Review

Functional Organic Materials for Photovoltaics: The Synthesis as a Tool for Managing Properties for Solid State Applications

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Abstract: The continuous increase in the global energy demand deeply impacts the environment. Consequently, the research is moving towards more sustainable forms of energy production, storage and saving. Suitable technologies and materials are fundamental to win the challenge towards a greener and more eco-friendly society. Organic π-conjugated materials, including small molecules, oligomers and polymers are a wide and versatile class of functional materials with great potentiality, as they can be used as active matrixes in the fabrication of lightweight, flexible, cheap and large area devices. Their chemical and physical properties, both at a molecular level and mainly in the solid state, are a result of many factors, strictly related to the conjugated structure and functional groups on the backbone, which control the intermolecular forces driving solid state aggregations. The synthesis, through the molecular design, the choice of conjugated backbone and functionalization, represents the first and most powerful tool for finely tuning the chemico-physical properties of organic materials tailored for specific applications. In the present review, we report an overview of our works focused on synthetic methodologies, characterization, structure-properties correlation studies and applications of organic materials designed for energy-involving solid-state applications, organic photovoltaics in particular. The impact of functionalization on electro-optical properties and performance in device are discussed, also in relation to the specific applications.

Keywords: energy; organic photovoltaics; functional organic materials; synthetic methodologies; structure-properties correlation

1. Introduction

The energetic demand can be approached both from the production and storage point of view as well as from the energy saving side, and the potential of organic materials can meet both strategies. Organic materials with an extended π-conjugated backbone have semiconducting properties exploitable in a number of applications [1,2], such as organic light emitting diodes [3,4], organic photovoltaics [5–7], photocatalysis [8,9], batteries [10,11], and smart applications [12,13]. Due to their great chemical versatility, organic materials can be suitably addressed by synthesis, to be incorporated into various devices for many applications. On the other hand, their chemico-physical properties can be easily tuned by the synthesis, working on molecular conjugated skeleton and functionalization. These structural characteristics, in fact, drive electronic and steric effects, then electro-optical properties and molecular interactions in the solid state, determining the performance of organic materials in device. HOMO and LUMO energy levels, energy gap, charge mobility, thermal and chemical stability, solubility in different solvents, processability, and solid-state aggregation are all properties underlying synthetic control. In this view, clarifying the relationship between chemical structure and the properties of organic semiconductors plays a key role in order to extract their best performance in technological applications.
Although the molecular conjugated backbone drives the basic electro-optical properties, functionalization deeply affects these properties, and the type of functional group and its position onto the skeleton can drastically change the behaviour of organic semiconductors and then their final performance. In line with these considerations, in this review we present our contribution in the field of organic semiconductor materials for photovoltaic energy, highlighting synthetic methodologies, electrical and optical properties, intermolecular interactions, solid-state organization and performance in device. Organometallic synthetic methodologies were carried out to prepare the organic materials, exploiting their wide versatility, mild reaction conditions and stereoselectivity which allow a rigorous control of the stereochemistry of final compounds. The synthetic methodologies were also selected in order to realize a proper functionalization of the conjugated skeleton, by regarding both the type of functional group and its position on the polymer or molecular skeleton. Electrical, optical and theoretical investigations were performed to deeply understand the effect of chemical structures and functionalization on chemico-physical properties, in order to optimize the performance of the final compound in photovoltaic devices. Then, the performance of synthesized organic materials was investigated by BHJ or DSSC solar cells. Finally, the chemico-physical properties and photovoltaic performance were analyzed and discussed in relation to the chemical structure of materials.

2. Results

2.1. Fluorine Impact on Arylenevinylene Materials: Synthesis, Chemico-Physical Properties and Application in BHJ Solar Cells

Solar energy is the most abundant and cleanest renewable energy sources available. Organic solar cells (OSCs) as one of the most promising technologies for harvesting solar energy have attracted broad interest thanks to their advantages such as low cost, flexibility, lightweight, semi-transparency, large area, non-toxicity and roll-to-roll large-scale processability. The bulk heterojunction BHJ is a very simple configuration of OSC in which a π-conjugated donor material is blended with an electron acceptor (very often a fullerene derivative) to yield the active layer sandwiched between the electrodes [14,15]. On the way to technological applications, some issues have to be overcome, since critical drawbacks limit the performance of BHJ solar cells, such as efficiency and the low chemical stability of the organic active layer [16,17]. The efficiency of OSCs is attributed to a combination of several features, such as absorption capability, charge-carrier mobility, exciton diffusion length, and the morphology of the active layer plays a key role affecting all these features. To gain good solar cell efficiencies, the electron donor material should possess some characteristics: an extended light absorption matching, as much as possible, the solar spectrum for effective harvesting of the solar photons; high extinction coefficient; HOMO and LUMO energy levels, correctly aligned with those of the acceptor; good hole and exciton transport properties; thermal and chemical stability. So that an ideal donor material blended with PCBM in a BHJ should have a low band gap (between 1.2 and 1.9 eV), with the HOMO energy level in the range $-5.2/-5.8$ eV and the LUMO energy level in the range $-3.8/-4.0$, to ensure efficient charge separation and high Voc [18]. Polyarylenevinylenes are a class of materials with low band gap, potentially suitable as donor system in BHJ [19,20]. Following our first results on synthetic methods aimed to prepare arylenevinylene oligomers and polymers with fluorinated double bond [21–25], and on investigations of the effect of fluorination on their chemico-physical properties [26–30], we became interested in studying the impact of the fluorinated vinylene units on the photovoltaic performance of a polyarylenevinylene as donors for BHJ solar cells [31]. For this purpose, we synthesized two soluble conjugated polymers alternating double bonds (either fluorinated or non-fluorinated) with the bis-thienyl-bis-alkoxybenzothiadiazole unit, PDTBTFV and PDTBTV (Figure 1).
2.1.1. Synthesis of Polyarylenevinylenes for BHJ Solar Cells

In order to synthesize arylenevinylene materials with fluorinated and non-fluorinated double bonds, we followed a methodological approach based on the organometallic chemistry, exploiting the Stille reaction, a palladium-catalysed cross-coupling between organo-stannane reagents and organo-halogen derivatives. The Stille reaction is a general and versatile synthetic protocol, which requires very mild reaction conditions, tolerates many functional groups and allows a rigorous stereo-chemical control of the reaction and then of the chemical structure of the final product. The general strategy draws inspiration from the synthetic procedure to prepare the (E)-1,2-difluoro-1,2-bis(tributystannyl)ethene M1, as reported in Scheme 1.

![Scheme 1. Synthetic route to reagent M1 and intermediates 1–4.](image)

We adopted a modified literature synthetic procedure, which allows preparing, in addition to the final difluorovinyl-bis-stannane M1, useful fluorinated vinyl intermediates (1–4) which can also be exploited, in turn, as organometallic reagents to prepare arylenevinylene oligomers containing fluorinated double bonds [21]. Compound M1 is a very useful organo-stannane reagent, suitable to undergo cross-coupling reactions in the Stille conditions, to yield polyarylenevinylenes with all-trans fluorinated double bonds. The stereo-regularity of the vinylene units in an arylenevinylene polymer is a key factor, since it determines the effective conjugation length of the backbone and then the chemico-physical and structural properties of the material. Compared to the Stille reaction, traditional synthetic protocols such as the Wittig [32–34] reaction, the Knoevenagel [35–38] condensation, the precursor routes Gilch [39] or Wessling [40] used to prepare arylenevinylene polymers do not allow an accurate control of the stereochemistry of the vinylene units, introducing defects in the conjugated backbone that are detrimental for electrical and optical performance. Materials with semiconductor properties require a high degree of structural regio- and stereo-regularity to exhibit their best performance in a device, so that the Stille reaction, with its high regio- and stereo-selectivity is very attractive as a synthetic methodology to prepare organic semiconducting materials. The analogous of M1 without fluorine atoms, i.e., (E)-1,2-bis(tributyldistannyl)ethene M2, used to prepare the non-fluorinated polyarylenevinylene, is a well-known reagent, easy to synthesize and exploited in many Stille reactions to prepare a lot of polyarylenevinylenes [25,41–43]. As the aryldihalide reagent partner of the organostannanes M1 and M2, we used 4,7-bis(5-
iodothiophen-2-yl)-5,6-bis(octyloxy)benzo[c][1,2,5]thiazole M3, which was synthesized as shown in Scheme 2.

Scheme 2. Synthetic route to reagent M3.

The synthesis of the two polymers, with (PDTBTFV) and without (PDTBTV) fluorinated vinylene units, was carried out as shown in Scheme 3, via the Stille cross-coupling, using Pd(PPH₃)₄ as the catalyst and CuI in stoichiometric amount, in DMF/THF as the solvent, at room temperature, for one week.

Scheme 3. Synthetic route to polymers PDTBTFV and PDTBTV.

After purification, PDTBTFV and PDTBTV were obtained as dark-violet and dark-blue powders, respectively, with good yields and good average molecular weights (yield = 75%, Mn = 43,700 and Mn/Mw = 4.46 for PDTBTFV, yield = 70%, Mn = 36,900 and Mn/Mw = 4.67 for PDTBTV). Both polymers are soluble in chlorinated solvents, such as chloroform, chlorobenzene and 1,2-dichlorobenzene.

