# Molecular Level Factors Affecting the Efficiency of Organic Chromophores for p-Type Dye Sensitized Solar Cells

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#### Abstract:

A series of mono- and di-branched donor-?-acceptor charge-separated dyes incorporating triphenylamine as a donor and either Dalton's or benzothiadiazole group as strong acceptors was synthesized and its fundamental properties relevant to the sensitization of nanocrystalline NiO investigated. The dyes exhibited an intense visible absorption band with a strong charge transfer character favorable to NiO sensitization, shifting the electron density from the donor to the acceptor branches. Nevertheless, the computed exciton binding energy is circa twice that of a common literature standard (P1), suggesting a more difficult charge separation. When tested in p-type dye-sensitized solar cells the dyes successfully sensitized NiO electrodes, with photocurrent densities about half than that of the reference compound. Being recombination kinetics comparable, the larger photocurrent generated by P1 agrees with the superior charge separation capability originating by its smaller exciton binding energy.

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### Article Molecular Level Factors Affecting the Efficiency of Organic Chromophores for *p*-Type Dye Sensitized Solar Cells

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**Abstract:** A series of mono- and di-branched donor- $\pi$ -acceptor charge-separated dyes incorporating triphenylamine as a donor and either Dalton's or benzothiadiazole group as strong acceptors was synthesized and its fundamental properties relevant to the sensitization of nanocrystalline NiO investigated. The dyes exhibited an intense visible absorption band with a strong charge transfer character favorable to NiO sensitization, shifting the electron density from the donor to the acceptor branches. Nevertheless, the computed exciton binding energy is *circa* twice that of a common literature standard (P1), suggesting a more difficult charge separation. When tested in *p*-type dye-sensitized solar cells the dyes successfully sensitized NiO electrodes, with photocurrent densities about half than that of the reference compound. Being recombination kinetics comparable, the larger photocurrent generated by P1 agrees with the superior charge separation capability originating by its smaller exciton binding energy.

**Keywords:** dyes; heteroaromatic; visible absorption; dipolar; donor–acceptor; triphenylamine; branched; Suzuki coupling; DFT-TDDFT

#### 1. Introduction

With the increasing demand of clean, secure, cost-effective, and renewable energy sources, the exploitation of solar light as a major source has clearly emerged as a key strategic priority. Following the first publication of Grätzel and O'Regan in 1991 [1], dye-sensitized solar cells (DSSCs) were recognized as a relatively cheap and easy-to-scale approach to direct solar-to-electrical power conversion. Furthermore, their transparency, versatile design and wide color palette offer unique structural and architectural possibilities in the emerging field of building integration [2,3], for example through the realization of photovoltaic windows and façades. The key feature of DSSCs is the sensitization of a porous wide-band-gap semiconductor thin film with a photoactive dye, which, following excitation, is able to transfer either electrons (n-type sensitization) or holes (p-type sensitization) to a semiconductor substrate. n-type DSSCs have been intensively investigated over the last two decades, recently reaching optimized power conversion efficiencies up to 13% with a single photoactive junction (dye/n-TiO<sub>2</sub>) [4]. Despite this recent progress, power conversion efficiencies in n-type DSSCs seem to have reached a plateau, urging the search for new viable approaches to further increase DSSCs efficiencies. A particularly promising strategy to produce DSSCs with significantly

enhanced power conversion efficiencies is the connection of an *n*-type photoelectrode (*n*-dye/TiO<sub>2</sub>) with a *p*-type photoelectrode (*p*-dye/NiO), affording a tandem cell composed by two serially connected photoactive electrodes, each contributing to the total photovoltage delivered by the cell. Applying such a concept, organic-based photovoltaic devices with up to 40% conversion efficiency could be theoretically achievable [5]. Unfortunately, thus far, sensitized *p*-type systems have been much less investigated and much lower photocurrents, compared to their *n*-type counterparts, have been reported [6]. One of the main limitations in *p*-type systems, commonly based on NiO as a hole semiconductor [7], arises from the fast charge recombination [8] between the photoinjected hole in NiO and the reduced dye. Therefore, it is essential to develop new *p*-type chromophores, which could produce a long-lived charge separated state and minimize back recombination.

Several families of *p*-type dyes have been so far synthesized for this purpose. Those based on coumarin, porphyrin and peryleneimide scaffolds showed incident monochromatic photon-to-current conversion efficiencies (IPCE) only up to 4% and overall efficiencies lower than 0.2% [9–12]. One of the best examples of improved charge separation through dye design is represented by push–pull systems based on a di-branched D-( $\pi$ -A)<sub>2</sub> (where D = electron-donor group,  $\pi = \pi$ -spacer, and A = electron-acceptor group) structures, like **P1**, firstly reported in 2008 by Sun and coworkers [13]. This prototypical *p*-type dye had a carboxylic anchoring group on the triphenylamine donor moiety bridged to a dicyanovinyl acceptor by a thienylene ring acting as a  $\pi$ -linker. The di-branched architecture, constituted by one donor and two  $\pi$ -spacer and acceptor units, follows the same design strategy used for recent *n*-type DSSC sensitizers [14–16]. Using the same general and successful principles, more complex dyes based on much more elaborate triphenylamine/oligothiophene dyes have soon followed [17–22]. Nevertheless, even in the best cases, the power conversion efficiencies obtained by *p*-type devices.

We were thus triggered to explore new structures for potentially efficient chromophores for *p*-type devices, by considering that the intramolecular charge transfer, at the basis of efficient charge separation in donor–acceptor dyes, is strongly dependent on the electron-withdrawing ability of the acceptor. We have therefore designed and investigated new *p*-type dyes where **P1**-like structures have been optimized by inserting stronger and more efficient electron-acceptor groups compared to the conventional dicyanovinyl moiety. First, we have designed the **SK2** dye, where the dicyanovinyl group of **P1** has been replaced by the much stronger 3-cyano-2-(dicyanomethylene)-4,5,5-trimethyl-2,5-dihydrofuran (Dalton's acceptor) group, widely used in other materials science fields [23]. Second, mono- and di-branched dyes (**SK3** and **SK4**), characterized by a D- $\pi$ -A- $\pi$  architecture, were realized. In the latter dyes, an additional benzothiadiazole-based acceptor-spacer unit was introduced both in linear (**SK3**) and branched (**SK4**) geometrical motifs for the manifold purpose of increasing the transition dipole moment, improving the spatial separation between hole and electron, and favoring the electron transfer to the electron mediator ( $I^-/I_3^-$ ), resulting in improved interfacial charge separation.

Herein, we report the synthesis of the new *p*-type chromophores as well as their computational, electrochemical and photoelectrochemical properties in comparison with the literature standard **P1**. We have investigated the new dyes (Figure 1) as *p*-type sensitizers in *p*-type DSSC and the results were compared with the reference dye **P1**.



Figure 1. Structure of the investigated and reference (P1) dyes.

