Phys. Chem. Chem. Phys.*, 2010, **12**, 14710–14719.
- 50 H. Fang, J. Ma, M. J. Wilhelm, B. G. DeLacy and H. L. Dai, *Part. Part. Syst. Charact.*, 2021, 38, 2000220.
- 51 G. Consiglio, A. Gorcynski, S. Petralia and G. Forte, *RSC Adv.*, 2023, **13**, 1019–1030.
- 52 G. Consiglio, A. Gorcynski, S. Petralia and G. Forte, *Dalton Trans.*, 2023, **52**, 15995–16004.
- 53 G. Forte, L. D'Urso, E. Fazio, S. Patanè, F. Neri, O. Puglisi and G. Compagnini, *Appl. Surf. Sci.*, 2013, **272**, 76–81.
- 54 E. Fazio, L. D'Urso, G. Consiglio, A. Giuffrida, G. Compagnini, O. Puglisi, S. Patanè, F. Neri and G. Forte, J. Phys. Chem. C, 2014, 118, 28812–28819.
- 55 N. R. Agarwai, A. Lucotti, M. Tommasini, W. A. Chalifoux and R. R. Tykwinski, *J. Phys. Chem. C*, 2016, **120**(20), 11131-11139.
- 56 P. Marabotti, A. Milani, A. Lucotti, L. Brambilla,
 M. Tommasini, C. Castiglioni, P. Mecik, B. Pigulski,
 S. Szafert and C. S. Casari, *Carbon Trends*, 2021, 5, 100115.
- 57 S. Eisler, A. D. Slepkov, E. Elliott, T. Luu, R. McDonald, F. A. Hegmann and R. Tykwinski, *J. Am. Chem. Soc.*, 2005, 127, 2666–2676.
- 58 L. K. Zaitriand and S. M. Mekelleche, *Mol. Phys.*, 2020, **118**, 1–10.

- 59 C. S. Casari, M. Tommasini, R. R. Tykwinski and A. Milani, *Nanoscale*, 2016, **8**, 4414–4435.
- 60 F. Cataldo, Polyynes: Synthesis, Properties, and Applications, 2005.
- 61 W. H. Liu, I. C. Wu, C. H. Lai, C.-H. Lai, P. T. Chou, Y. T. Li, C. L. Chen, Y. Y. Hsu and Y. Chi, *Chem. Commun.*, 2008, 5152–5154.
- 62 C. Bonaccorso, F. De Rossi, M. Panigati, C. G. Fortuna, G. Forte, T. M. Brown, G. M. Farinola and G. Musumarra, *Tetrahedron*, 2015, **71**, 7260–7266.
- 63 M. J. Frisch, G. W. Trucks, H. B. Schlegel, G. E. Scuseria, M. A. Robb, J. R. Cheeseman, G. Scalmani, V. Barone, G. A. Petersson, H. Nakatsuji, X. Li, M. Caricato, A. V. Marenich, J. Bloino, B. G. Janesko, R. Gomperts, B. Mennucci, H. P. Hratchian, J. V. Ortiz, A. F. Izmaylov, J. L. Sonnenberg, D. Williams-Young, F. Ding, F. Lipparini, F. Egidi, J. Goings, B. Peng, A. Petrone, T. Henderson, D. Ranasinghe, V. G. Zakrzewski, J. Gao, N. Rega, G. Zheng, W. Liang, M. Hada, M. Ehara, K. Toyota, R. Fukuda, J. Hasegawa, M. Ishida, T. Nakajima, Y. Honda, O. Kitao, H. Nakai, T. Vreven, K. Throssell, J. A. Montgomery, Jr., J. E. Peralta, F. Ogliaro, M. J. Bearpark, J. J. Heyd, E. N. Brothers, K. N. Kudin, V. N. Staroverov, T. A. Keith, R. Kobayashi, J. Normand, K. Raghavachari, A. P. Rendell, J. C. Burant, S. S. Iyengar, J. Tomasi, M. Cossi, J. M. Millam, M. Klene, C. Adamo, R. Cammi, J. W. Ochterski, R. L. Martin, K. Morokuma, O. Farkas, J. B. Foresman and D. J. Fox, Gaussian 16 Rev C.01, Gaussian, Inc., Wallingford CT, 2016.