2.1.2. Chemico-Physical Properties and Photovoltaic Performance in BHJ Solar Cells of PDTBTFV and PDTBTV: Impact of Fluoro-Functionalization of Vinylene Units

About the fluoro-functionalization, some papers demonstrated favourable effects of fluorine substitution on the performance of polymers used as donors in blend with...
PCBM derivatives in BHJ solar cells [44–47]. Fluorine as a substituent of conjugated materials is appropriate for lowering both the HOMO and LUMO energy levels; this can cause an increase of the open circuit voltage, Voc, in a donor material for BHJ. At the same time, the fluoro-functionalization increase the thermal and chemical stability of organic materials, matching the requirement of organic-based long-lifetime devices. In this direction, the insertion of fluorine atoms on the vinylene unit can be used as a strategic tool to increase the stability of double bonds to the photo-oxidation processes in arylenevinylene materials [28], affecting at the same time electronic properties both in solution [48] and solid state [26–28]. In the solid state, in fact, the C-F bonds promote π-stacking interactions [29,49–53]. Experimental and theoretical studies carried out on arylenevinylene polymers and oligomers fluorinated on the vinylene unit highlighted a strong impact of fluorine on the chemico-physical properties of this class of materials, showing a very large blue shift in absorption and emission (with respect to the non-fluorinated homologous), in both solution and solid state. All theoretical studies lead to assign the large blue shift to a remarkable molecular distortion induced by the repulsive interactions between fluorine atoms on the vinylene units and the oxygen atoms of alkoxy groups on the aromatic rings [26,27,29]. Furthermore, all investigations carried out on the arylenevinylene fluorinated on the double bond highlighted the ability of fluorine, as the substituent, to improve the stability of these materials against the photo-oxidation processes, as well as to deeply affect the electronic properties, disclosing it as one of the powerful tools to obtain blue emission from this class of materials, both in solution and solid state [28].

Polymers PDTBTFV and PDTBTV were used as donor materials blended with PCBM in BHJ solar cells and the photovoltaic performance were recorded on non-optimized devices [31]. Different donor/acceptor (polymer/PCBM) weight ratios were tested and the results of cell performance are reported in Table 1.

Table 1. Photovoltaic performance of BHJ solar cells of PDTBTFV and PDTBTV in blend with PCBM: open circuit voltage Voc; short circuit current Isc; fill factor FF and power conversion efficiency η.

| Polymer     | Polymer/PCBM Weight Ratio | Voc [V] | Isc [mA cm⁻²] | FF [%] | η [%] |
|-------------|---------------------------|---------|---------------|--------|-------|
| PDTBTFV     | 2:7                       | 0.73    | 2.94          | 38.57  | 0.83  |
|             | 3:7                       | 0.75    | 3.55          | 37.97  | 1.05  |
|             | 1:1                       | 0.83    | 4.32          | 34.56  | 1.24  |
| PDTBTV      | 2:7                       | 0.55    | 3.33          | 28.83  | 0.53  |
|             | 3:7                       | 0.56    | 2.42          | 24.50  | 0.33  |
|             | 1:1                       | 0.43    | 3.15          | 33.51  | 0.46  |

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Cell performance data shows the superiority of the fluorinated polymer PDTBTFV as a donor over the non-fluorinated PDTBTV, with higher values of Voc, FF and then η. For both polymers, the thermal annealing of the blend in the range 70–150 °C substantially does not affect the photovoltaic performance, according to the data of the differential scanning calorimetry (DSC), which shows the lack of any transition.

In order to shed light on the results achieved regarding the capability of fluorine to enhance the photovoltaic properties of the arylenevinylene structure investigated, we carried out theoretical, electrochemical and optical characterization. The absorption spectra of PDTBTFV and PDTBTV recorded in a chloroform solution are shown in Figure 2.
Both polymers absorb in a wide range of the visible region, with maxima picked at 548 nm (2.26 eV) for PDTBTFV and 569 nm (2.18 eV) for PDTBTV, with a blue-shift of about 21 nm (0.08 eV) induced by fluorination of vinylene units. According with the results of the TD-DFT calculations, the blue-shift is a result of the mesomeric and inductive effects of fluorine atoms. In the solid state, ellipsometry investigations showed the same trend in the absorption spectrum observed in solution, with an optical band gap of the films of polymers found at 1.79 eV for PDTBTFV and 1.69 eV for PDTBTV. The HOMO and LUMO energy levels were investigate by cyclic voltammetry, and based on the Koopmans’ theorem, they were estimated to be HOMO $-5.20$ eV for PDTBTFV and $-5.01$ for PDTBTV, LUMO $-3.14$ for PDTBTFV and $-3.03$ for PDTBTV. These data reveal an overall stabilization of both HOMO and LUMO energy levels induced by fluorination of vinylene units, in agreement with the results of DFT calculations. The ellipsometric investigations were very interesting, revealing a remarkable effect of fluoro-functionalization of the vinylene units inducing a significant enhancement of thin film light absorption in PDTBTFV with respect to PDTBTV. This result can be attributed to a different packing of the polymer chains in the solid state, with the fluorination of the vinylene units being responsible for the increased intermolecular interactions by π-stacking interactions and then for the observed increased absorption coefficient. Moreover, ellipsometry analysis carried out on the blend of polymers and PCBM used for the BHJ solar cells revealed that the absorption coefficient of the PDTBTFV-PCBM blend still remains higher than that of the PDTBTV-PCBM blend in the region of the main optical transition of the polymers (Figure 3).
Performance in Device

The higher capability of light absorption of the fluorinated polymer PDTBTFV might be a main reason for its better photovoltaic performance with respect to the non-fluorinated PDTBTV, since a higher light absorption has a greater chance of generating more charge carriers in the active blend. In summary, two main synergetic effects can be considered the basis of the favourable impact of fluorination of vinylene units on the photovoltaic performance of the polyarylenevinylene structure investigated: (a) the lowering of HOMO and LUMO energy levels, leading to an increase in Voc; (b) the increase in the molar extinction coefficient. The latter is very attractive in the perspective of reduction of film thickness in BHJ solar cells, which is necessary to enhance the charge carrier generation at the interface and enable the collection of carriers at the electrodes against their tendency to recombine.

2.2. Small Molecules for Dye-Sensitized Solar Cells DSSCs: From Synthesis to Photovoltaic Performance in Device

In the framework of the new generation of photovoltaics which can provide an alternative for the exploitation of solar energy, dye-sensitized solar cells (DSSCs) are hybrid photovoltaic devices with great potentiality, being inexpensive and environmentally friendly. Starting from the pioneering work of O’Regan and Grätzel in 1991 [54], they have been attracting a lot of interest as technically and economically credible alternative to the p-n junction solar cells. A DSSC is substantially a photoelectrochemical solar cells consisting of an organic dye sensitizing a mesoporous TiO$_2$ working electrode (WE), a redox electrolyte and a counter electrode (CE). Since both the WE and CE can be semi-transparent, both sides of the solar cell can be exploited for illumination. The efficiency of a DSSC is strictly connected with the nature of organic dye, WE, CE and electrolyte, and the interactions between all these components. The organic dye plays a key role in the performance and stability of the device, and the improvement of its performance is fundamental to gain advances in cell efficiency. Since high extinction coefficients are needed to gain high photovoltaic efficiencies, pure organic dyes are preferred with respect to traditional and first used metal-organic complexes [55–58]. Organic dyes, in fact, have unique advantages such as being low cost,
non-toxic, having a high extinction coefficient, a flexible molecular design, easy tunability of molecular structure and additionally of optical and electrochemical properties [59–62]. Despite these attractive features, drawbacks such as a low open circuit voltage and high recombination losses limit the efficiencies in the final device [63–65]. An ideal organic dye should possess features such as a high efficiency in device, chemical and thermal stability, and an easy and low cost of the synthetic procedure used for its preparation. The best performing organic dyes have a donor-π-bridge-acceptor (D-π-A) structural architecture, where A is mainly the cyanoacrylic group [66–71], while D and π are the units on which most of the efforts have been focused, in order to gain improvements of photovoltaic performances. As donor systems, the aryl amines resulted to be the best performing, and the research efforts have been devoted to the strengthening of the electron-donating power and to the introduction of bulky chains preventing the aggregation and recombination processes on the TiO₂. However, the π-bridge, linking the donor and acceptor units, plays a crucial role in the whole performance of the organic dye, by adjusting the HOMO and LUMO energy levels and extending the absorption range [60]. Moreover, since the π-bridge has to favour the electron migration from the donor to the acceptor, a planar geometry (fused rings) should be preferred. In fact, the literature reports, in general, high performances for organic dyes containing a fused ring as the π-bridge [72–78].

2.2.1. Benzothienobenzothiophene (BTBT)-Based Organic Dyes

In the framework of our interest in organic dyes for DSSCs, we focused our attention on a D-π-A molecular architecture based on an intriguing fused-ring planar system as the π-bridge, namely the [1]benzothieno[3,2-b]benzothiophene (BTBT). BTBT has been largely investigated for its excellent charge mobility in field-effect transistors [79–81], but not as a component of materials for photovoltaic applications. The charge-transport properties and the very easy and cheap synthetic access suitable to large scale production of BTBT make this molecular unit very attractive for photovoltaic purposes, especially in DSSC devices. We selected the cyanoacrylic group as the acceptor unit A, which was linked to the BTBT unit by the 3-methylthiophene ring (see Figure 4 for the units to connect for the synthesis of organic dyes).

![Figure 4. Units to be reacted to prepare the organic dyes.](image)

As donor D we selected three electron-donor groups, characterized by a different donating power and long alkyl chains (octyl chains) to prevent aggregation processes [82].

Synthesis of BTBT-Based Organic Dyes

The synthetic strategy carried out to prepare the three intermediates containing the donor unit D₁, D₂ and D₃ is shown in Scheme 4.
D1 was synthesized by reacting the fluorene derivatives 10 and 11 in the Buchwald-Hartwig reaction conditions. D2 was synthesized by reacting fluorenyl-boronate 12 with 2,3-dibromothiophene, in the Suzuki-Miyaura reaction conditions. Following the intermediate 14 was converted in the stannyl derivative D2 by lithiation and quenching with tributylstannylchloride. D3 was synthesized by reacting 12 with the phenothiazine derivative 13 in the Suzuki-Miyaura reaction conditions and following deprotection of the N-boc group.

