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Beneficial Effect of Electron Withdrawing Groups on the Sensitizing Action of Squaraines for \( p \)-Type Dye Sensitized Solar Cells

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Abstract

The three squaraines (SQs) here indicated as VG1-C8, VG10-C8, VG11-C8 (Figure 1) have been synthesized and employed as dye sensitizers of $p$-type dye-sensitized solar cells ($p$-DSCs) when screen-printed NiO is the nanoporous photocathode. The photoelectrochemical performance of the three differently sensitized $p$-DSCs is compared in terms of the structural diversities among the three SQs. The $p$-DSC with VG11-C8 as sensitizer, i.e. the sole SQ with a dicyano–vinyl substituent on the central squaric ring, results the most efficient device because of the highest values of overall conversion efficiency ($\eta > 0.04 \%$ for VG11 vs $\eta < 0.02 \%$ for VG1 and VG10). Moreover, VG11 is the sole SQ of the C8 series which exerts a favourable synergistic effect on the external quantum efficiency (EQE) of screen-printed NiO. The anti-aggregating agent CDCA (= chenodeoxycholic acid) in the sensitizing solution of NiO is beneficial on the performance of the corresponding $p$-DSCs when the ratio [CDCA]/[dye] is 1:10 for all three SQs. Electrochemical impedance spectroscopy (EIS) of the SQs sensitized $p$-DSCs under illumination reveals that the most performing photoelectrochemical cell, i.e. the one with VG11 sensitizer, is characterized by relatively lower resistances of charge transport (~ 30 Ω) and interfacial recombination (~ 90 Ω), shorter hole diffusion times (~ 2 ms), and larger hole diffusion coefficient (~ $2*10^{-7}$ cm$^2$ s$^{-1}$) with respect to the $p$-DSCs with VG1 and VG10 dyes.
1. Introduction

As alternative to the photovoltaic technologies based on \( pn \) junctions the photoelectrochemical dye-sensitized solar cells (DSCs), or Grätzel cells,\(^3\) have received the greater attention. These can be divided into \( n \)- and \( p \)-type DSCs when oxidation and reduction are respectively the photoactivated electrochemical processes.\(^4\)\(^{13}\) The \( n \)-DSCs are comparatively better than the \( p \)-type counterparts because \( n \)-DSCs are capable to reach efficiencies of about 14%\(^8\) whereas \( p \)-DSCs\(^{15-19}\) display values still below 2%\(^{20-23}\) at their best \([\eta = 1.67\%]\) when NiO is the photocathode, PMI-6T-TPA the dye-sensitizer and Co(II)/Co(III) the redox couple\(^24\)\(^{25}\). Current research efforts on \( p \)-DSCs are dedicated to the preparation of photocatodes with large surface area thus affording high dye loading and large current flows \( via \) the exploration of new and improved deposition methods of the photoelectrode\(^{26-30}\) (typically NiO).\(^{15-19}\) Another important line of research is directed towards the development of dye-sensitizers appositely designed for DSC photocathodes with the aim of ameliorating the performances of the corresponding \( p \)-DSCs. When designing dye-sensitizers for \( p \)-DSCs\(^3\) and eventually tandem DSCs (\( t \)-DSCs)\(^32\) with anodes and cathodes both photoelectrochemically active, the following key requirements have to be fulfilled: (i) broad absorption spectrum comprising visible and NIR with high extinction coefficient; (ii) highest-occupied molecular orbital (HOMO) energy of the sensitizer lying below the valence band (VB) upper edge of the \( p \)-type semiconductor; (iii) lowest-unoccupied molecular orbital (LUMO) energy of the sensitizer placed above the redox potential of the electrolyte to ensure efficient hole injection into the semiconductor VB and regeneration from the redox species; (iv) anchoring groups to link strongly the dye on the oxide surface and allow electronic coupling between the Fermi level of the semiconductor and the HOMO of the dye; (v) electron withdrawing groups located far away from the semiconductor surface in order to minimize recombination phenomena and accelerate electron transfer (\( et \)) from the excited dye to the redox shuttle. In particular, it has been the last issue which has motivated the synthesis of the squaraine denominated VG11 (Figure 1, \textit{vide infra}) here proposed for the first time for \( p \)-DSCs.\(^{33-37}\) In fact, organic dyes like SQs\(^{33-37}\) can be particularly
suitable for photocathodes sensitization of \( p \)- and mostly \( t \)-DSCs since in the latter device the red shifted absorption of the SQs would complement the spectrum of the traditional organometallic and organic dyes for \( n \)-DSCs. Among NIR-absorbing dyes, SQs are particularly important due to high absorption coefficient in the NIR (> 550 nm, Figure ESI1), and the tunability of the structure and electronic properties. Some successful employment of SQs in \( n \)-DSC has been reported in the recent past, but few examples of SQs have been actually designed on purpose for NiO based \( p \)-DSCs. In the present contribution we synthesized the three symmetrical squaraines VG1-C8, VG10-C8 and VG11-C8 (Figure 1) for their employment as dye-sensitizers of screen-printed NiO photocathodes for \( p \)-DSCs.