#### 2. Results and Discussion

#### 2.1. Synthesis

SK2, SK3 and SK4 were obtained following a synthetic pathway shown in Scheme 1. 4-[Bis-(4-bromophenyl)amino]benzaldehyde (1) was oxidized to the corresponding benzoic acid 2 by standard silver mirror reaction and then submitted to the Suzuki coupling with 5-formyl-2-thienylboronic acid for introducing the two thienyl linkers [13]. Knoevenagel reaction between the resulting bis-aldehyde 3 and 2-(3-cyano-4,5,5-trimethylfuran-2(5H)-ylidene)malononitrile afforded the desired chromophore SK2. It should be noted that the Knoevenagel condensation of 3 could be successfully accomplished only in acidic conditions at relatively high temperatures (>75 °C), while the reaction under conventional basic conditions (piperidine/EtOH) did not afford the condensation product, likely for the presence of the terminal COOH substituent on the donor core.

Preparation of **SK3** and **SK4** started from bromination of 4,7-di(thiophen-2-yl)benzo[c][1,2,5] thiadiazole (4) to the mono-bromide 5 using *N*-bromosuccinimide (NBS) in presence of a 1:1 solution of  $CH_2Cl_2$  and acetic acid [24]. Since such reaction gave mono- and di-substituted products possessing close polarities, the separation of these two compounds by column chromatography had to be performed with care in order to afford the mono-derivative in moderate yields. Borination of 5 with *bis*(pinacolato)diboron resulted in the key intermediate boronic ester 6. Different reaction conditions (solvent, base, and temperature) were investigated for Suzuki condensation between 6 and triphenylamine derivatives. The best choice in both cases was dimethoxyethane (DME) as a solvent and aqueous solution of  $K_2CO_3$  as a base. Under these conditions, the coupling reaction started immediately after mixing up the reagents and within a considerably short time afforded the desired targets 7 and 8 with fairly good yields. **SK3** and **SK4** were then obtained by following the previously described procedure for **SK2**.



Scheme 1. Synthesis of SK2, SK3 and SK4. Reagents and conditions: (i) Ag<sub>2</sub>O, EtOH, NaOH, r.t.; (ii) Pd(dppf)Cl<sub>2</sub>, 5-formyl-2-thienylboronic acid, K<sub>2</sub>CO<sub>3</sub>, toluene/MeOH, microwave; (iii) 2-(3-cyano-4,5,5-trimethylfuran-2(5*H*)-ylidene)malononitrile, NH<sub>4</sub>OAc/AcOH, EtOH, reflux; (iv) NBS, CH<sub>2</sub>Cl<sub>2</sub>/AcOH, r.t.; (v) bis(pinacolato)diboron, Pd(dppf)Cl<sub>2</sub>, KOAc, dioxane, reflux; (vi) 4-bromo-*N*,*N*-bis(4-formylphenyl)aniline, Pd(dppf)Cl<sub>2</sub>, aq. K<sub>2</sub>CO<sub>3</sub>, DME, reflux; (vii) Ag<sub>2</sub>O, EtOH, NaOH, r.t.; (viii) 4-(bis(4-bromophenyl)amino)benzaldehyde, Pd(dppf)Cl<sub>2</sub>, aq. K<sub>2</sub>CO<sub>3</sub>, DME, reflux; and (ix) Ag<sub>2</sub>O, EtOH, NaOH, r.t.

#### 2.2. Spectroscopic and Electrochemical Properties

The absorption spectra of **SK2**, **SK3** and **SK4** in dimethyl sulfoxide (DMSO) are depicted in Figure 2 together with that of the reference dye **P1**. The normalized absorption and emission spectra of the SK dye series are reported in Figure S1 (Supplementary Materials). All dyes under investigations exhibited similar general spectral features, summarized by two intense ( $\varepsilon > 3 \times 10^4 \text{ M}^{-1} \cdot \text{cm}^{-1}$ ) relatively broad absorption bands extending the light harvesting up to 700 nm in the case of **SK2**, where the presence of an efficient intramolecular charge transfer to the strong electron-withdrawing groups resulted in a significant bathochromic shift (*ca.* 70 nm) of the visible absorption maximum. **SK3**, **SK4**, and **P1** are *ca.* 100 nm blue-shifted with respect to **SK2**, showing a visible absorption maximum at 480–490 nm and an absorption peak in the interval 350–370 nm, where **SK3** and his di-branched analogue **SK4**, incorporating the same donor–acceptor side arm, showed the sharpest and most intense band. The main optical and electrochemical parameters together with HOMO/LUMO energies estimated by cyclic voltammetry in 0.1 M TBAClO<sub>4</sub> in DMSO are collected in Table 1.



Figure 2. Absorption spectra of P1, SK2, SK3 and SK4 in dimethyl sulfoxide (DMSO).

Dye	$\lambda_{abs}(nm)$ ( $\epsilon$ ) (10 <sup>4</sup> M <sup>-1</sup> ·cm <sup>-1</sup> )	λ <sub>em</sub> (nm)	E°° (eV) <sup>b</sup>	E <sub>(HOMO)</sub> vs. NHE(V) (vs. Vacuum) (eV) <sup>c</sup>	E <sub>(LUMO)</sub> vs. NHE (V) (vs. Vacuum) (eV) <sup>c</sup>
P1	372 (3.97); 489 (5.17)	611	2.14	1.34(-6.0)	-0.62(-4.0)
SK2	373 (3.12); 559 (4.90)	674	1.91	1.26(-5.9)	-0.22(-4.4)
SK3	357 (5.87); 488 (3.28)	609	2.21	1.17(-5.8)	-0.85(-3.8)
SK4	365 (4.76); 498 (4.11)	621	2.17	1.16(-5.8)	-0.86(-3.7)

Table 1. Optical and electrochemical parameters of the dyes. <sup>a</sup>

<sup>a</sup> in DMSO; <sup>b</sup> Calculated from the onset of the normalized absorption spectra of the dyes; <sup>c</sup> Evaluated from the DPV oxidation and reduction peak potentials in the presence of ferrocene as an internal reference ( $E_{1/2}$  Fc<sup>+</sup>/Fc = 0.68 V vs. NHE) [25] and using a potential value of -4.6 eV for NHE vs. vacuum [26].