The synthetic strategy carried out to prepare the organic dyes ODy1, ODy2 and ODy3 is detailed in Scheme 5.
Scheme 5. Synthetic route to organic dyes ODy1, ODy2 and ODy3.

In the first step, the dibromo-[1]benzothieno[3,2-b]benzothiophene (BTBT-Br$_2$) derivative was reacted with the three intermediates D1, D2 and D3 to yield the intermediate compounds 16, 17, 18. BTBT-Br$_2$ was coupled with D2 in the Stille reaction conditions, and was coupled with D1 and D3 in the Buchwald-Hartwig reaction conditions. The resulting intermediates 16, 17 and 18 were submitted to the Stille cross-coupling with the stannylthiophene derivative 19, to yield compounds 20, 21 and 22. Finally, compounds 20, 21 and 22 were reacted with cyanoacrylic acid to yield the final organic dyes ODy1, ODy2 and ODy3, which were obtained as red-orange, red and light-red powders, respectively.
Optical and Electrochemical Properties, Photovoltaic Performance of BTBT-Based Organic Dyes

Optical properties of the synthesized organic dyes ODy1, ODy2 and ODy3 were investigated by recording molar absorption spectra in chloroform solution (Figure 5).

All dyes show broad absorption spectra: ODy1 in the range 250–500 nm, with the maximum picked at 453 nm and an extinction coefficient of 14,190; ODy2 in the range 300–550 nm, with two main bands picked at 372 and 464 nm, and extinction coefficients over 60,000 and 42,000, respectively; ODy3 in the range 250–500 nm, with two main bands picked at 320 and 420 nm, and extinction coefficients over 100,000 and 69,000, respectively. As evident, the introduction of stronger electron-donor arylamine units in ODy1, strongly impacts the absorption capability of the dyes with a strong enhancement of the extinction coefficient. The increase in the absorption capability leads to an enhancement of photovoltaic performance (as reported below). Moreover, the introduction of a more complex donor system, based on the phenothiazine unit in ODy3, further increases the absorption capability of the related dye, beside a blue shift of the absorption spectrum. Nevertheless, the photovoltaic performance of ODy3 does not follow the strong improvement of the extinction coefficient, due to the blue shift of the absorption spectrum.

Cyclic voltammetry CV was performed to investigate the redox behaviour of the dyes and the HOMO and LUMO levels, together the energy gap Eg were estimated, as reported in Table 2.

Table 2. Electrochemical data, absorption maxima and molar extinction coefficients.

| Dye   | λmax (nm) a | ε (M⁻¹ cm⁻¹) b | HOMO (V) (vs. NHE) | LUMO (V) (vs. NHE) | Eg (eV) |
|-------|-------------|----------------|--------------------|-------------------|--------|
| ODy1  | 453         | 14,190         | 1.22               | −1.16             | 2.38   |
| ODy2  | 454/372     | 42,130/60,890  | 0.85               | −1.36             | 2.21   |
| ODy3  | 420/320     | 69,800/108,120 | 0.75               | −1.75             | 2.50   |

a absorption maximum of dyes in CHCl₃ with conc. ca. 2 μM. b Molar extinction coefficient at λmax. HOMO calculated by CV in CH₂Cl₂ with tetrabutylammonium hexafluorophosphate (0.1 M) as the supporting electrolyte, Ag/AgCl as the reference electrode, and Pt as the counter electrode. Scanning rate: 100 mV s⁻¹. LUMO was calculated as the difference HOMO—Eg. Eg was estimated from the absorption spectral onsets of the dyes.

Figure 5. Molar absorption spectra of organic dyes ODy1, ODy2 and ODy3 in CHCl₃. Reprinted (adapted) with permission from ref. [82]. Copyright [2016] American Chemical Society.
The HOMO energy levels of the dyes are more positive than the redox potential of I\(^-\)/I\(_3\)^- (0.4 V vs. NHE) [83], therefore the oxidized dyes obtained after electron injection into the conduction band of TiO\(_2\) could be regenerated by the reducing species in the electrolyte solution. The LUMO energy levels of the dyes are sufficiently more negative than the conduction band edge of TiO\(_2\) (−0.5 V vs. NHE), therefore there is a sufficient driving force for electron injection from the excited dyes to the conduction band of TiO\(_2\) [84]. These electrochemical data suggest these organic dyes could be suitable as sensitizer in DSSCs.

Since the optical and electrochemical properties of the organic dyes ODy1, ODy2 and ODy3 resulted suitable for photovoltaic applications, we tested them in DSSC solar cells and the results are shown in Table 3.

Table 3. Photovoltaic parameters for DSSCs based on ODy1, ODy2 and ODy3 dyes.

| Dye | CDCA (mM) | PCE (%) | Voc (V) | Jsc (mA/cm\(^2\)) | FF | Dye Loading (mol/cm\(^2\)) |
|-----|-----------|---------|---------|-----------------|----|--------------------------|
| ODy1 | 0         | 4.20    | 0.704   | 9.18            | 0.65 | 2.9 × 10\(^{-7}\) |
|     | 10        | 5.16    | 0.735   | 10.33           | 0.68 | 2.4 × 10\(^{-7}\) |
| ODy2 | 0         | 5.76    | 0.736   | 11.50           | 0.68 | 3.0 × 10\(^{-7}\) |
|     | 10        | 6.25    | 0.765   | 11.85           | 0.69 | 2.4 × 10\(^{-7}\) |
| ODy3 | 0         | 4.54    | 0.689   | 9.56            | 0.69 | 2.8 × 10\(^{-7}\) |
|     | 10        | 5.00    | 0.719   | 9.67            | 0.72 | 2.2 × 10\(^{-7}\) |

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Taking in account the high planarity of the BTBT system, which in turn favours the planarity of the dyes, unfavourable π−π stacking and fast intramolecular charge transfer should be expected, which may lead to intermolecular quenching or back transfer of the injected electrons from the TiO\(_2\) conduction band. In these cases, the addiction of chenoxycholic acid (CDCA) to the solution containing the dyes could be very useful. CDCA competing with the dye in binding the titania surface, can insert itself between the molecules of sensitizer, preventing or minimizing the detrimental aggregation effects [85–87]. In addition, the introduction of coadsorbents such as CDAC allows for a more uniform coverage of the inorganic semiconductor surface, hence lowering the probability of recombination processes between electrons injected and I\(_3\)^- or other acceptor species. Actually, the introduction of the coadsorbent CDCA favourably impacted on the photovoltaic performance, with an increase in Voc and Jsc parameters and then of efficiencies.

The organic dyes ODy1, ODy2 and ODy3 based on the BTBT π-bridge showed good photovoltaic parameters, with efficiencies in the range 4–6%. Although these values are not as high as other optimized structures [72–74,77,88] based on fused ring systems, they are comparable [76,89–92] and sometimes competitive [67,93–97] with other dyes. These achievements support the choice of the BTBT planar system as π-bridge with good potentiality in organic dyes for DSSCs.

2.2.2. Dibenzofulvene (DBF)-Based Organic Materials for Dye-Sensitized Solar Cells

DSSCs: Synthesis, Structure-Properties Correlation and Cell Performance

In the framework of our interest in organic small molecules for DSSC solar cells, especially for new π-bridges in D-π-A structures, we devoted our attention on the dibenzofulvene (DBF) system, having a planar configuration and whose chemical structure allows to connect two donor units per DBF molecule in a new structure of 2D-π-A type. In such a dye structure, two electron-donating units can push electrons to the electron-acceptor anchoring group. The DBF unit has been used as π-conjugated system in advanced functional materials for optoelectronic devices, such as organic field effect transistors (OFETs) [98], organic light emitting diodes (OLEDs) [99] and organic photovoltaics [100–102], but until our work, to the best of our knowledge, was lacking a systematic and in-depth investigation of its potential as a component of dyes for DSSC solar cells. By exploiting the electron delocalization along the ylidene system, the electron-donating effect of donor groups linked
to the DBF skeleton can be directed to the acceptor group, thus allowing the electron injection into the TiO$_2$ component of the cell. The position of donor groups onto the DBF skeleton, as well as their electron donating power can be exploited as chemical tools to tune the electronic properties of the organic dyes, and at the whole, their performance in cell. We selected diarylamines as donor groups, the cyanoacrylic acid as the acceptor group and the $\pi$-bridges consisting of DBF alone or DBF connected to thiophene or bithiophene systems [75,103–105]. The structures of the synthesized dyes are shown in Figure 6.

![Figure 6. Dibenzofulvene (DBF)-based organic dyes.](image)

**Synthesis of Dibenzofulvene (DBF)-Based Organic Dyes**

Organic dyes **TK1**, **TK2** and **TK3** were synthesized as reported in Scheme 6 [75].
To yield the intermediate compound 23, 3,6-dibromofluorenone 22 was reacted with diphenylamine in microwave-assisted Buchwald-Hartwig reaction conditions. Compound 23, when reacted with cyanoacrylic acid in the presence of TiCl₄ and pyridine yields the dye TK1, when submitted to the reduction of the carbonyl group by tert-butylamine-borane complex and aluminum chloride yields the intermediate 24. Compound 24 was reacted with thiophene-2,5-dicarbaldehyde and 2,2'-bithiophene-5,5'-dicarbaldehyde in a microwave reactor to yield intermediates 25 and 26, respectively. Finally, compounds 25 and 26 reacted with cyanoacrylic acid yield the dyes TK2 and TK3, respectively. The dyes TK1, TK2 and TK3 differ for the π-bridge, which is the DBF unit alone in TK1, the DBF-thiophene unit in TK2 and the DBF-bithiophene unit in TK3.