![Figure 1](image)

**Figure 1.** Structures of the three SQs VG1, VG10 and VG11 (C8 series) employed as sensitizers of \( p \)-DSCs with screen-printed NiO photocathodes.

Due to their symmetric structure the three SQs of the C8 series present easiness of preparation, simpler and cheaper procedures of purification in comparison to their unsymmetrical analogues. In particular, VG1 and VG10 were in precedence employed by us in \( p \)-DSCs when NiO cathode was prepared via screen-printing with \( \text{CH}_3\text{CO}_2\text{H} \) as anti-aggregating agent, whereas VG11 is
reported here for the first time as sensitizer of a $p$-DSCs. The three SQs differ for the extent of electronic conjugation and the electronic character, i.e. if electron-withdrawing or releasing, of the substituents on the central squaric ring. VG1, VG10 and VG11 possess alkyl chains of the same length (C8) to exclude a priori differences of sensitizing action which are ascribable to these non electroactive groups. Therefore, the main purpose of the present paper is to provide new directions in terms of SQ structural design for improving the still weak performances of the NiO-based $p$-DSC with SQ sensitizers. SQs generally suffer of intermolecular aggregation in the surface anchored state due to their large size and the presence of the flat squaric central unit. Such phenomena are generally undesired in DSCs since they lower the rate of dye photoexcitation with consequent inhibition of photocharge injection. To prevent these, we have considered the use of the co-adsorbent CDCA (chenodeoxycholic acid) during the preparation of the sensitizing solutions with VG1, VG10 and VG11. Erythrosine B (ERY) has been also utilized as benchmark sensitizer for comparative purposes. The analysis of the kinetics of electron transfer processes in these photoelectrochemical devices differing for the nature of the SQ sensitizer will be given employing EIS.

2. Experimental

2.1 Screen-printing paste preparation

The preparation of the paste, i.e. the precursor utilized successively in the process of screen-printing deposition of NiO, has been realized through the procedure described schematically in Table ESI1. This procedure is a slightly modified version of the one reported by Ito et al. for the preparation of the paste to be employed in the screen-printing deposition of TiO$_2$ for $n$-DSCs. All chemicals were purchased from Sigma-Aldrich or Fluka at the highest degree of purity available and were used without any further purification. Anhydrous terpineol was used as a mixture of enantiomers. Grinding was realized at room conditions in a mortar with 15 cm of diameter. Stirring process was carried out with a 4 cm long magnetic stirrer at the rotation speed of 250 rpm. Ultrasonic
homogenization was performed with a Ti horn-equipped sonicator (Vibracell 72408 from Bioblock scientific). The paste we obtained had the appearance of a viscous slurry with a shelf-life of about one year. The originality of the procedure of paste preparation reported in Table ESI1 resides on the utilization of preformed NiO microspheres (max diameter = 50 nm) instead of NiO chemical precursors such as NiCl₂ or Ni(OH)₂ the use of which renders generally more elaborated the procedures of purification and requires more drastic conditions of sintering.