As often observed with organic D- $\pi$ -A dyes [27,28], the electrochemical behavior (Figure S2a, Supplementary Materials) of the series under investigation is dominated by irreversible processes that complicate a rigorous thermodynamic evaluation of the redox levels relevant to NiO sensitization. For these reasons, we have used the Differential Pulsed Voltammetry (DPV) to evaluate orbital energies. DPV shows higher sensitivity to Faradaic currents compared to other techniques (Figure S2b, Supplementary Materials). At anodic potentials, all chromophores presented a similar oxidation behavior resulting in an oxidation wave having a peak potential at ca. 0.5 V vs. Fc<sup>+/o</sup>, which is related to the oxidation of the electron-rich TPA group [29]. The reductive behavior is generally more complex, showing for all dyes a first, weak and irreversible wave likely originated by the reductive chemisorption of the acidic protons onto the electrode surface [30]. The following current wave corresponds to the injection of the electron into the LUMO orbital and was used to estimate the corresponding energy level. In agreement with the structural similarity of their acceptor group, SK3 and SK4 showed close LUMO energies at -3.8 eV. This estimate is in reasonable agreement with the quasi-reversible wave observed at -1.55 V vs. Fc<sup>+/ $\circ$ </sup>. The reduction of **SK2** dye, bearing a stronger acceptor group, is comparatively more anodically shifted with a lower energy LUMO at -4.5 eV. Thus, no thermodynamic limitation to hole transfer to NiO are expected: in all cases, the HOMO energies are similar to the P1 dye and sufficiently lower than the upper valence band edge of NiO, located at 0.5–0.6 V vs. NHE [17,31]. Electron transfer to  $I_3^-$ , relevant to dye regeneration, is in all cases, hexoergonic, with  $\Delta G^{\circ}$  values larger than -0.2 eV, allowing to predict favorable kinetics.

#### 2.3. Computational Investigation

To gain insights into the electronic and optical properties of the investigated sensitizers, we performed Density Functional Theory (DFT) calculations. All the calculations have been performed

using the Gaussian 09 program package [32]. The dye structures at the ground state were optimized at the B3LYP level employing the 6-31G\* basis set, starting from a semi-empirical PM3 pre-optimized geometry. Similar structural features were observed throughout the whole series. The nitrogen atom of the triphenylamine core gives rise to a distorted tetrahedral motif, resulting in a dihedral angle between the branches of the order of 40°. The  $\pi$ -A unit, constituted by thienylene bridges and dicyanovinly groups in **P1** and **SK2** and benzothiadiazole fragments in **SK3** and **SK4** (Figure 3), is essentially planar in agreement with the presence of an extended  $\pi$ -electron delocalization. Each branch is linked to the electron donating group via a benzene ring, with a dihedral angle of about 20°–27°. Such twisting angle is expected to be beneficial for decoupling holes and electrons, once the initial charge separation is achieved.



**Figure 3.** Equilibrium geometries of the *p*-type dyes under investigation calculated at the DFTB3LYP6-31G\* level.

It is notoriously difficult for Time-Dependent DFT (TDDFT) methods to reliably describe charge transfer states, particularly those involving spatially separated orbitals and long range excitations as in the case of our *p*-type systems [33,34]. Therefore, in the attempt to drive reasonable insights on the electronic structure and excited state energetics of the dye series under investigation, the excitation energies (Table 2), resulting from calculations with B3LYP [35] and BH & H (Half and Half) [36,37] functionals were compared. Frontier orbitals isodensity maps (isovalue of 0.02) of the selected dyes are depicted in Figure S3 (Supplementary Materials).

**Table 2.** Comparison of experimental and calculated lowest absorption maximum of the investigated dyes in DMSO.

Dye	E <sub>exp</sub> (eV) [λ <sub>max</sub> (nm)]	E <sup>∘p</sup> <sub>BH &amp; H</sub> (eV) [λ <sub>max</sub> (nm)]	E <sup>op</sup> <sub>B3LYP</sub> (eV) [λ <sub>max</sub> (nm)]	∆E <sub>BH &amp; H</sub>   (eV)	∆E <sub>B3LYP</sub>  (eV)
P1	2.53 [489]	2.79 [443]	2.30 [540]	0.26	0.23
SK2	2.22 [559]	2.86 [433]	2.08 [595]	0.64	0.14
SK3	2.54 [488]	2.62 [473]	2.14 [580]	0.08	0.40
SK4	2.49 [498]	2.48 [501]	1.92 [614]	0.01	0.57

The BH & H is considerably successful in predicting the optical transition energies of dyes **SK3** and **SK4**, where deviations from the experimentally measured spectrum are as low as 0.08 and 0.01 eV, but less successful with dyes **P1** and **SK2**, where the calculated  $E^{\circ p}$  is higher than the experimental value by

0.26 and 0.64 eV, respectively. The lowest transition has, in all cases, a major HOMO-LUMO component (55%–87%) in the Kohn–Sham basis, with minor components arising from higher energy excitations, typically HOMO-1 $\rightarrow$ LUMO, HOMO-1 $\rightarrow$ LUMO + 1 and HOMO $\rightarrow$ LUMO + 2. The description of the lowest singlet state (S1) by the B3LYP calculation is roughly similar to that of the BH & H, although the HOMO-LUMO component to the excitation is largely predominating (>93%). The lowest transition in P1 and SK2 is identified by B3LYP with a lower absolute error (0.21 and 0.11 eV) than the BH & H functional, whereas larger errors are found with SK3 and, particularly with SK4, which shows long range charge transfers with a high degree of spatial charge separation, as indicated by Electron Density Difference Maps (EDDM) (Scheme 2). Both the B3LYP and BH & H functional agree in the description of the EDDMs, showing a shift in electron density from the triphenylamine core to the electron acceptor branches, where the maximum density is localized in the proximity of the cyano groups (P1 and SK2) or on the benzothiadiazole acceptor (SK3 and SK4). It can also be appreciated that, in branched dyes (P1, SK2, and SK4), the electronic excitation involves the two acceptor branches simultaneously. Thus, the electronic distribution resulting from the lowest, and usually most intense, electronic transition is favorable to a successful charge separation. The hole density is localized in the immediate proximity of the surface interacting with NiO via COOH groups. This causes photoexcitation of the electron to occur predominantly on the farthest from the NiO part of the molecule where scavenging of the electron by  $I_3^-$  is more favorable.



**Scheme 2.** Electron Density Difference Maps (EDDMs) (isovalue = 0.001) of the main transition in the visible region calculated with a 6311 G,d,+ basis set obtained with the BH & H (top row) and B3LYP (bottom row) functionals. DMSO solvent was described as a polarizable continuum model (PCM). Light blue and violet indicate an increased and a depleted electron density respectively.

The exciton binding energy (EBE) [38] was calculated [39] from the B3LYP data set, since BH & H was found to give unrealistically high HOMO-LUMO gaps (Table S1, Supplementary Materials) as well as binding energies (>1 eV). For the **SK** dyes, similar binding energies of the order of 0.33–0.38 eV are found, being comparable to that of some efficient *n*-type push pull charge separators [40] recently reported in the literature. **P1** displays the smallest EBE within the series with a value as small as 0.17 eV, comparable to some of the best *n*-type charge separators reported [41]. Smaller EBE favor separation of the electron–hole pair, and, consequently, charge injection into the semiconductor. The largest binding energy value of 0.38 eV is found for the dye **SK3**, which means that a higher energy, usually provided by the local electric potential at the semiconductor/dye interface, or by collision with phonons, is needed in order to promote charge injection. The larger EBE of the **SK** series may constitute a disadvantage compared to the reference dye **P1** for obtaining hole injection with high quantum yields.