Organic dyes TK4, TK5 and TK6 were synthesized as reported in Scheme 7 [104].
Compound 22 was submitted to the reduction of the carbonyl group to yield intermediate 27, which in turn was reacted with compounds 28a and 28b to yield intermediates 29a and 29b, respectively (see reference [104] for details about the synthesis of compounds 28a, 28b, 30a and 30b). Compounds 29a and 29b were then reacted with 30 in the microwave-assisted Buchwald-Hartwig reaction conditions, to yield the intermediates 31a, 31b and 31c, respectively. Finally, 31a, 31b and 31c were reacted with cyanoacrylic acid to yield the organic dyes TK4, TK5 and TK6.

Organic dyes TK7, TK8 and TK9 were synthesized as reported in Scheme 8 [103].

Scheme 8. Synthetic route to dyes TK7, TK8 and TK9.

For details about the synthesis of intermediates 30a, 30b and 32 see reference [103]. Compound 22 was submitted to the reduction of the carbonyl group to yield intermediate 27, which was reacted with 32 to yield intermediate 33. Following, intermediate 33 was reacted with intermediates 30, 30a–b in the microwave-assisted Buchwald-Hartwig reaction conditions, to yield intermediates 34a–c. Finally, intermediates 34a–c were reacted with cyanoacrylic acid to yield the organic dyes TK7, TK8 and TK9.

Organic dyes TG1, TG2, TG3 and TG4 were synthesized as reported in Scheme 9 [105].

Scheme 9. Synthetic route to dyes TG1, TG2, TG3 and TG4.
Fluorene was reacted with intermediates 28a and 28b (see reference [105] for details about their synthesis) to yield intermediates 35a and 35b, which in turn were reacted with cyanoacrylic acid to yield the organic dyes TG1 and TG2. 2,7-dibromofluorene was reacted with intermediates 28a and 28b to yield intermediates 36a and 36b, which in turn were reacted with diphenylamine, in the microwave-assisted Buchwald-Hartwig reaction conditions, to yield intermediates 37a and 37b. Finally, compounds 37a and 37b were reacted with cyanoacrylic acid to yield the organic dyes TG3 and TG4.

**Electro-Optical Properties, Structure-Properties Correlation and Photovoltaic Performance of Dibenzofulvene (DBF)-Based Organic Dyes**

In view of design and synthesis of DBF-based organic dyes with structure 2D-π-A for DSSC, the DBF molecule is a versatile chemical structure which can be easy functionalized in the 2,7- and 3,6-positions by electron-donor groups. This allows for the design of two different classes of dyes, with different structural and chemico-physical behaviour, in which a different electronic conjugation is expected between the donor and the acceptor units, the last being connected to the donor by the ylidene system and the π-spacer of the π-bridge. In line with these assessments, we studied the effects of donor position on DBF-based organic dyes for DSSC [105], starting our investigation from structurally simply dyes, shown in Figure 7.

**Figure 7.** Dibenzofulvene (DBF)-based organic-dyes.

TG1 and TG2 are two basic structures without donor groups on the DBF unit, and containing one or two thiophene rings, respectively, as spacer between the ylidene system and the cyanoacrylic acceptor unit. TG3 and TG4 can be considered as deriving from TG1 and TG2, respectively, with the introduction of the diphenylamine donor unit in the 2,7-positions of the DBF structure. TK2 and TK3, instead, can be considered as deriving from TG1 and TG2, respectively, with the introduction of diphenylamine donor unit in the 3,6-positions of DBF structure. The absorption spectra of dyes recorded in CH$_2$Cl$_2$ are reported in Figure 8 as molar absorption coefficient $\varepsilon$. 
From the analysis of absorption features (absorption spectra in solvent with different polarity, such as DMSO and chlorobenzene, were also recorded), some interesting behaviours can be observed, strictly depending on the position of the donor groups on the DBF structure. In fact, going from TG1 to TG3 (as well as from TG2 to TG4) the lowest-energy absorption peak undergoes a slight perturbation, since the excitation in both dyes is associated to a single particle transition between orbitals essentially localized on the DBF core and the \( \pi \)-conjugated bridge. Moreover, in TG3 and TG4 an additional peak appears at about 350 nm, which can be attributed to \( \pi-\pi^* \) transitions localized on the diphenylamine and the DBF core, as suggested by TD-DFT calculations and the low influence of solvent polarity. When comparing TG1/TG3 and TG2/TG4 with TK2 and TK3, respectively, the situation is different. TK2 and TK3 differ from TG2 and TG4 for the position of diphenylamino groups on the DBF structure. For TK2 and TK3 the lowest-energy peak has a marked charge-transfer character, with the HOMO localized mainly on the diphenylamino substituents and the LUMO localized mainly on the cyanoacrylic group and the linker. Therefore, this peak is red shifted with respect to that of TG1/TG3 and TG2/TG4. Furthermore, the peak at about 350 nm results reduced in TK2 and TK3, with respect to TG3 and TG4, since the excitation associated to a single particle transition is no more localized only on the \( \pi \) system of diphenylamines and DBF. These optical features are a proof of a better coupling between diphenylamines and DBF core for 3,6-functionalized dyes with respect to 2,7-functionalized ones, so that the electron donating capacity of donors is enhanced as well as the electron delocalization in TK2 and TK3 with respect to TG3 and TG4. Moreover, the effect of the bridge length on the absorption properties of dyes also turns out to be noteworthy. Comparing the spectra of TG1, TG2, TG3 and TG4, a red shift in the absorption is observed with the introduction of an additional thioephene ring in the bridge, so that the effective conjugation length is increased in TG2 and TG4 with respect to TG1 and TG3. Conversely, TK2 and TK3 dyes showed opposite behaviour, since despite a longer conjugated bridge with two thienyl rings, the latter absorbs at higher energies compared to the former. This is because the effective conjugation length is mainly affected by the functionalization in 3,6-positions, while an additional thiophene ring substantially does not impact on the global effective conjugation, but on the contrary, it appears to negatively affect it. HOMO and LUMO energy levels of dyes were estimated by cyclic voltammetry and the results highlight they are suitable to be used as sensitizers in DSSC.
The results suggest the functionalization of 3,6-positions of DBF molecule as more favorable to achieve higher photovoltaic performance in this class of organic dyes. In order to pull out the best photovoltaic potential from this class of DBF-based dyes, our investigation continued focusing the attention on three structures, labelled as TK1, TK2 and TK3, characterized by diphenylamine donor groups linked to the 3,6-positions of DBF core and different for the number of thiophene rings between the DBF and the electron-acceptor group (cyanoacrylic): no thiophene ring in TK1, one thiophene ring in TK2 and two thiophene rings in TK3. The absorption spectra of such dyes in CH$_2$Cl$_2$ solution are shown in Figure 9.

![Figure 9](image-url)

**Figure 9.** Absorption spectra of TK1, TK2 and TK3 in CH$_2$Cl$_2$ solution. Reprinted (adapted) with permission from ref. [75]. Copyright [2014] “The Royal Society of Chemistry”.

TK1 shows the maximum absorption peak at 449 nm ($\varepsilon = 7.3 \times 10^3$). The introduction of one thiophene ring as a spacer between the DBF structure and the cyanoacrylic acceptor unit yields the dye TK2, which shows a bathochromic shift of 73 nm of the absorption maximum and an increase in the molar extinction coefficient ($\varepsilon = 1.2 \times 10^4$). The introduction

Table 4. Photovoltaic parameters for DSSCs based on dyes TG1–4 and TK2–3.

| Dye   | $\eta$ (%) | Voc (V) | Jsc (mA/cm$^2$) | FF  |
|-------|------------|---------|-----------------|-----|
| TG1   | 1.23       | 0.56    | 3.09            | 0.71|
| TG2   | 2.07       | 0.61    | 5.22            | 0.65|
| TG3   | 3.91       | 0.73    | 7.50            | 0.71|
| TG4   | 2.68       | 0.65    | 5.37            | 0.77|
| TK2   | 5.01       | 0.66    | 10.85           | 0.70|
| TK3   | 5.42       | 0.67    | 11.55           | 0.70|

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The photovoltaic performance of dyes is consistent with the optical and electrochemical properties. TG1 and TG2 show very poor efficiencies (1.23% and 2.07%, respectively), as expected, due to the lack of donor groups. TG3 and TG4 show enhanced efficiencies, with values of 3.91% and 2.68%, respectively, thanks to the introduction of donor groups in 3,6-positions of DBF molecule as more favourable to achieve higher photovoltaic performance in this class of organic dyes.

The absorption spectra of such dyes in CH$_2$Cl$_2$ solution are shown in Figure 9.
of two thiophene rings yields the dye TK3, which shows a lower bathochromic shift of 36 nm of the absorption maximum and a higher increase in the molar extinction coefficient \( \varepsilon = 1.7 \times 10^4 \). HOMO and LUMO energy levels were estimated by cyclic voltammetry and the data are reported in Table 5, together with those of absorption.