2.2 Screen-printing deposition of NiO thin films

The paste precursor was spread over a FTO-covered glass panel (item no. TCO227 from Solaronix), which has been previously cleaned in ultrasonic bath with acetone for 10 minutes and with ethanol for 10 minutes. The paste was spread via screen-printing through a 90.48 T mesh screen on dried FTO/glass panels. After a pre-drying period of 15 minutes at 100°C in oven (WHT 5/120 from Welland), the temperature of the oven was increased with a temperature ramp of 15°C/min till 450 °C. The screen-printed slurry of NiO nanoparticles was kept at the highest temperature for 30 minutes to complete sintering. After sintering the resulting mesoporous film of NiO was cooled down to ambient temperature with the attainment of a 2 μm thick film of NiO. The thickness (l) of the NiO film was estimated with a Dektat 150R profilometer from Veeco. The imaging of the surface morphology of mesoporous NiO film (Figure 2) was realized with a Scanning Electron Microscope (SEM, Auriga Zeiss 170 instrument). The advantage of such a procedure is the lower temperature of processing (≤ 400–450 °C) with respect to sol–gel method (> 400-500 °C). The presence of pre-formed nanoparticulated NiO in the starting paste forces the heat to sinter solely the metal oxide nanoparticles with no need of activating chemical reactions for metal oxide formation. At the adopted temperature of sintering the cellulose-based binders of the paste are completely combusted and removed as a volatile product of thermolysis from the oxide layer. The resulting oxide in the pristine state generally presents non stoichiometric features for the co-presence of
nickel sites with valence II and III. For this reason it would be more appropriate to report it with the formula NiO$_x$.

Figure 2. SEM picture of the surface morphology of mesoporous NiO thin film ($l = 2 \mu m$) deposited via screen-printing.

2.3 Synthesis of squaraines

The details of the synthesis of VG1-C8, VG10-C8 and VG11-C8 have been previously reported by our groups in the studies presented in refs.33 and 67.

2.4 p-DSC assembly and testing

Screen-printed NiO photocathode and the platinised FTO counter electrode were assembled together in a sandwich configuration using as a Bynel® thermoplastic polymeric film having the dual function of spacer and sealant. Platinized counter electrodes were prepared by screen printing platinum paste Ch01 from Chimet through a 100T mesh screen onto FTO-coated glass.$^{68}$ Sensitization of screen-printed NiO photocathodes was realized by dipping the NiO films in a 0.2 mM dye-sensitizer solution (either with or without CDCA) using ethanol as solvent. All screen-printed NiO samples were sensitized at room temperature for 2 hours. After sandwiching the electrodes, the triiodide/iodide based electrolyte [commercial Iodolyte Z-150 from Solaronix, Aubonne (CH)] was injected inside the cell by means of the vacuum backfilling
technique. The hole for electrolyte injection was sealed with a liquid resin [TB3035B from ThreeBond, Tokyo (JP)], which becomes a hard paste upon UV curing treatment. The photoactive area of the samples was 0.25 cm².

Optical transmittance was measured with a double ray spectrometer [UV-2550 by Shimadzu, Kyoto (JP)]. Photoelectrochemical performances consisting in the determination of the characteristic curves current-potential ($JV$) were evaluated using the solar simulator Solar Test 1200 KHS (class B) at 1000 W m⁻² with artificial solar spectrum AM 1.5 G. The curves of IPCE (incident photon to current conversion efficiency, see ESI for fundamental photoelectrical equations) were recorded using a computer controlled set-up consisting of a Xe lamp (Mod.70612, Newport) coupled to a monochromator (Cornerstone 130 from Newport), and a Keithley 200 2420 light-source meter.