#### 2.4. Photoelectrochemical Investigation in DSSCs

The **SK** dyes together with reference compound **P1** were tested as photosensitizers in p-type DSSCs. NiO photocathodes were fabricated by blading a NiO colloidal paste, obtained by dispersing commercial 25-nm NiO nanoparticles in terpineol, with ethylcellulose as an organic

binder/densificating agent, followed by high temperature sintering on FTO (Fluorine-doped Tin Oxide) coated glass. The resulting photocurrent (J)/photovoltage (V) curves under AM 1.5 illumination (1 sun) are shown in Figure 4A and their main efficiency parameters are listed in Table 3.



**Table 3.** Photovoltaic parameters of investigated chromophores in *p*-type dye-sensitized solar cells (DSSCs).

**Figure 4.** Photoelectrochemical and spectral properties of *p*-type DSSCs in the presence of 1 M LiI and 0.1 M I<sub>2</sub> in acetonitrile: (**A**) current–voltage characteristics; (**B**) Light Harvesting Efficiency (LHE) (LHE( $\lambda$ ) = 1 – 10<sup>-A( $\lambda$ )</sup>), where A( $\lambda$ ) is the background-subtracted absorbance of the dyed NiO films; (**C**) photoaction spectra (IPCE *vs.*  $\lambda$ ); and (**D**) absorbed photon conversion efficiency spectra (APCE *vs.*  $\lambda$ ).

The photoelectrochemical investigation was carried out with the iodide/triiodide redox couple (solution of 1.0 M LiI and 0.1 M I<sub>2</sub> in CH<sub>3</sub>CN). Dye **SK4** produces the highest open circuit voltage (133 mV), that is *ca*. 40 mV higher than that of device sensitized by the other reference and novel dyes. The **SK2** and **SK3** molecules produced comparable  $J_{sc}$  (*ca*. 0.5 mA/cm<sup>2</sup>), which was, however, approximately half that of the reference **P1** (*ca*. 1.1 mA/cm<sup>2</sup>). As a comparison, blank NiO cells without the sensitizer generated *ca*. 0.25 mA/cm<sup>2</sup> at 0 V, that is less than half of the cells sensitized by the SK series and *ca*. <sup>1</sup>/<sub>4</sub> of the reference P1 (Figure S4, Supplementary Materials). In all cases the sloped J/V characteristic at reverse (negative) bias is consistent with a small shunt resistance of the NiO films, causing charge leaks and losses by recombination. For instance, in the case of **P1** the photoinjected charge extracted from the photocathode under short circuit conditions (0 V) is roughly one half that available at -0.2 V. Such a photocurrent loss upon moving from negative to positive potential is less marked for the **SK** series, which, giving rise to a smaller extent of hole injection, result in fewer charge

leaks to the electrolyte. The overall PCE of **P1**-sensitized cells is about twice as large as that of the SK-based devices, mainly due to their lower photocurrents. In general, although not optimized, the efficiencies of the NiO sensitized solar cells recorded in this work are in agreement with the average performances reported so far in the literature for similar types of sensitized *p*-type electrodes.

To understand the nature of photocathodic current, the IPCE spectra of the *p*-type cells, under short circuit conditions were measured (Figure 4 C). Consistent with their spectral properties, all dyes displayed a strongest response in the blue region of the spectrum for  $\lambda < 450$  nm, reaching the best values of *ca*. 12%–14% for **P1** and **SK2**. The higher IPCE in this narrow wavelength Vis region is motivated by both a direct contribution from the I<sup>-</sup>/I<sub>3</sub><sup>-</sup> photochemistry and by a more efficient collection of the charge carriers generated in closer proximity of the electron collector by photons having a comparatively short penetration length into the film. The photoconversion then decreases by moving from the blue to the red region of the visible spectrum, showing a shoulder the 500–600 nm region, with values of 4%–5% in the case of **P1** and **SK4**. Despite the opacity of the NiO electrodes, causing significant light scattering, and the strong competitive absorption of the iodine based electrolyte, in general the photoconversion appeared to be in a relatively good agreement with the Light Harvesting Efficiency (LHE) (Figure 4B) of the sensitized NiO photoelectrodes, confirming that the excited state of the dye is responsible for the observed photocurrent in the visible region of spectrum.

The LHE plot, with values of 50%–60% in the 550–450 nm region, showed that **SK4** has here the best light harvesting performance, consistent with its relatively good IPCE response at these wavelengths. The absorbed monochromatic photon-to-current conversion efficiencies (APCE) spectra, obtained by dividing the corresponding IPCE spectra by the LHE of the respective photocathodes, are shown in Figure 4D. The low energy response of **P1** and **SK4** are similar, with APCE ~10% at 500 nm. Interestingly, **SK4** is able to yield a slightly more efficient conversion for  $\lambda \ge 550$  nm, while **P1** prevails below 480 nm. As for the IPCE, the best APCE response is observed in the blue region of the spectrum, with values of ~18%–20% for **P1** and **SK2**.

Electrochemical impedance spectroscopy (EIS) was used to further investigate the comparative behavior of the chromophores in photovoltaic devices. All dyes were stable during potentiostatic measurements, which were performed at potentials between -10 mV and the  $V_{oc}$  of the cells. The J/V curves after EIS measurements showed essentially the same performances found before the measurements, which allows to rule out substantial dye or cell degradation affecting the results. The EIS spectra (Figure S5, Supplementary Materials) could be well fitted with the equivalent circuit, comprising the serial resistance, the counter-electrode interface, the diffusion element accounting for electrolyte transport impedance in a thin layer cell, and the transmission line [42] (DX1 in Zview) which describes the transmission network of the NiO, applied by Zhang *et al.* [17,18] to *p*-type DSSCs, comprising both the transport and the charge transfer resistance (R<sub>ct</sub> (NiO)) (recombination resistance) at the NiO/electrolyte interface.

The impedance response of the NiO based DSSC exhibited similar features, appearing as a small high frequency semicircle, due essentially to the counter-electrode electrochemical interface, followed by a much larger loop at lower frequencies, which bears the main contribution of the NiO/electrolyte interface. Indeed, the J/V curves are essentially dominated by the charge transfer resistance of the NiO, as can be appreciated by the comparison of the derivative  $(\frac{\partial i}{\partial V})$  of the J/V curves and the inverse of the interfacial charge transfer resistance (R<sub>ct</sub>(NiO))<sup>-1</sup> (Figure 5B). This agreement is particularly good in the case of the **SK** series, for which the charge transfer resistance was the largely predominant contribution to the total cell resistance at each potential and varied, typically between 1000 and 200  $\Omega$ , depending on the voltage. Obviously, the lowest R<sub>ct</sub> (NiO) are in all cases found at the open-circuit voltage, due to hole accumulation into the mesoscopic film under open-circuit conditions.