Table 5. Absorption and electrochemical data of TK1, TK2 and TK3.

| Dye   | \( \lambda_{\text{abs}} \) (nm) | \( \varepsilon \times 10^3 \) (M\(^{-1}\) cm\(^{-1}\)) | HOMO/LUMO (eV) | \( E_{0.0} \) b | \( E_{\text{ox}} \) c (V) |
|-------|-------------------------------|---------------------------------|----------------|--------------|----------------|
| TK1   | 449                           | 7.3                             | \(-5.35/\sim3.17\) | 2.18         | 1.29           |
| TK2   | 522                           | 12                              | \(-5.08/\sim3.13\) | 1.96         | 1.03           |
| TK3   | 485                           | 17                              | \(-4.99/\sim2.91\) | 2.08         | 0.94           |

\( \lambda_{\text{abs}} \) Absorption maximum measured in CH\(_2\)Cl\(_2\). b Bandgap estimated from the onset of absorption spectrum. c Oxidation potential in CH\(_2\)Cl\(_2\) solution containing 0.1 M of NBu\(_4\)PF\(_6\) with a scan rate of 100 mV s\(^{-1}\) (vs. NHE) and calibrated against ferrocene. Reprinted (adapted) with permission from ref. [75]. Copyright (2014) “The Royal Society of Chemistry”.

The absorption and electrochemical data indicate the dyes are suitable for application as sensitzers in DSSC solar cells; then, they were used to fabricate photovoltaic devices, in order to investigate their performance. Chenodeoxylic acid was also used as a co-adsorbent to prevent recombination phenomena which lead to the detriment of cell efficiency. The photovoltaic parameters are summarized in Table 6.

Table 6. Photovoltaic parameters for DSSCs based on dyes TK1, TK2 and TK3.

| Dye   | CDCA (mM) | \( \eta \) (%) | FF | Voc (V) | Jsc (mA cm\(^{-2}\)) | Dye Loading (10\(^{-7}\) mol cm\(^{-2}\)) |
|-------|-----------|----------------|----|---------|-----------------------|------------------------------------------|
| TK1   | 0         | 2.14           | 0.75 | 0.63    | 4.52                  | 2.1                                       |
|       | 10        | 1.08           | 0.70 | 0.70    | 2.21                  | 1.5                                       |
| TK2   | 0         | 5.01           | 0.70 | 0.66    | 10.85                 | 2.5                                       |
|       | 10        | 4.72           | 0.73 | 0.73    | 8.85                  | 1.8                                       |
| TK3   | 0         | 5.42           | 0.70 | 0.67    | 11.55                 | 2.6                                       |
|       | 10        | 7.45           | 0.71 | 0.70    | 14.98                 | 2.0                                       |
| N719  | 0         | 8.11           | 0.70 | 0.79    | 14.68                 | 1.7                                       |

\( \text{CDCA} = \) chenodeoxycholic acid. Reprinted (adapted) with permission from ref. [75]. Copyright (2014) "The Royal Society of Chemistry.

When switching from TK1 to TK2 to TK3 the power conversion efficiency increases from 2.1% to 5.0% to 5.4%, respectively. This is the result of an increase mainly of Jsc, which goes from 4.52 mA cm\(^{-2}\) (TK1) to 10.82 mA cm\(^{-2}\) (TK2) to 11.55 mA cm\(^{-2}\) (TK3), respectively. The IPCE spectra partially explains what happens: the DSSC based on TK1 shows a lower photoresponse with a maximum of about 20% at 430 nm, compared with the value of about 50% in the region 450–550 nm obtained with the DSSCs based on TK2 and TK3, according with the absorption spectra. Moreover, regarding the dye loading, TK1 shows a lower adsorption amount compared to TK2 and TK3, probably due to the bulkier molecular structure (the addition of CDCA leads to lower efficiencies), providing a reason for the lower efficiency, assigned to the increased probability of detrimental charge recombination processes. So, the addition of CDCA causes a detrimental effect on the photovoltaic efficiency. To more deeply investigate the relationship between molecular structure and photovoltaic performance, the capacitance, the charge transfer resistance at the TiO\(_2\)/dye/electrolyte interface (Rct) and electron lifetime were measured by electrochemical impedance spectroscopy. For the TK1-based DSSC a lower Rct was found, indicating the injected electrons easily undergo to recombination processes. The introduction of one or two thiophene rings in the structure of the dyes (TK2 and TK3, respectively) allows to extend the \( \pi \)-bridge, favouring an increase in the interfacial resistance (also confirmed by the higher Voc value), suggesting a larger number of photo-generate electrons can reach the contact and justifying the higher photocurrent. Furthermore, TK1 shows a
shorter apparent electron lifetime, to confirm an easier recombination of injected electron from TK1. Taking into account the planarity of dyes TK1−3, favoured by the planar core of DBF, a fast intramolecular charge transfer as well as an unfavourable π−π stacking are expected, which may lead to recombination processes. So, the use of CDAC as a coadsorbent is expected to be useful to improve the cell performance, since it can be able to minimize the detrimental dye aggregation by competing with the dye in binding the TiO2 surface. DSSCs built by using CDCA as a co-adsorbent yielded 1.1%, 4.72% and 7.45% efficiencies for TK1, TK2 and TK3 dyes, respectively, highlighting a marked improvement for TK3. In the case of TK1, the addition of CDAC lowers the efficiency of device, since the only effect of the coadsorbent results in an unfavourable competition with the sensitizer in binding the TiO2 surface, without the benefit of minimizing the dye aggregation, since the high steric hindrance of TK1 directly acts to prevent aggregation processes. The results achieved in this work [75] highlights simple modifications in the molecular structure of a dye, such as the introduction of one or two thiophene rings, can have a strong impact on the chemico-physical properties and on the performance of photovoltaic devices, improving the efficiency from 2.14% to 7.45%.

Continuing further with our interest in deepening the knowledge of the correlation between structure and chemico-physical properties, aimed to improving the photovoltaic performance of DBF-based dyes, we focused our attention on the enhancement of the electron donor power of the diphenylamine units used as donor systems and on the modulation of the π-bridge of the 2D-π-A structure. For this purpose, we synthesized the dyes TK4, TK5 and TK6, which can be considered as deriving from the best performing dyes TK2 and TK3, by introduction of para-alkoxy substituted diphenylamines as donor units. The strategy was devoted to structures to pursue characteristics such as high electron-donating ability, reduced dye aggregation and charge recombination, aimed to enhance photovoltaic performances.

Photophysical properties were investigated by recording absorption spectra of TK4, TK5 and TK6 dyes in CH2Cl2 solution, as reported in Figure 10.

![Figure 10](image-url)  
**Figure 10.** Absorption spectra of TK4, TK5 and TK6 in CH2Cl2. Reprinted (adapted) with permission from ref. [104]. Copyright [2016] “Elsevier”.

All dyes show strong absorption bands in the ranges 260–350 nm and 350–600 nm, where the higher energy bands are attributed to local π−π* transitions, while the lower energy ones are attributed to charge transfer (CT) transitions. TK4 shows two maxima peaks at 360 and 548 nm, with extinction coefficients ϵ = 13,200 and 10,100, respectively.
TK5, which differs from TK4 only for the presence of two thiophene rings, instead of one, between the DBF core and the acceptor unit, shows two maxima peaks at 413 and 513 nm, with extinction coefficients $\varepsilon = 24,800$ and 17,800, respectively. TK6, which differs from TK5 for the octyloxy groups, instead of the methoxy groups on the diphenylamine units, shows two maxima peaks at 420 and 516 nm, with extinction coefficient $\varepsilon = 34,500$ and 25,100, respectively. The HOMO energy level was calculated by cyclic voltammetry and the LUMO energy level was derived by the formula $E_{\text{HOMO}} - E_{0-0}$, where the last is the optical band gap estimated from the absorption edge. The data are reported in Table 7.

Table 7. Absorption and electrochemical data of TK4, TK5 and TK6.

| Dye  | $\lambda_{\text{abs}}$ ($\varepsilon \times 10^3$ M$^{-1}$ cm$^{-1}$) a (nm) | HOMO b | $E_{0-0}$ c | LUMO d |
|------|-------------------------------------------------|--------|-------------|--------|
| TK4  | 548 (10.1)                                      | 0.64   | 1.82        | −1.18  |
| TK5  | 513 (17.8)                                      | 0.68   | 1.90        | −1.22  |
| TK6  | 516 (25.1)                                      | 0.73   | 1.88        | −1.15  |

a Absorption maximum in CH$_2$Cl$_2$ (10$^{-5}$ M) solution. b HOMO (vs. NHE) of the dyes by cyclic voltammetry in CH$_2$Cl$_2$ solution containing 0.1 M of tetrabutylammonium hexafluorophosphate, with a scan rate of 100 mV$^{-1}$ and calibrated against ferrocene. c The bandgap, $E_{0-0}$ was determined from the onset of absorption spectrum. LUMO = HOMO $- E_{0-0}$. Reprinted (adapted) with permission from ref. [104]. Copyright (2016) “Elsevier”.

The electrochemical data allow us to consider the dyes suitable to be used as sensitizers in DSSCs, so they were tested in liquid cells, and their photovoltaic parameters and performances are reported in Table 8.

Table 8. Photovoltaic performance of TK4, TK5 and TK6.

| Dye  | $\eta$ (%) | FF | Voc | Jsc | Dye Loading a |
|------|------------|----|-----|-----|---------------|
| TK4  | 5.9        | 0.67 | 0.667 | 13.29 | 2.0           |
| TK5  | 7.5        | 0.64 | 0.653 | 17.85 | 1.9           |
| TK6  | 7.8        | 0.69 | 0.663 | 17.19 | 1.6           |

a [1 × 10$^{-7}$ mol·cm$^{-2}$]. Reprinted (adapted) with permission from ref. [104]. Copyright (2016) Elsevier.