The EIS measurements were performed with an AUTOLAB PGSTAT12® from Metrohm at the condition of open circuit potential in dark conditions and under solar simulator illumination. For the sole EIS experiments conducted under illumination the applied potential was modulated within ± 20 mV with respect to the value of open circuit potential. This was made in order evaluate the variations of all key parameters characterizing the equivalent circuit fitting the electrochemical impedance spectra (vide infra). In the EIS experiments the potential perturbation had an amplitude of $10^{-2}$ V. The frequency range of the impedance perturbation varied depending on the state of the cell, i.e. if illuminated or not. Impedance spectra were recorded within the frequency ranges $10^{-1}$–$10^5$ Hz and $10^{-2}$–$10^5$ Hz when the cell was under illumination and in dark conditions, respectively. Electrochemical impedance spectra were fitted using Z-View software from Scribner Associates Inc.. In the context of impedance spectra simulation of NiO based $p$-DSCs we developed a new equivalent circuit adapting the transmission line model first proposed Bisquert et al. for the impedance analysis of TiO₂-based $n$-DSCs with gel electrolytes. 
3. Results and discussion

3.1 Optical characterization of sensitized NiO

The absorption spectra of VG1, VG10 and VG11 in ethanol at room temperature (see Figure ESI1) indicate 643 nm, 671 nm and 710 nm as values of $\lambda_{\text{MAX}}$ for VG1, VG10 and VG11, respectively, with $\Delta(\lambda_{\text{MAX}})$ progressively decreasing upon increase of $\lambda_{\text{MAX}}$ in the triad of chosen dyes. The red shift observed in passing from VG1 to VG10/VG11 is related to the larger extent of electronic conjugation in passing from the indoleninic unit of VG1 to the benzoindoleninic unit of VG10/VG11 (Figure 1), whereas the further red shift observed on going from VG10 to VG11 is ascribed to the presence of the dicyano-vinyl cromophoric unit. The resulting effect of sensitization of the NiO cathode (Figure 3) reflects the optical absorption differences already observed in the solution spectra of VG1, VG10 and VG11 (see Figure ESI1).

![Figure 3](image-url)

**Figure 3.** Transmittance variation of screen-printed NiO in passing from the pristine to the sensitized state with the C8 series of the three SQs under consideration (Figure 1), and with ERY as comparative dye.

The efficacious sensitization of screen-printed NiO is a consequence of its open and scale-like morphology (Figure 2), which is characterized by cavities and apertures having a linear size in the order of few hundred of nanometers. Such voids warrant the wettability of NiO by the solution of
sensitization as well as liquid electrolytes throughout the whole thickness of the oxide film. As main consequence, this morphological feature brings about the observation of the direct proportionality between layer thickness and the extent of electrical current exchanged by the NiO electrode when the latter constitutes the actual electroactive species. When CDCA (Figure ESI2) is added to the solution of sensitization, the corresponding variations of NiO optical transmittance (Table ESI2) are directly correlated to the relative concentration of the dye-sensitizer with respect to CDCA (Figure 4). Upon increase of anti-aggregating agent concentration in the sensitizing solution, the amount of chemisorbed CDCA correspondingly increases with consequent diminution of the quantity of chemisorbed dye as evinced by the increase of transmittance ion correspondence of the characteristic absorption peaks of the colorants (Figure 4 and Table ESI2). It is here anticipated that for these SQs a higher dye loading is not necessarily associated to an improvement of the photoelectrochemical performance in the corresponding p-DSC, as previously observed with analogous systems, and an optimum concentration ratio does exist (vide infra). At the maximum concentration of CDCA (200 mM) employed by us in the sensitization solution, the transmittance spectrum of VG1- and VG10-sensitized oxide does not indicate any optical evidence of the presence of anchored sensitizer (Figure 4, top and middle frames). This is not the case of VG11-sensitized NiO the spectrum of which shows the typical absorption features of VG11 [Figures 3 and 4 (bottom frame)]. A possible reason at the basis of such observed differences could be due to a different geometry of anchoring for the diverse SQs for which calculations in the gas phase have predicted that VG11 possesses a trans form with respect to the bonds linking the squaraine core to the benzoindoleninic moieties (Figure 1) as most stable configuration, while VG1 and VG10 have the cis form with respect to the same type of bonds as most favourable configuration. As a consequence of these theoretical considerations we expect that the geometry of chemisorption for the three SQs under investigation follows the two main schemes presented in Figure 5. According to this depiction the geometry of anchoring of one molecular unit of trans
VG11 does not impede the occupancy of the vicinal sites of anchoring to a second molecule of \textit{trans} VG11.