The NiO capacitance (Figure 5A), calculated according to  $C = CPE(\omega)^{n-1}$ , where n is the exponent of the CPE admittance from the transmission line, in all cases variable between 0.9 and 0.84, and  $\omega$  is the frequency at the maximum of the large low frequency loop, was reasonably linear on a logarithmic

scale, indicating a chemical capacitance [43] behavior determined by empty states in the valence bands or in surface states following a Boltzmann distribution near the valence band edge. The inspection of Figure 5 reveals that **P1** is the dye which exhibits the lowest charge transfer resistance and highest capacitance, indicating a superior capability of hole injection.



**Figure 5.** (**A**) Capacitance of sensitized NiO films at increasing potentials in the forward direction. (**B**) Reciprocal of the charge transfer resistance at the NiO/electrolyte interface (squares) compared to the derivative of the J/V curve (solid lines). (**C**) Chemical capacitance of NiO *vs.* interfacial charge transfer resistance in *p*-type DSSCs under study.

The representation of  $R_{ct}$ (NiO) *vs.* C(NiO) on a logarithmic scale indicates that all the dyes behave intrinsically similarly to each other, as far as recombination kinetics are concerned, when compared at the same chemical capacitance of the NiO. In other terms, the charge loss pathways involving direct dye<sup>-</sup>/h<sup>+</sup> and hole recapture by I<sup>-</sup> occurred at comparable rates for all the dyes under investigation. Accordingly, the higher photovoltage observed for the **SK4**-sensitized device is likely due to a positive shift of the valence band edge of the NiO by *ca.* 30–40 mV, induced by the adsorption of the dye sensitizer, as can be observed by comparing the different voltages at which the similar capacitances are found, rather than by slower interfacial recombination kinetics. The voltage shift observed for **SK4** in EIS under illumination is quantitatively consistent with the more positive onset of the anodic current observed for this moiety in the dark (Figure S6, Supplementary Materials).

#### 3. Experimental Section

#### 3.1. General Remarks

NMR spectra were recorded with an instrument operating at 500.13 (<sup>1</sup>H) and 125.77 MHz (<sup>13</sup>C). Flash chromatography was performed with silica gel 230–400 mesh (60 Å). Reactions conducted under a nitrogen atmosphere were performed in oven-dried glassware and monitored by thin-layer chromatography by using UV light (254 and 365 nm) as a visualizing agent. 4-[Bis-(4-bromo-phenyl)-amino]-benzaldehyde and 4/4'-(4-bromophenylazanediyl)dibenzaldehyde were synthesised following procedure known in literature [44,45]. All other reagents were obtained from commercial suppliers at the highest purity and used without further purification. Anhydrous solvents were purchased from commercial suppliers and used as received. Extracts were dried with anhydrous Na<sub>2</sub>SO<sub>4</sub> and filtered before removal of the solvent by evaporation.

#### 3.2. Synthesis of the p-Type Sensitizers

#### 3.2.1. 4-[Bis-(4-bromo-phenyl)-amino]-benzoic Acid (2)

A procedure reported in the literature was adapted with some modifications [13]. The suspension of sodium hydroxide (62 mmol, 2.5 g) in 45 mL of ethanol was vigorously stirred for 20 min and then silver (I) oxide (9.5 mmol, 2.19 g). Resulting mixture was stirred for another 20 min before 4-[*bis*-(4-bromo-phenyl)-amino]-benzaldehyde (1) (1.2 mmol, 550 mg) in 4 mL of toluene was slowly added. The resulting suspension was stirred at room temperature for 24 h and then neutralized with HCl (15%, 25 mL) until formation of grey silver chloride precipitate was observed. The resulting solution was decanted and extracted with ethyl acetate ( $3 \times 40$  mL). The organic phase was washed with water and dried. Thereafter the solvent was removed by rotary evaporation giving the desired acid **2** (0.8 g, 78%) as white solid, which was used without further purification. <sup>1</sup>H NMR (500 MHz, DMSO-*d*<sub>6</sub>):  $\delta$  6.90 (d, *J* = 8.5 Hz, 2H), 6.93 (d, *J* = 8.8 Hz, 4H), 7.44 (d, *J* = 8.8 Hz, 4H), 7.79 (d, *J* = 8.5 Hz, 2H).

#### 3.2.2. 4-{Bis-[4-(5-formyl-thiophen-2-yl)-phenyl]-amino}-benzoic Acid (3)

The solution of benzoic acid **2** (0.4 mmol, 200.0 mg) and Pd(dppf)Cl<sub>2</sub> (0.1 mmol, 79 mg) in toluene (5 mL) was mixed with the solution of 5-formyl-2-thienylboronic acid (1.2 mmol, 187 mg) and K<sub>2</sub>CO<sub>3</sub> (2.5 mmol, 355 mg) in methanol (5 mL). The resulting mixture was irradiated in the microwave oven at 80 °C for 20 min and then poured into a saturated solution of ammonium chloride (60 mL). After extraction with ethyl acetate (3 × 100 mL), the combined organic phases were washed with brine, dried and then filtered over Celite and concentrated under reduced pressure. The crude residue was purified by column chromatography over silica gel using dichloromethane (DCM)/methanol 1:9 mixture as the eluent to give the product (145 mg, 62%) in form of yellow solid. m.p. 137–139 °C, <sup>1</sup>H NMR (500 MHz, acetone-*d*<sub>6</sub>):  $\delta$  7.21 (d, *J* = 8.8 Hz, 2H), 7.28 (d, *J* = 8.7 Hz, 4H), 7.66 (d, *J* = 4.0 Hz, 2H), 7.83 (d, *J* = 8.7 Hz, 4H), 7.98 (d, *J* = 4.0 Hz, 2H), 8.00 (d, *J* = 8.8 Hz, 2H), 9.94 (s, 2H), 11.14 (s, 1H).

### 3.2.3. 4-(*Bis*(4-(5-((*E*)-2-(4-cyano-5-(dicyanomethylene)-2,2-dimethyl-2,5-dihydrofuran-3-yl)vinyl) thiophen-2-yl)phenyl)amino)benzoic Acid (**SK2**)

To a solution of the intermediate **4** (100 mg, 0.2 mmol) and 2-(3-cyano-4,5,5-trimethylfuran-2 (5*H*)-ylidene)malononitrile (77.2 mg, 0.4 mmol) in ethanol (20 mL) was added catalytic amount of ammonium acetate (8 mg, 0.1 mmol) in 2 mL of acetic acid. Reaction mixture was refluxed under nitrogen atmosphere for 100 h. Afterwards, organic solvent was evaporated under reduced pressure and resulted crude product was washed with ethyl acetate to give the desired dye **SK2** (75 mg, 74%) in form of dark violet solid. m.p. 252–253 °C. <sup>1</sup>H NMR (500 MHz, DMSO-*d*<sub>6</sub>):  $\delta$  12.75 (s, 1H), 8.17 (d, *J* = 16 Hz, 2H), 7.92 (d, *J* = 8 Hz, 2H), 7.87 (d, *J* = 4 Hz, 2H), 7.84 (d, *J* = 8.0 Hz, 4H), 7.73 (d, *J* = 4 Hz, 2H), 7.2 (d, *J* = 9 Hz, 4H), 7.16 (d, *J* = 9 Hz, 2H), 6.83 (d, *J* = 16 Hz, 2H), 1.81 (s, 12 H). <sup>13</sup>C NMR (acetone-*d*6):  $\delta$  26.7, 54.9, 98.3, 99.2, 100.2, 112.4, 113.4, 114.2, 114.3, 123.9, 126.5, 127.2, 129, 129.6, 132.5,