- 64 J. Preat, C. Michaux, D. Jacquemin and E. A. Perpete, J. Phys. Chem. C, 2009, 113, 16821–16833.
- 65 M. Pastore, E. Mosconi, F. De Angelis and M. Gratzel, J. Phys. Chem. C, 2010, 114, 7205–7212.
- 66 D. Fadili, Z. M. E. Fahim, S. M. Bouzzine, O. T. Alaoui and M. Hamidi, *Comput. Theor. Chem.*, 2022, **1210**, 113645.
- 67 C. O'Rourke and D. Bowler, *J. Phys.: Condens. Matter*, 2014, 26, 195302.
- 68 M. Wagstaffe, A. G. Thomson, M. J. Jackman, M. Torres-Molina, K. L. Syres and K. Handrup, *J. Phys. Chem. C*, 2016, 120(3), 1693–1700.

- 69 R. Sánchez-de-Armas, M. A. San Miguel, J. Oviedo and J. F. Sanz, *Phys. Chem. Chem. Phys.*, 2012, 14(1), 225–233.
- 70 R. Sánchez-de-Armas, M. A. San-Miguel, J. Oviedo, A. Márquez and J. F. Sanz, *Phys. Chem. Chem. Phys.*, 2011, 13(4), 1506–1514.
- 71 R. Sánchez-de-Armas, J. OviedoLópez, M. A. San-Miguel, J. F. Sanz, P. Ordejón and M. Pruneda, *J. Chem. Theory Comput.*, 2010, 6(9), 2856–2865.
- 72 T. Lu and F. Chen, J. Comput. Chem., 2012, 33, 580-592.
- 73 L. L. Estrella, M. P. Balanay and D. H. Kim, *J. Phys. Chem. A*, 2016, **120**, 5917–5927.
- 74 R. A. Marcus, J. Chem. Phys., 1956, 24, 966-978.
- 75 R. A. Marcus and N. Sutin, *Rev. Bioenerget.*, 1985, **811**, 265–322.
- 76 M. P. Balany and D. H. Kim, THEOCHEM, 2009, 910, 20-26.
- 77 T. P. Nguyen, J. H. Shim and J. Y. Lee, J. Phys. Chem. C, 2015, 119, 11301–11310.
- 78 J. C. Cave and M. D. Newton, *Chem. Phys. Lett.*, 1996, 249, 15–19.
- 79 J. Preat, A. Hagfeldt and E. A. Perpete, *Energy Environ. Sci.*, 2011, 4, 4537–4549.
- 80 J. B. Asbury, Y. Q. Wang, E. C. Hao, H. N. Ghosh and T. Q. Lian, *Res. Chem. Intermed.*, 2001, 27, 393–406.
- 81 G. R. Hutchison, M. A. Ratner and T. J. Marks, J. Am. Chem. Soc., 2005, 127, 2339–2350.
- 82 H. Geng, Y. Niu, Q. Peng, Z. Shuai, V. Coropceanu and J. L. Bredas, J. Chem. Phys., 2011, 135, 104703.
- 83 A. Amini and A. Harriman, *J. Photochem. Photobiol.*, *C*, 2003, 4, 155–177.
- 84 Y. K. Lan and C. I. Huang, J. Phys. Chem. B, 2008, 112, 14857–14862.
- 85 A. Hagfeldt, G. Boschloo, L. Sun and H. Pettersson, *Chem. Rev.*, 2010, **110**, 6595–6663.
- 86 M. Pastore, S. Fantacci and F. De Angelis, J. Phys. Chem. C, 2013, 117(8), 3685–3700.
- 87 T. Manzoor, S. Asmi, S. Niaz and A. H. Pandith, *Int. J. Quantum Chem.*, 2017, **117**, 1–14.
- 88 D. Unny, G. R. Kandregula and K. Ramanujam, New J. Chem., 2021, 45, 16989–17001.