\textbf{Figure 4.} Transmittance spectra of SQ-sensitized NiO films when different concentrations of CDCA were used in the sensitizing solution. Dyes: VG1 (top), VG10 (middle) and VG11 (bottom).
The red trace refers to the spectrum of NiO sensitized in CDCA-free solution. The orange profile refers to the spectrum of pristine NiO. The ratios in the insets indicate the concentration ratio [dye]/[CDCA] in the solution of sensitization.

![Diagram of anchoring sites and molecules](image)

**Figure 5.** Expected geometry of SQs chemisorption on NiO for the most stable configurations of VG1, VG10 and VG11. The differently colored boxes indicate the sites of anchoring on NiO surface. For sake of simplicity we have omitted the second carboxylic unit of VG1 and VG10 systems, which is employed for molecular anchoring.

This would keep a relatively higher surface concentration of the anchoring sites actually accessible to the other molecules of *trans* VG11 (perpendicularly oriented with respect to NiO surface) in comparison to the situation with *cis* VG1 and *cis* VG10. In fact, upon chemisorption the two latter systems are expected to prevent further occupancy of the vicinal sites of anchoring because of their hypothetical parallel orientation with respect to the NiO surface as sketched in Figure 5. It is also expected that the process of *trans* VG11 chemisorption is favoured kinetically with respect to the chemisorption of *cis* VG1 and VG10 (Figure 5). This is because the rate of dye chemisorption (a heterogeneous process of second order) is directly proportional to the product [SQ]·[NiO]surf, where [NiO]surf represents the surface concentration of the NiO anchoring sites available to the dye.
The parameter \([\text{NiO}]_{\text{s}}\) varies with the nature of the SQ and its geometry of chemisorption, and is supposed to be larger for trans VG11 with respect to cis VG1 and VG10 by virtue of the given hypothetical geometry of dye anchoring (Figure 5). Under the conditions of sensitization with molar ratio SQ/CDCA = 1/1000, the SQs with slower kinetics of chemisorption, i.e. VG1 and VG10 (vide supra), apparently are not fast enough to be chemisorbed by the NiO surface because of the relatively very large concentration of CDCA which occupies in a shorter time all the anchoring sites available on the NiO surface. This hypothesis would explain the lack of any evidence of SQ optical signature in the spectrum of NiO electrode sensitized at the condition \([\text{SQ}]/[\text{CDCA}] = 1/1000\) with SQ = VG1 and VG10 (Figure 4, top and middle plots). In contrast, VG11 would manage to get anchored on NiO surface also from the solution more highly concentrated in CDCA prior to the saturation of NiO surface sites by CDCA because of the faster kinetics of VG11 chemisorption with respect to the analogous processes of VG1 and VG10. At the moment we are not in possess of a clear experimental evidence of the actual molecular orientation of this series of SQs on the NiO surfaces for p-DSCs. This type of studies can be conducted with XPS and/or ATR-FTIR techniques which, in turn, require a flat surface morphology as well as exposed single crystal faces. As previously evidenced, the sensitized NiO samples here investigated are polycrystalline and present mesoporous morphology (Figure 1) with no preferred surface orientation. These structural and morphological features somewhat limit the meaningfulness of such surface studies that should be apt to provide an indication of the molecular orientation for these complex sensitized systems. The anchoring of CDCA on mesoporous NiO cannot be considered innocent in optical terms: NiO films immersed in highly concentrated solutions of CDCA and in absence of dye-sensitizer presented a spectrum with lower optical transmission than pristine NiO within the window 330-650 nm (not shown). Moreover, the observed shifts of transmission minima for the SQ-sensitized spectra obtained from the sensitization solutions with the largest concentrations of CDCA (see Table ESI2) could be the consequence of dye-protonation (halochromic effect of SQ dyes).
3.2 Photoelectrochemical characterization of SQ-sensitized p-DSCs