139.5, 140.3, 141.7, 148.2, 151.3, 152.2, 168.1, 175.9, 178.2.  $\nu_{max}(cm^{-1})$  3583, 3476, 3007, 2310, 1738, 1590, 1373, 1217. Found: C, 70.11; H, 4.19; N, 11.60. C<sub>51</sub>H<sub>33</sub>N<sub>7</sub>O<sub>4</sub>S<sub>2</sub> requires: C, 70.25; H, 3.81; N, 11.24.

#### 3.2.4. 4-(5-Bromothiophen-2-yl)-7-(thiophen-2-yl)benzo[1,2,5]thiadiazole (5)

To a solution of 4,7-di-2-thienyl-2,1,3-benzothiadiazole (4) [46] (350 mg, 1.17 mmol) in a 100 mL of acetic acid-CH<sub>2</sub>Cl<sub>2</sub> 1:1 mixture, was added N-bromosuccinimide (NBS) (229 mg, 1.28 mmol) in DCM by small portions over 30 min. The resulting reaction mixture was stirred overnight, then poured into water (100 mL) and the separated organic layer was dried, filtered and concentrated in vacuum. Obtained crude product was purified by flash column chromatography on silica gel (DCM-hexane = 1:4) giving a target benzothiadiazole derivative (5) in form of bright orange solid with 70% yield (310 mg, 0.8mmol). <sup>1</sup>H NMR (500 MHz, CDCl<sub>3</sub>):  $\delta$  8.12 (d, 1H,), 7.88 (d, 1H), 7.8 (m, 2H), 7.46 (d, 1H), 7.22 (d, 1H), 7.15 (d, 1H).

3.2.5. 4-(5-(4,4,5,5-Tetramethyl-1,3,2-dioxaborolan-2-yl)thiophen-2-yl)-7-(thiophen-2-yl)benzo[c] [1,2,5]thiadiazole (6)

Schlenk flask was charged with 4,7-di(thiophen-2-yl)benzo[c][1,2,5]thiadiazole (5) (550 mg, 1.45 mmol), potassium acetate (440 mg, 4.48 mmol) and bis(pinacolato)diboron (490 mg, 1.93 mmol) together with the catalytic amount of Pd(dppf)Cl<sub>2</sub>(0.05 mmol, 39 mg) and 15 mL of dry dioxane. Resulting mixture was heated at reflux under argon atmosphere for 24 h then poured in water, extracted with ethyl acetate (3 × 50 mL) and dried. After organic solvent was evaporated under reduced pressure, crude product was purified on silica gel using CH<sub>2</sub>Cl<sub>2</sub>-cyclohexane 1:4 as eluent. Product was obtained in form of brownish orange solid (318 mg, 55% yield). m.p. 168 °C–170 °C. <sup>1</sup>H NMR (500 MHz, CDCl<sub>3</sub>):  $\delta$  = 8.18 (d, *J* = 4 Hz, 1H) 8.13 (m, 1H), 7.92 (d, *J* = 8 Hz, 1H), 7.87 (d, *J* = 8 Hz, 1H), 7.72 (d, *J* = 3 Hz, 1H), 7.45 (m, 1H), 7.21 (m, 1H), 1.38 (s, 12H). <sup>13</sup>C NMR (CDCl<sub>3</sub>):  $\delta$  = 152.66 (CH), 152.63 (CH), 145.89 (CH), 139.32 (CH), 137.89 (C), 128.65 (C), 128.03 (C), 127.67 (C), 126.97 (C), 126.48 (CH), 126.37 (C), 125.83 (CH), 125.75 (C), 84.26 (CH3), 24.79 (C) ppm. Found C, 56.39; H, 4.60; N, 6.29. C<sub>20</sub>H<sub>19</sub>BN<sub>2</sub>O<sub>2</sub>S<sub>3</sub> requires: C, 56.34; H, 4.49; N, 6.57.

3.2.6. 4,4'-(4-(5-(7-(Thiophen-2-yl)benzo[c][1,2,5]thiadiazol-4-yl)thiophen-2-yl)phenylazanediyl) dibenzaldehyde (7)

4,4'-(4-Bromophenylazanediyl)dibenzaldehyde (114 mg, 0.3 mmol), benzothiadaizole derivative 6 (130 mg, 0.31 mmol), a catalytic amount of Pd(dppf)Cl<sub>2</sub> (0.019 mmol, 18 mg) and a solution of K<sub>2</sub>CO<sub>3</sub> (400 mg, 2.9 mmol) in water (3 mL) were charged into Schlenk flask and then dissolved in 15 mL of dry dimethoxyethane (DME). Reaction mixture was refluxed under argon atmosphere for 15 h and after cooling down to the room temperature it was washed with water, dried and concentrated under vacuum. Crude product was purified by column chromatography on silica gel using ethyl acetate-cyclohexane = 2:7 as the eluent. The pure intermediate 7 (55 mg, 30% yield) was obtained in form of bright red solid. m.p. 173 °C–175 °C. <sup>1</sup>H NMR (500 MHz, CDCl<sub>3</sub>): δ = 9.91 (s, 2H), 8.12 (dd, J = 8.0; 15 Hz, 2H), 7.86 (s, 1H), 7.81 (d, J = 17.0 Hz, 4H), 7.68 (d, J = 17.0 Hz, 2H), 7.45 (d, J = 8.0 Hz, 1H), 7.39 (d, J = 8.0 Hz, 1H), 7.24 (d, J = 17.0 Hz, 4H), 7.21 (d, J = 8.0 Hz, 1H), 7.17 (d, J = 17.0 Hz, 2H).<sup>13</sup>C NMR (CDCl<sub>3</sub>):  $\delta = 191.4, 153.5, 153.4, 152.6, 145.8, 140.18, 139.9, 132.7, 132.4, 132.2, 129.5, 128.9, 128.5, 128.2, 127.89, 127.83, 126.9, 126.6, 126.5, 126.3, 125.2, 123.9 ppm. Found: C, 67.84; H, 3.63; N, 7.33. C<sub>34</sub>H<sub>21</sub>N<sub>3</sub>O<sub>2</sub>S<sub>3</sub> requires: C, 68.09; H, 3.53; N, 7.01.$ 