Different solvents were tested, reaching the best performance using THF for TK5 and TK6, and CH$_3$CN/CHCl$_3$ (1/0.01 v/v) for TK4. Different amounts of CDCA were also tested, reaching the best results with a concentration 15 mM for TK4 and TK5 and 5 mM for TK6. The reduced amount of CDCA needed for TK6 is consistent with the structure of such a dye, whose long octyl chains, as expected, act as hindrance to the aggregation of the molecules, lowering the amount of CDCA needed to gain the best photovoltaic performance. Starting from TK4, whose maximum efficiency was found to be 5.9%, the addition of a thiyl ring in the $\pi$-bridge of the molecular structure leads to TK5, which shows a significant improvement of performance with an efficiency of 7.5%. The additional structural variation by long octyloxy chains in TK6 results in a further increase in photovoltaic performance, reaching a maximum efficiency of 7.8%, together a reduced amount of CDCA needed in the device. The enhancement of photovoltaic efficiencies is mainly the result of an increase in the circuit current $J_{\text{sc}}$, which value goes from 13.29 mA/cm$^2$ for TK4 to 17.85 and 17.19 mA/cm$^2$ for TK5 and TK6, respectively. The behaviour of dyes is confirmed also by the IPCE spectra, in which the TK4-based device shows the lower photoresponse, with a maximum of about 55% at 450–550 nm, compared to the plateau of about 70% and 75% in the same region obtained with TK5- and TK6-based devices, respectively. Interestingly, comparing the photovoltaic parameters of the dyes TK4 and TK5 with those of their analogous without methoxy-groups TK2 and TK3, the improvement of the optical properties of the new dyes positively affects the efficiency of the devices. The introduction of the methoxy groups determines a red shift and a widening of the absorption spectra of TK4 and TK5, so their IPCE curves result higher and extended as well as higher result their photocurrent densities. In fact, the Jsc value goes from 10.85 and 14.98 mA/cm$^2$ for TK2 and TK3, to 13.29 and 17.85 mA/cm$^2$ for TK4
and TK5, respectively, probably for the enhanced light harvesting capability. The results highlight as a simple structural variation consisting in the introduction of alkoxy groups onto the donor units, providing a fine synthetic tool to modulate photovoltaic parameters and enhance the efficiency of device.

During our ongoing investigations, we finally became interested in studying the effects of a combined functionalization by alkoxy groups onto the donor units and alkyl chains onto the π-bridge, on the optical and photovoltaic properties of DBF-based organic dyes for DSSC. In detail, we focused our attention on the dye TK3, one of the best performing in terms of photovoltaic efficiency, using it as starting molecular structure to get new derivatives by functionalization with methoxy or octyloxy groups of the diphenylamine donor units, and with n-hexyl chains of one of the two thienyl rings connecting the DBF core to the acceptor unit. Three new dyes were then obtained, TK7, TK8 and TK9, where TK7 can be considered as deriving from TK3 by a simple introduction of a n-hexyl chain on the thienyl ring closer the acceptor unit, TK8 as a further evolution of TK7 by the introduction of methoxy chains in the para position of diphenylamines, and TK9 as a further evolution of TK7 by the introduction of octyloxy chains in the para positions of diphenylamines.

The optical properties of TK7, TK8 and TK9 were investigated by recording absorption spectra in CH2Cl2 solution (Figure 11); electrochemical properties were recovered by cyclic voltammetry and the data were collected in Table 9.

![Figure 11. Absorption spectra of TK7, TK8 and TK9 in CH2Cl2. Reprinted (adapted) with permission from ref. [103]. Copyright (2016) Elsevier.](image)

| Dye | \( \lambda_{\text{max}} \) (nm) \( \times 10^3/M^{-1} \text{cm}^{-1} \) | HOMO (V) vs. NHE | LUMO (V) vs. NHE | \( E_{0-0} \) (eV) |
|-----|------------------|------------------|------------------|---------------|
| TK7 | 422 (33.9)        | 0.98             | -1.04            | 2.02          |
| TK8 | 416 (30.6)        | 0.94             | -1.04            | 1.98          |
| TK9 | 428 (17.5)        | 0.93             | -1.07            | 2.00          |

\( a \) Absorption maximum in CH2Cl2 (10^{-5} M) solution. \( b \) HOMO of dyes by cyclic voltammetry in 0.1 M TBAPE6 in CH2Cl2 solutions as supporting electrolyte, Ag/AgCl as the reference electrode, and Pt as the counter electrode, and calibrated against ferrocene. Scanning rate: 100 mV s^{-1}. \( c \) LUMO was calculated by HOMO-\( E_{0-0} \). \( d \) \( E_{0-0} \) was estimated from the onset of the absorption spectrum. Reprinted (adapted) with permission from ref. [103]. Copyright (2016) Elsevier.
The dyes show two bands, one at longer wavelengths, in the range 360–600 nm, which is mainly attributed to the charge transfer (CT) from diarylamine donors to cyanoacrylic acid acceptor, and the other in the region 270–350 nm, originating from DBF and thiophene units, which can be attributed to $\pi-\pi^*$ transitions.

$TK7$ shows two absorption maxima at 304 and 422 nm, with extinction coefficient $\varepsilon = 49,000$ and 33,900, respectively. $TK8$ and $TK9$, characterized by methoxy and octyloxy groups, respectively, on the diphenylamines, show a reduced absorption capability with respect to $TK7$. $TK8$ shows two absorption maxima at 293 and 416 nm, with extinction coefficient $\varepsilon = 42,500$ and 30,600, respectively. $TK9$ shows two absorption maxima at 270 and 428 nm, with extinction coefficient $\varepsilon = 45,180$ and 17,500, respectively. By cyclic voltammetry was estimated the HOMO energy level of $TK7$, $TK8$ and $TK9$, which was found to be 0.98, 0.94 and 0.93 V, respectively, while the energy gap ($E_{0-0}$) was estimated by the onset wavelength of the absorption spectra, and found to be 2.02, 1.98 and 2.00 eV, respectively. The LUMO energy level was then calculated as the difference HOMO $- E_{0-0}$ and found to be $-1.04$, $-1.04$ and $-1.07$ V (vs NHE), respectively. The optical and electrochemical data prove the dyes are suitable as sensitizer in DSSCs, so their photovoltaic potential was tested in such devices, and the parameters are collected in Table 10.

Table 10. Photovoltaic parameters of $TK7$, $TK8$ and $TK9$ in DSSC devices.

| Dye | $\eta$ % | FF | Voc (V) | $J_{sc}$ (mA/cm$^2$) | Dye Loading (10$^{-7}$ mol/cm$^2$) |
|-----|----------|----|---------|----------------------|-------------------------------------|
| $TK7$ | 7.88     | 0.66 | 0.67    | 17.82                | 2.3                                 |
| $TK8$ | 6.35     | 0.67 | 0.66    | 14.33                | 2.1                                 |
| $TK9$ | 6.14     | 0.67 | 0.68    | 13.48                | 1.7                                 |

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$TK7$ shows the higher efficiency, 7.88%, mainly due to the higher photocurrent density ($J_{sc}$ 17.82 mA/cm$^2$). $TK8$ and $TK9$ show efficiencies of 6.35% and 6.14%, respectively. The higher value of $J_{sc}$ in the $TK7$-based device is related to its higher molar extinction coefficient. The IPCEs spectra recorded for the DSSCs are substantially in accordance with the optical properties of the dyes and in agreement with the higher light-harvesting capability of $TK7$, whose spectrum shows a plateau at ~80% between 400 and 550 nm, considerably wider with respect to the spectrum observed for $TK8$ and higher with respect to the response obtained for $TK9$. Correlating the optical and photovoltaic properties with the structural characteristics, the role of the hexyl chain inserted on the thienyl ring of the $\pi$-bridge can be highlighted by comparing the photovoltaic performance of $TK7$ and its homologous without the hexyl chain $TK3$, where the maximum efficiency in the former ($\eta$ 7.88%) is reached with a lower amount of co-adsorbent CDCA (5 mM), with respect to the latter ($\eta$ 7.45%, CDCA 10 mM). In the comparison between dyes, results show that the alkoxy groups (in $TK8$ and $TK9$) on the diphenylamine donor units negatively impact photovoltaic performance, suggesting that in such a branched dye structure, although they space the molecules preventing undesirable dye aggregation, on the other side they can lead to a less-uniform coverage of the TiO$_2$ surface.

3. Conclusions

In the present review, we have reported some of our contributions in the field of functional organic materials for photovoltaic applications. We have highlighted the synthetic aspects inherent to the synthetic methodologies exploited for their preparation, the photophysical properties and the performance in photovoltaic devices. Organometallic methodologies were mainly used as synthetic protocols, which allow for a rigorous stereocontrol of reaction, then of the stereochemistry of the final products. These protocols are characterized by mild reaction conditions and tolerate a wide variety of functional groups, resulting in key routes to access to a widespread variety of organic semiconductor materials. Afterwards, we have described the optical and electrochemical properties of
the synthesized materials and their performance in a device. Finally, we investigated the correlation between chemical structure and properties, highlighting the conjugated structure and the functionalization, understood as a type and position of a functional group onto the conjugated skeleton, which can be used as key tools to finely tune properties, both at a molecular level and in solid state, and then, performance in technological applications. Today, power conversion efficiency, stability and cost are the main open challenges in the OPV technology. In recent years, tremendous strides have been made in all three fields, and it seems reasonable to think the progress will continue in the near future. The development of new materials and the improvement of the understanding of the key physical processes governing the device operation are the two areas where the progress has been most significant. Power conversion efficiency near 20% has been reached in the best OPVs and it is reasonable to expect improvements for the years to come, taking into account the infinite number of potential materials offering endless possibilities and the enhancement in device building. On the other hand, machine learning is a field of computer science with high potential and is very promising for organic photovoltaics research [106,107]. Machine learning methods exploiting their data-analysis capability have the potential to guide researchers to discover and develop new high-performance materials [108,109]. Since the OPV technology is form-free, compatible with large-scale printing methods, requires low-cost production investments, is economically and environmentally sustainable, and although the performance of device is not still comparable with the inorganic counterparts, its implementation in “niche” applications such as building integrated environments or wearable electronics can be very competitive and commercially attractive. Furthermore, the Internet of Things (IoT) is a rapidly developing sector that requires cost-effective, low-energy and maintenance-free power sources for use in offices, human wellness and retail. Photovoltaic technologies, especially OPVs, are interesting energy-harvesting systems for IoT, thanks to their potential for low-power output and portability in miniaturized and grid-independent applications. The OPVs have the potential to be one piece of the puzzle of the future’s energy landscape.