The characteristic $JV$ curves of the $p$-DSCs differing for the conditions of NiO photocathode sensitization are presented in Figure ESI3. The corresponding parameters of short circuit current density ($J_{SC}$), open circuit potential ($V_{OC}$), fill factor (FF) and overall efficiency ($\eta$) are reported in Table 1. The conditions of NiO electrode sensitization with ERY colorant have been previously reported in refs. 51 and 52.

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<th>Dye</th>
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<th>$V_{OC}$ (mV)</th>
<th>$J_{SC}$ (mA/cm$^2$)</th>
<th>FF (%)</th>
<th>$\eta$ (%)</th>
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Table 1: Comparison of the main photovoltaic parameters obtained from the $JV$ curves of the $p$-DSCs with bare NiO photocathode and differently sensitized NiO electrodes (see Figure ESI3). The effect of CDCA concentration in the sensitization solution on the corresponding $p$-DSC performance with SQ dyes is shown for four predefined values of the molar ratio CDCA:SQ in the solution of sensitization. ERY is used as benchmark sensitizer for comparative purposes.

From the $JV$ data summarized in Table 1, it results that the $p$-DSCs with the largest overall efficiencies were those sensitized with VG11 ($\eta = 0.043 \%$) and the dye-benchmark ERY ($\eta = 0.032 \%$). VG1 and VG10 as photocathode sensitizers inhibited the intrinsic photoelectrochemical activity of screen printed NiO in the bare state as evinced from the decrease of the overall efficiencies (0.020 % for bare NiO vs 0.018% of VG1- and VG10-sensitized NiO, Table 1). This trend could be ascribed to a photoinjection not sufficiently sustained by excited VG1 and VG10.
with respect to pristine NiO, Dye anchoring thus produces a counterproductive effect. The sole colorant that does not exert an inhibiting effect on the photoelectrochemical activity of bare NiO resulted the system VG11. To our knowledge this is the first case of a SQ sensitizer which does not produce the inhibition of substrate photoactivity when a NiO based $p$-DSC is analysed.