3.2.7. 4-(*Bis*(4-(5-(7-(thiophen-2-yl)benzo[c][1,2,5]thiadiazol-4-yl)thiophen-2-yl)phenyl)amino) benzaldehyde (8)

4-[*Bis*-(4-bromo-phenyl)-amino]-benzaldehyde (1) (100 mg, 0.232 mmol) and benzothiadaizole derivative **6** (213 mg, 0.5 mmol) and K<sub>2</sub>CO<sub>3</sub> (691 mg, 5 mmol)were dissolved in 5 mL of water and charged into Schlenk flask together with the catalytic amount of Pd(dppf)Cl<sub>2</sub> (0.02 mmol, 19 mg) and then dissolved in 15 mL of dry DME. Reaction mixture was refluxed under argon atmosphere for

15 h. After cooling to the room temperature it was washed with water, dried and concentrated under vacuum. Crude product was washed with ethyl acetate and filtered. The desired intermediate **8** was obtained as dark red solid (70 mg, yield 45%). m.p. 189 °C–190 °C (decomp.). <sup>1</sup>H NMR (500 MHz, CDCl<sub>3</sub>):  $\delta$  = 9.88 (s, 1H), 8.14 (d, *J* = 8.0, 4H), 7.9 (s, 1H), 7.77 (d, *J* = 17.0 Hz, 2H), 7.69 (d, *J* = 17.0 Hz, 4H), 7.47 (d, *J* = 8.0 Hz, 2H), 7.41 (d, *J* = 8.0 Hz, 2H), 7.23 (d, *J* = 17.0 Hz, 4H), 7.22 (d, *J* = 8.0 Hz, 2H), 7.19 (d, *J* = 17.0 Hz, 2H). <sup>13</sup>C NMR (CDCl<sub>3</sub>):  $\delta$  = 191.4, 153.6, 153.5, 153.4, 146.4, 145.3, 140.2, 139.5, 132.3, 131.7, 130.9, 129.6, 128.9, 128.4, 127.9, 127.8, 127.7, 127.0, 126.8, 126.6, 126.2, 124.9, 121.6 ppm. Found C 64.50; H 2.88; N 7.91. C<sub>47</sub>H<sub>27</sub>N<sub>5</sub>OS<sub>6</sub> requires: C, 64.87; H, 3.13; N, 8.05.

3.2.8. 4,4'-(4-(5-(7-(Thiophen-2-yl)benzo[c][1,2,5]thiadiazol-4-yl)thiophen-2-yl)phenylazanediyl) dibenzoic Acid (**SK3**)

A suspension of sodium hydroxide (600 mg, 15 mmol) in 10 mL of ethanol was vigorously stirred for 15 min before Ag<sub>2</sub>O (208 mg, 0.8 mmol) was added. After another 15 min, dibenzaldehyde 7 (50 mg, 0.08 mmol) in toluene was added dropwise to the solution. Reaction mixture was stirred overnight at room temperature and quenched by the slow addition of 1 M HCl solution, then it was extracted with CH<sub>2</sub>Cl<sub>2</sub> (3 × 75 mL). Combined organic layers were washed with water and dried. After evaporation of solvent under reduced pressure resulting crude product was washed with diethyl ether to give final compound **SK3** in form of dark red solid (40 mg. 87% yield). m.p. 215 °C–217 °C. <sup>1</sup>H NMR (500 MHz, DMSO-*d*6):  $\delta$  = 8.20 (t, *J* = 9 Hz, 2H), 8.17 (d, *J* = 8 Hz, 1H), 8.12 (d, *J* = 8.0 Hz, 1H), 7.92 (d, *J* = 8.7 Hz, 4H), 7.80 (d, *J* = 8.5 Hz, 2H), 7.78 (d, *J* = 5.1 Hz, 1H), 7.67 (d, *J* = 4.0 Hz, 1H), 7.29 (d, *J* = 8.7 Hz, 1H), 7.22 (d, *J* = 8.5 Hz, 2H), 7.15 (d, *J* = 8.7 Hz, 4H). <sup>13</sup>C NMR (DMSO-*d*6):  $\delta$  = 168.1, 153.0, 152.9, 151.4, 146.7, 145.7, 139.8, 138.9, 132.5, 131.4, 130.1, 129.7, 129.5, 128.8, 128.3, 127.7, 127.2, 126.9, 126.4, 126.3, 126.0, 125.9, 124.0 ppm.  $\nu_{max}$  (cm<sup>-1</sup>) 3456, 2970, 2281, 1738, 1494, 1366, 1217. Found: C, 64.36; H, 3.74; N, 6.94. C<sub>34</sub>H<sub>21</sub>N<sub>3</sub>O<sub>4</sub>S<sub>3</sub> requires: C, 64.64; H,3.35; N, 6.65.

3.2.9. 4-(*Bis*(4-(5-(7-(thiophen-2-yl)benzo[c][1,2,5]thiadiazol-4-yl)thiophen-2-yl)phenyl)amino) benzoic Acid (**SK4**)

To the vigorously stirred suspension of NaOH (300 mg, 7.5 mmol) in 10 mL of ethanol was added silver (I) oxide (205 mg, 0.8 mmol). Mixture was stirred for 15 min before adding benzaldehyde **8** (50 mg, 0.057 mmol) in CH<sub>2</sub>Cl<sub>2</sub>. Reaction mixture was stirred overnight at the room temperature and quenched by the slow addition of 1 M HCl solution, extracted with ethyl acetate (3 × 75 mL) and then combined organic layers were washed with water and dried. After evaporation of solvents, crude product was washed with dichloromethane. **SK4** dye was obtained in form of black-red solid (40 mg, 90% yield). m.p. > 285 °C (decomp.). <sup>1</sup>H NMR (500 MHz, CDCl<sub>3</sub>):  $\delta$  = 8.22 (d, *J* = 3.9 Hz, 2H), 8.18 (s, 2H), 8.14 (d, *J* = 8.3 Hz, 2H), 7.88 (d, *J* = 8.7 Hz, 2H), 7.81 (d, *J* = 8.5 Hz, 4H), 7.79 (d, *J* = 5.5 Hz, 2H), 7.68 (d, *J* = 4.1 Hz, 2H), 7.31 (t, *J* = 8.8 Hz, 2H), 7.23 (d, *J* = 8.5 Hz, 4H), 7.11 (d, *J* = 8.7 Hz, 2H). <sup>13</sup>C NMR (DMSO-*d*<sub>6</sub>):  $\delta$  = 169.07, 153.5, 153.3, 151.8, 151.7, 146.97, 145.8, 142.2, 132.0, 131.5, 130.8, 129.6, 128.9, 128.2, 127.8, 127.7, 126.9, 126.7, 126.6, 126.3, 126.1, 124.6, 122.3 ppm.  $v_{max}$  (cm<sup>-1</sup>) 3472, 2976, 2110, 1738, 1546, 1372, 1254. Found C 63.88, H 3.44, N 7.68. C<sub>47</sub>H<sub>27</sub>N<sub>5</sub>O<sub>2</sub>S<sub>6</sub> requires: C, 63.70; H, 3.07; N, 7.90.