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References
1. Facchetti, A.; Marks, T.J.; Katz, H.E.; Veinot, J. Organic Semiconductor Materials. In Printed Organic and Molecular Electronics; Gamota, D., Brazis, P., Kalyanasundaram, K., Zhang, J., Eds.; Springer: Boston, MA, USA, 2004; pp. 83–159.
2. Bronstein, H.; Nielsen, C.B.; Schroeder, B.C.; McCulloch, I. The role of chemical design in the performance of organic semiconductors. Nat. Rev. Chem. 2020, 4, 66–77. [CrossRef]
3. Oksana, O. Handbook of Organic Materials for Electronic and Photonic Devices, 2nd ed.; Woodhead Publishing: Sawston, UK; Elsevier: Amsterdam, The Netherlands, 2019; pp. 873–891.
4. Zou, S.-J.; Shen, Y.; Xie, F.-M.; Chen, J.-D.; Li, Y.-Q.; Tang, J.-X. Recent advances in organic light-emitting diodes: Toward smart lighting and displays. Mater. Chem. Front. 2020, 4, 788–820. [CrossRef]
5. Inganas, O. Organic Photovoltaics over Three Decades. Adv. Mater. 2018, 30, 1800388. [CrossRef]
6. Wang, X.; Sun, Q.; Gao, J.; Wang, J.; Xu, C.; Ma, X.; Zhang, F. Recent Progress of Organic Photovoltaics with Efficiency over 17%. Energies 2021, 14, 4200. [CrossRef]
7. Iqbal, J.; Enevold, J.; Larsen, C.; Wang, J.; Revoju, S.; Barzegar, R.H.; Wagberg, T.; Eliasson, B.; Edman, L. An Arylene-vinylene Based Donor-Acceptor-Donor Small Molecule for the Donor Compound in High-Voltage Organic Solar Cells. Sol. Energy Mater. Sol. Cells 2022, 155, 348–355. [CrossRef]
8. Romero, N.A.; Nicewicz, D.A. Organic Photoredox Catalysis. Chem. Rev. 2016, 17, 10075–10166. [CrossRef] [PubMed]
9. Melchionna, M.; Fornasiero, P. Updates on the Roadmap for Photocatalysis. ACS Catal. 2020, 10, 5493–5501. [CrossRef]
37. Chen, S.-A.; Chang, E.-C. Structure and properties of cyano-substituted poly(2,5-dialkoxy-p-phenylene vinylene). *Macromolecules* 1998, 31, 4899–4907. [CrossRef]
38. Jin, Y.; Ju, J.; Kim, J.; Lee, S.; Kim, J.Y.; Park, S.H.; Sun, S.-M.; Jin, S.-H.; Lee, K.; Suh, H. Design, synthesis and electroluminescent property of CN-poly(dihexylfluorenevinylene) for LEDs. *Macromolecules* 2003, 36, 6970–6975. [CrossRef]
39. Glich, H.G.; Wheelwright, W.L. Polymerization of -halogenated p-xylenes with base. *J. Polym. Sci. A Polym. Chem.* 1966, 4, 1337–1349. [CrossRef]
40. Wessling, R.A. The polymerization of xylylene bisdiaryl sulphonium salts. *J. Polym. Sci. Polym. Symp.* 1985, 72, 55–66. [CrossRef]
41. Babudri, F.; Ciccio, S.R.; Farinola, G.; Naso, F. Synthesis, characterization and properties of a soluble polymer with a poly(phenylenevinylene) structure. *Macromol. Chem. Commun.* 1996, 17, 905–911. [CrossRef]
42. Babudri, F.; Cardone, A.; Chiavarone, L.; Ciccarella, G.; Farinola, G.M.; Naso, F.; Scamarcio, G. Synthesis and characterization of poly(2,3,5,6-tetrafluoro-4,4-phenylenevinylene). *Chem. Commun.* 2001, 1940–1941. [CrossRef]
43. Babudri, F.; Cardone, A.; Farinola, G.M.; Naso, F.; Cassano, T.; Chiavarone, L.; Tommasi, R. Synthesis and optical properties of a copolymer of tetrafluoro- and dialkoxy-substituted poly(p-phenylenevinylene) with a high percentage of fluorinated units. *Macromol. Chem. Phys.* 2003, 204, 1621–1627. [CrossRef]
44. Liang, Y.; Feng, D.; Wu, Y.; Tsai, S.-T.; Li, G.; Ray, C.; Yu, L. Highly efficient solar cell polymers developed via fine-tuning of structural and electronic properties. *J. Am. Chem. Soc.* 2009, 131, 7792–7799. [CrossRef] [PubMed]
45. Liang, Y.; Yu, L. A new class of semiconducting polymers for bulk heterojunction solar cells with exceptionally high performance. *MACRO* 2022, 15, 6333.
46. Price, S.C.; Steuart, A.C.; Yang, L.; Zhou, H.; You, W. Fluorine substituted conjugated polymer of medium band gap yields 7% efficiency in polymer-fullerene solar cells. *J. Am. Chem. Soc.* 2011, 133, 4625–4631. [CrossRef] [PubMed]
47. Schroeder, B.C.; Huang, Z.G.; Ashraf, R.S.; Smith, J.; D’Angelo, P.; Watkins, S.E.; Anthopoulos, T.D.; Durrant, J.R.; McCulloch, I. Silaindacenodithiophene-based low band gap polymers-the effect of fluorine substitution on device performance and film morphologies. *Adv. Funct. Mater.* 2012, 22, 1663–1670. [CrossRef]
48. Facchetti, A.; Yoon, M.H.; Stern, C.; Katz, H.E.; Marks, T.J. Building blocks for n-type organic electronics: Regiochemically modulated inversion of majority carrier sign in perfluoroarene-modified polythiophene semiconductors. *Angew. Chem. Int. Ed.* 2003, 42, 3900–3903. [CrossRef]
49. Curti, M.D.; Cao, J.; Kempf, J.W. Solid-state packing of conjugated oligomers: From -stacks to the herringbone structure. *J. Am. Chem. Soc.* 2004, 126, 4318–4328. [CrossRef]
50. Reichenbacher, K.; Suss, H.I.; Hullinger, J. Fluorine in crystals engineering-“the little atom that could”. *Chem. Soc. Rev.* 2005, 34, 22–30. [CrossRef]
51. Gierschner, J.; Ehni, M.; Egelhaaf, H.-J.; Mili-Medina, B.; Beljonne, D.; Bennounsour, H.; Bazan, G.C. Solid-state optical properties of linear polyconjugated molecules: -stack contra herringbone. *J. Chem. Phys.* 2005, 123, 144914. [CrossRef]
52. Wang, Y.; Parkin, S.R.; Gierschner, J.; Watson, M.D. Highly fluorinated benzobisbenzothiophenes. *Org. Lett.* 2008, 10, 3307–3310. [CrossRef] [PubMed]
53. Facchetti, A.; Yoon, M.H.; Stern, C.L.; Katz, H.E.; Marks, T.J. Building blocks for n-type organic electronics: Regiochemically modulated inversion of majority carrier sign in perfluoroarene-modified polythiophene semiconductors. *Angew. Chem. Int. Ed.* 2003, 42, 3900–3903. [CrossRef]
54. Curti, M.D.; Cao, J.; Kempf, J.W. Solid-state packing of conjugated oligomers: From -stacks to the herringbone structure. *J. Am. Chem. Soc.* 2004, 126, 4318–4328. [CrossRef]
55. Reichenbacher, K.; Suss, H.I.; Hullinger, J. Fluorine in crystals engineering-“the little atom that could”. *Chem. Soc. Rev.* 2005, 34, 22–30. [CrossRef]
56. Gierschner, J.; Ehni, M.; Egelhaaf, H.-J.; Mili-Medina, B.; Beljonne, D.; Bennounsour, H.; Bazan, G.C. Solid-state optical properties of linear polyconjugated molecules: -stack contra herringbone. *J. Chem. Phys.* 2005, 123, 144914. [CrossRef]
57. Wang, Y.; Parkin, S.R.; Gierschner, J.; Watson, M.D. Highly fluorinated benzobisbenzothiophenes. *Org. Lett.* 2008, 10, 3307–3310. [CrossRef] [PubMed]
58. Mathew, S.; Yella, A.; Gao, P.; Humphry-Baker, R.; Curchod, B.F.E.; Ashari-Astani, N.; Tavernelli, I.; Rothlisberger, U.; Nazeeruddin, M.K.; Grätzel, M. A Low-Cost, High-Efficiency Solar Cell Based on Dye-Sensitized Colloidal TiO2 Films. *Nature* 1991, 353, 737–740. [CrossRef]
59. Benkó, G.; Kallioinen, J.; Korppi-Tommola, J.E.I.; Yartsev, A.P.; Sundström, V. Photoinduced ultrafast dye-to-semiconductor electron injection from nonthermalized and thermalized donor states. *J. Am. Chem. Soc.* 2002, 124, 489–493. [CrossRef] [PubMed]
60. Sarto Polo, A.; Itokazu, M.K.; Iha, N.Y.M. Metal complex sensitizers in dye-sensitized solar cells. *Coord. Chem. Rev.* 2004, 248, 1343–1361. [CrossRef]