When the overall efficiency of the series of VG sensitized $p$-DSCs is analyzed, the most performing set of cells resulted the one that adopted the concentration ratio $[\text{CDCA}]:[\text{SQ}] = 10:1$ in the solution of sensitization no matter of the nature of the dye (Table 1). The existence of an optimal CDCA/SQ ratio for the effective inhibition of deleterious intermolecular interactions between the molecules of anchored dye could be a consequence of the existence of an allotted number of surface sites for the chemisorption of non (or poorly) interacting dye-sensitizers. This surface dependent effect would explain why larger values of the $[\text{CDCA}]:[\text{SQ}]$ ratio would drastically decrease the amount of anchored dye whereas the total lack of disaggregating agent in the sensitizing solution would not impede at all the creation of dye aggregates on the electrode surface. The existence of such aggregates is recognized as responsible of a photophysical behaviour that would follow altered and probably less efficient mechanisms of photoexcitation, photoinjection and dye regeneration with respect to non interacting dye molecules in the anchored state. Another possible influence of CDCA on the photoelectrochemical behaviour of SQ based $p$-DSCs could be related to the acidic nature of the disaggregating agent that can modify the state of charge of the colorant ($\text{vide supra}$). The protonation of the dye generally represents a beneficial event in the context of photoelectrochemical reduction since the positive charge of the dye is expected to promote the electron transfer from the valence band of NiO to the protonated dye thus realizing the process hole injection at a presumably higher rate with respect to a non protonated dye. The latter statement derives from elementary electrostatic considerations. The most efficient VG-based photoelectrochemical cells presented also the highest values of $J_{SC}$ values (Table 1), whereas FF values remained stable around 35-40% (as typically found for NiO based $p$-DSCs), and were thus poorly dependent on the amount of CDCA in the sensitizing solution (Table 1). The
parameter $V_{OC}$ tended to decrease as CDCA concentration increased in the sensitizing solution (Table 1). This fact indicated the direct proportionality between the open circuit photovoltage and amount of dye actually present on the electrode surface. It is worth to notice that the VG10-sensitized photoelectrode presented the generally largest values of $V_{OC}$, i.e. a parameter that is related principally to the lifetime of the holes once photoinjected in the oxide electrode. To the larger value of open circuit potential should correspond the larger surface concentration of photoinjected charges (and not necessarily to larger a dye-sensitizer surface concentration) and the smaller rate of recombination between photoholes and the photoproduct of reduction (I in the present case) because of the longer hole lifetime on the NiO surface. The reason why VG10 produces a larger value of $V_{OC}$ than VG1 is found in the larger size of the conjugated moiety present in VG10 with respect to VG1 (benzoindoleninic vs indoleninic unit). The larger benzoindoleninic unit in VG10 would than act as a more efficient spacer between the hole localized on the surface electrode and the site of recombination where iodide is present. The anion iodide is expected to be either directly absorbed on the positively charged NiO electrode surface, or in the electrolyte side of the electrode/electrolyte interface. Therefore, the dye in this case would act as a separator between hole and iodide in function of its molecular extension. This, in turn, would reduce the time-probability with which the hole recombines with the reduced form of the redox mediator and leads to the resulting increase of $V_{OC}$. This consideration is based on the assumption that all SQs here considered anchor through both carboxylic groups present in the structure. The given explanation founded on the different molecular sizes of VG1 and VG10 sensitizer does not longer hold when the trend of $V_{OC}$ refers to the comparison of VG10- and VG11-based photoelectrochemical cells for the supposition we made on the geometry of molecular anchoring shown in Figure 5. At the basis of the smaller value of $V_{OC}$ for the VG11-sensitized $p$-DSC (Table 1) it is believed that the presence of the strong electron-withdrawing dicyano-vinyl group plays a relevant role. As a matter of fact, an electron-attracting group tends to recall electrostatically the photoinjected positive charge in absence of a current flow which, in turn, would displace the
injected holes by diffusion towards the site of charge collection at the FTO/NiO interface. The attraction exerted by the dicyano-vinyl group on the photoinjected positive hole will favour kinetically the process of recombination between hole and iodide in absence of a current flow with respect to the situation occurring with the VG10- sensitized p-DSC. The best performing SQ p-DSCs, i.e. those obtained from sensitization solutions with [CDCA]:[dye] = 10:1, were compared with the photoelectrochemical cell utilizing the benchmark sensitizer ERY (Figure ESI3). The VG1 and VG10 showed lower efficiencies with respect to ERY mainly because of the lower $J_{SC}$ (Table 1). The sole SQ-based cell that presented larger and more efficient parameters of photoconversion with respect to the ERY-sensitized cell was the one with VG11 sensitizer. The efficiency of 0.043% reached by the VG11-sensitized p-DSC represents a record value for the NiO based p-DSCs with symmetrical SQ sensitizers when $I_3^{-}/I^{-}$ is the redox shuttle and SQ constitutes the sole chromophoric unit. This finding is quite encouraging because it evidences how determining has been the design of a SQ like the VG11 here proposed for NiO p-DSCs to obtain augmented conversion efficiencies and superior photoelectrochemical performances.