#### 3.3. Solar Cell Assembly

NiO films were prepared by grounding 7.5 g of NiO nanoparticles (Inframat, nominal size 25 nm) in a mortar in the presence of 200 mL of acetilacetone. 100 mL of terpineol were then added, followed by 10 g of ethlycellulose dissolved in 110 mL of absolute ethanol. The resulting mixture was homogenized by stirring and sonication in an ultrasonic bath. Ethanol was evaporated under reduced pressure leaving a dense paste constituted by terpineol, ethylcellulose and NiO nanoparticles. The resulting paste could be spread by blading onto well cleaned FTO glass to form, after drying and high temperature sintering, the NiO thin films. The temperature program adopted for drying and sintering in air was the following: 10 min at 120 °C, followed by a ramp (15 °C/min) to 450 °C which

were maintained for 30 min. A subsequent ramp (10 °C/min) brought the temperature from 450 °C to 550 °C which were kept constant for further 20 min. Cooling occurred naturally, by interrupting the heating. Sensitization of the resulting NiO electrodes, having an active surface of 0.25 cm<sup>2</sup>, was carried out overnight in the dark, by immersion in the DMSO solutions at the concentration of  $2 \times 10^{-4}$  M of the selected sensitizers. Solar cells were fabricated by clamping the sensitized NiO photocathode with a platinum coated FTO counter electrode (Chimet, Badia al Pino, Italy). The electrolyte was 1 M LiI/0.1 M I<sub>2</sub> in acetonitrile.

#### 3.4. Spectroscopic, Electrochemical, and Photoelectrochemical Measurements

UV-Vis spectra of sensitized NiO thin films, collected in transmission mode, were recorded on a JASCO V570 UV-Vis-NIR spectrophotometer (Jasco Analytical instruments, Easton, MD, USA). UV-Vis spectra in solution were obtained with a V-570 JASCO spectrophotometer (Jasco Analytical instruments, Easton, MD, USA). Emission spectra were recorded FP6200JASCO spectrofluorometer. TBAICIO<sub>4</sub> 0.1 DMSO.

Cyclic voltammetry of the dye dissolved in 0.1 M solution of tetrabutylammonium perchlorate (Fluka, electrochemical grade, 99.0%) in anhydrous DMSO (Sigma-Aldrich) as the supporting electrolyte was carried out at a glassy carbon working electrode with a PARSTA2273 potentiostat in a two-chamber three-electrodeelectrochemical cell with a scan rate of 50 mVs<sup>-1</sup>. Potentials are referred to the ferrocenium/ferrocene (Fc<sup>+</sup>/Fc) couple as internal reference. An Ecochemie PGSTAT 302/N electrochemical workstation equipped with the FRA 2 Frequency Response Analyzer and running under either GPES or Nova software was used for collecting both the JV characteristics and the EIS response of the p-DSSCs. Solar cells were illuminated under simulated solar conditions (Am 1.5 G 100 mW/cm<sup>2</sup>) generated by an ABET sun simulator. J/V curves were recorded by linear scan voltammetry, using a scan rate of 5 mV/s. Every cell was left to equilibrate under illumination until a superimposable J/V response was obtained upon subsequent scans. Curves reported in this work, representative of the average performance of three cells, are those measured after steady state was achieved. Potentiostatic EIS data were collected on a single cell at various voltages under illumination by applying a 10 mV sinusoidal frequency variable from 10<sup>5</sup> to 10<sup>-2</sup> Hz. Impedance data were fitted with Zview.

IPCE data were collected under monochromatic illumination, having a band pass of 10 nm, generated by an Applied Photophysics monochromator coupled to a 175 W Luxtel Xe arc lamp. Short circuit photocurrents were recorded on an Agilent 34401A multimeter. Incident irradiance was measured with a calibrated Centronic OSD 7Q silicon photodiode having an active area of 1 cm<sup>2</sup>.

#### 3.5. Computational Studies

Optimized ground state geometry of the *p*-type dyes were obtained at the DFT-B3LYP level by using a 631G\* basis set. The DFT calculation was carried out on pre-optimized structures at the PM6 level. Time dependent (TDDFT) calculations in the presence of DMSO solvent, treated as continuum polarizable medium, were carried out on the optimized structures by considering the two different B3LYP and BH & H functionals with the 6311 G\* basis set incorporating also diffuse functions (+). All calculations were carried out with Gaussian 09 A02 [32] on multi-core computers. Structures and isodensity surfaces were graphically visualized with Gaussview 5. Electron Density Difference Maps (EDDMs) were generated with GaussSum 2.2 [47].

#### 4. Conclusions

New *p*-type photosensitizers based on organic linear and di-branched donor–acceptor systems bearing tryphenylamine as a donor and strong electron-acceptor Dalton's (**SK2**) or benzothiadiazole (**SK3** and **SK4**) groups were synthesized and characterized by steady state spectroscopic, electrochemical, and computational studies. All the dyes under investigation exhibited strong charge transfer bands in the visible regions with ground and excited state energetic which are favorable to the

sensitization of NiO electrodes. The computational investigation revealed a clear directionality of the lowest excited state exhibiting a marked charge transfer character, shifting the electron density from the donor core to the acceptor branches, an electronic situation that is favorable to the hole injection in *p*-type semiconductors such as NiO. However, the exciton binding energy, *i.e.*, the energy of the bound electron-hole pair, is about twice as large as that of a known literature standard (**P1**), suggesting that a more difficult charge separation in the new dyes might occur.

When tested as photosensitizers in *p*-type DSSCs, the **SK** series was able to successfully sensitize NiO electrodes, resulting in photocurrents that are about half that of **P1**-based cells. The charge recombination kinetics, probed by considering the charge transfer resistance at the NiO/electrolyte interface at a comparable chemical capacitance, showed that the dyes behaved similarly under this respect and that the higher photovoltage observed for the device based on **SK4** dye is seemingly due to a positive shift of the valence band edge, consistent with the shift in the anodic threshold measured in the dark. The fact that similar recombination resistances were found at a comparable photohole density in the NiO corroborates the indication, gained by DFT studies, that the superior performance, particularly in photocurrent, of **P1** may be ascribed to a superior charge separation capability originating by its smaller exciton binding energy. Future designs of dyes for *p*-type sensitization will take this parameter into consideration for achieving a more effective charge separation.

**Supplementary Materials:** CV plot, normalized absorption and emission spectra, DFT isodensity plots of HOMOs and LUMOs, HOMO-LUMO energy gaps and excition binding energies, dark currents and EIS spectra of p-DSSC, <sup>1</sup>H and <sup>13</sup>C NMR spectra of intermediates and dyes. The following are available online at www.mdpi.com/1996-1073/9/1/33/s1.

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