61. Yum, J.-H.; Jung, I.; Baik, C.; Ko, J.; Nazeeruddin, M.K.; Grätzel, M. High efficient donor-acceptor ruthenium complex for dye-sensitized solar cell applications. *Energy Environ. Sci.* 2009, 2, 100–102. [CrossRef]
62. Mathew, S.; Yella, A.; Gao, P.; Humphry-Baker, R.; Curchod, B.F.E.; Ashari-Astani, N.; Tavernelli, I.; Rothlisberger, U.; Nazeeruddin, M.K.; Grätzel, M. Dye-sensitized solar cells with 13% efficiency achieved through the molecular engineering of porphyrin sensitizers. *Nat. Chem.* 2014, 6, 242–247. [CrossRef]
63. Qu, S.Y.; Wu, W.J.; Hua, J.L.; Kong, C.; Long, Y.T.; Tian, H. New Diketopyrrolopyrrole (DPP) Dyes for Efficient Dye-Sensitized Solar Cells. *J. Phys. Chem. C* 2010, 114, 1343–1349. [CrossRef]
64. Hagfeldt, A.; Boschloo, G.; Sun, L.C.; Kloo, L.; Pettersson, H. Dye-Sensitized Solar Cells. *Chem. Rev.* 2010, 110, 6595–6663. [CrossRef]
65. Kozma, E.; Concina, I.; Braga, A.; Borgese, L.; Depero, L.E.; Vomiero, A.; Sberveglieri, G.; Catellani, M. Metal-free organic sensitizers with a sterically hindered triphenyl borate unit for efficient dye-sensitized solar cells. *J. Mater. Chem.* 2011, 21, 13785–13788. [CrossRef]
66. Jiao, Y.; Zhang, F.; Grätzel, M.; Meng, S. Structure–Property Relations in All-Organic Dye-Sensitized Solar Cells. *Adv. Funct. Mater.* 2013, 23, 424–429. [CrossRef]
67. Kim, D.; Ghhoov, A.; Albu, S.P.; Schmuki, P. Bamboo-Type TiO2 Nanotubes: Improved Conversion Efficiency in Dye-Sensitized Solar Cells. *J. Am. Chem. Soc.* 2008, 130, 16454–16455. [CrossRef]
88. Yao, Z.; Zhang, M.; Li, R.; Yang, L.; Qiao, Y.; Wang, P. A Metal-Free N-Annulated Thiencyclopentaperylene Dye: Power Conversion Efficiency of 12% for Dye-Sensitized Solar Cells. Angew. Chem. Int. Ed. 2015, 54, 5994–5998. [CrossRef]
89. Baldoli, C.; Bertuolo, S.; Licandrello, E.; Viglianti, L.; Musinisi, P.; Marotta, G.; Salvatori, P.; De Angelis, F.; Manca, P.; Manfredi, N.; et al. Benzothiophene-based organic dyes for DSSC: Effect of alkyl chain substitution on dye efficiency. Dyes Pigm. 2015, 121, 351–362. [CrossRef]
90. Baheti, A.; Gajjela, S.R.; Balaya, P.; Justin Thomas, K.R. Synthesis, optical, electrochemical and photovoltaic properties of organic dyes containing trifluoromethylamine donors. Dyes Pigm. 2015, 113, 78–86. [CrossRef]
91. Justin Thomas, K.R.; Venkateswararao, A.; Lee, C.-P.; Ho, K.-C. Organic dyes containing fluoreneamine donor and carbazole \( \pi \)-linker for dye-sensitized solar cells. Dyes Pigm. 2015, 123, 154–165. [CrossRef]
92. Kumar, S.; Justin Thomas, K.R.; Li, C.-T.; Ho, K.-C. Synthesis and photovoltaic properties of organic dyes containing \( N \)-fluoren-2-yl dithieno[3,2-b:2′,3′-d]pyrrole and different donors. Org. Electron. 2015, 26, 109–116. [CrossRef]
93. Liu, J.; Sun, X.; Li, Z.; Jin, B.; Lai, G.; Li, H.; Wang, C.; Chen, Y.; Hua, J. New D-π-A system dye based on dithienosilole and carbazole: Synthesis, photo-electrochemical properties and dye-sensitized solar cell performance. J. Photochem. Photobiol. A 2014, 294, 54–61. [CrossRef]
94. Chung, C.-L.; Chen, C.-H.; Tsai, C.-H.; Wonga, K.-T. Novel organic dyes containing \( N \)-bridged oligothiophene coplanar cores for dye-sensitized solar cells. Org. Electron. 2015, 18, 8–16. [CrossRef]
95. Kumar, D.; Justin Thomas, K.R.; Lee, C.-P.; Ho, K.-C. Triarylamine-Free Pyrenoidimidazole-Containing Organic Dyes with Different \( \pi \)-Linkers for Dye-Sensitized Solar Cells. Asian J. Org. Chem. 2015, 4, 164–172. [CrossRef]
96. Baheti, A.; Justin Thomas, K.R.; Li, C.-T.; Lee, C.-P.; Ho, K.-C. Fluorene-Based Sensitizers with a Phenothiazine Donor: Effect of Mode of Donor Tethering on the Performance of Dye-Sensitized Solar Cells. ACS Appl. Mater. Interfaces 2015, 7, 2249–2262. [CrossRef] [PubMed]
97. Wang, D.; Ying, W.; Zhang, X.; Hu, Y.; Wu, W.; Hua, J. Near-infrared absorbing isoindigo sensitizers: Synthesis and performance for dye-sensitized solar cells. Dyes Pigm. 2015, 112, 327–334. [CrossRef]
98. Heeney, M.; Bailey, C.; Giles, M.; Shkunov, M.; Sparrowe, D.; Tierney, S.; Zhang, W.; McCulloch, I. Alkylidene Fluorene Liquid Crystalline Semiconducting Polymers for Organic Field Effect Transistor Devices. Macromolecules 2004, 37, 5250–5256. [CrossRef]
99. Li, Q.; Blancafort, L. A conical intersection model to explain aggregation induced emission in diphenyl dibenzofulvene. Chem. Commun. 2013, 49, 5966–5968. [CrossRef]
100. Du, C.; Li, C.; Li, W.; Chen, X.; Bo, Z.; Veit, C.; Ma, Z.; Wuefei, U.; Zhu, H.; Hu, W.; et al. 9-Alkylidene-9H-Fluorene-Containing Polymer for High-Efficiency Polymer Solar Cells. Macromolecules 2011, 44, 7617–7624. [CrossRef]
101. Liu, L.-Q.; Zhang, G.-C.; Liu, P.; Zhang, J.; Dong, S.; Wang, M.; Ma, Y.-G.; Yap, H.-L.; Huang, F. Donor–Acceptor-Type Copolymers Based on a Naphtho[1,2-c:5,6-c]bis(1,2,5-thiadiazole) Scaffold for High-Efficiency Polymer Solar Cells. Chem. Asian J. 2014, 9, 2104–2112. [CrossRef]
102. Choi, M.-H.; Song, K.W.; Moon, D.K. Alkylidenefluorene–isoindigo copolymers with an optimized molecular conformation for spacer manipulation, \( \pi \)-\( \pi \) stacking and their application in efficient photovoltaic devices. Polym. Chem. 2015, 6, 2636–2646. [CrossRef]
103. Capodilupo, A.L.; Giannuzzi, R.; Corrente, G.A.; De Marco, L.; Fabiano, E.; Cardone, A.; Gigli, G.; Ciccarella, G. Synthesis and photovoltaic performance of dibenzofulvene-based organic sensitizers for DSSC. Tetrahedron 2016, 72, 5788–5797. [CrossRef]
104. Capodilupo, A.L.; De Marco, L.; Corrente, G.A.; Giannuzzi, R.; Fabiano, E.; Cardone, A.; Gigli, G.; Ciccarella, G. Synthesis and characterization of a new series of dibenzofulvene based organic dyes for DSSCs. Dyes Pigm. 2016, 130, 79–89. [CrossRef]
105. Corrente, G.A.; Fabiano, E.; De Marco, L.; Accorsi, G.; Giannuzzi, R.; Cardone, A.; Gigli, G.; Ciccarella, G.; Capodilupo, A.L. Effects of donor position on dibenzofulvene-based organic dyes for photovoltaics. J. Mater. Sci. Mater. Electron. 2017, 28, 8694–8707. [CrossRef]
106. Mahmood, A.; Wang, J.-L. Machine learning for high performance organic solar cells: Current scenario and future prospects. Energy Environ. Sci. 2021, 14, 90–105. [CrossRef]
107. Mahmood, A.; Irfan, A.; Wang, J.-L. Machine learning for Organic Photovoltaic Polymers: A Minireview. Chin. J. Polym. Sci. 2022, 40, 870–876. [CrossRef]
108. Mahmood, A.; Irfan, A.; Wang, J.-L. Machine learning and molecular dynamics simulation-assisted evolutionary design and discovery pipeline to screen efficient small molecule acceptors for PTB7-Th-based organic solar cells with over 15% efficiency. J. Mater. Chem. A 2022, 10, 4170–4180. [CrossRef]
109. Mahmood, A.; Irfan, A.; Wang, J.-L. Developing Efficient Small Molecule Acceptors with sp\(^2\)-Hybridized Nitrogen at Different Positions by Density Functional Theory Calculations, Molecular Dynamics Simulations and Machine Learning. Chem. A Eur. J. 2022, 28, e202103712. [CrossRef]