The complete set of the IPCE spectra for the differently sensitized p-DSCs has been presented in Figure ESI4. From the latter a selection of the IPCE spectra produced by the best performing p-DSCs with all the dye-sensitizers here considered has been made. These spectral profiles are shown in Figure 6.
The curves of photoelectrochemical spectral efficiency (Figures ESI4 and 6) are consistent with the observed trends of overall efficiencies and $J_{SC}$ determined by the characteristic $J\!V$ curves (Figure ESI3 and Table 1). Moreover, the peaks of EQE corresponded to the absorption peaks obtained from the conversion of the transmittance data presented in Figure ESI1 into absorption data through the relationship $A(\lambda) = \log [1/T(\lambda)]$ being $T \leq 1$. The use of the most efficient sensitizer, i.e. VG11, brings about two main consequences on the IPCE spectrum of the corresponding $p$-DSC: i) induces photoelectrochemical activity in the spectral range of VG11 optical absorption, as expected for a normal completion of the function of sensitization (Figures 3 and ESI1); ii) it does not inhibit the intrinsic photoelectrochemical activity of bare NiO centred at about 370 nm (Figure 6). Therefore, dye-sensitizers VG1, VG10 and ERY actually behave as agents of photoelectrochemical passivation towards NiO substrate itself. When high concentration of CDCA are employed (practically for [CDCA] > 0.01 M) in the solution of NiO pseudo-sensitization, i.e. no colorant is present, a dramatic decrease of the characteristic peak of bare NiO photoelectrochemical activity in
the near UV (Figure 6) is observed in the relative IPCE spectra (Figure ESI5) and a general worsening of the photoelectrochemical performance of NiO is observed in the associated $JV$ curves (Figure ESI6).

In the case of NiO sensitization with VG10 it was clear that the intrinsic photoelectrochemical activity of NiO in the near UV dropped more sensibly for the photocathode treated with the sensitizing solution having the largest ratio $[\text{CDCA}]:[\text{VG10}]$ ( = 1000:1). This is an indication of the fact that the SQ dyes present a generally higher affinity of anchoring onto NiO surface with respect to the anti-aggregating agent CDCA. Beside the occupancy of the allotted sites of anchoring (Figure 5) , the CDCA would play also a non negligible role of passivating agent towards the photoelectrochemical activity of NiO (Figures ESI5 and ESI6).

The IPCE spectra obtained with the NiO-based $p$-DSCs here presented which utilize triiodide/iodide as redox couple (Figures ESI4 and ESI5), include contributions from the photoexcitation of different light-absorbing species that act (either in a synergistic or in independent fashion) in quite distinct portions of the spectrum. Upon determination of the IPCE spectra of VG11-sensitized $p$-DSC with sensitization molar ratio $[\text{CDCA}]:[\text{SQ}] = 100:1$ in both back and front modalities of illumination (see Figure ESI7) the comparison of the IPCE profiles allows the distinction of the various contributions to the photocurrent of reduction due to the different species with specific light-absorbing properties: triiodide in the range 350-400 nm, NiO in the near UV range 300-480 nm, and VG11 in the wavelength range 580-800 nm (Figures ESI4 and ESI7) taking into account the specific optical absorption properties of the triiodide/iodide-based electrolyte (Figure ESI8). We stress here the importance of NiO as main photoelectrochemically active species in this type of photoconversion devices for the presence of superficial Ni(III) centres as defective and photoactive sites of the oxide surface. It is widely acknowledged that these defective centres in non stoichiometric nickel oxide originate the near UV absorption displayed by bare nickel oxide (see Figure 3, orange trace) to which is associated the observed photoelectrochemical activity in the near UV, no matter of the nature of the redox shuttle. With these considerations we are
prone to minimize the importance of the triiodide species with respect to NiO as main responsible of the photoelectrochemical activity detected with NiO-based $p$-DSCs. Another finding which supports the relative importance of NiO vs triiodide as main photoelectrochemically active species in the near UV is the observation of a strong dependence of the photoelectrochemical activity of NiO when this is treated with CDCA. It has been previously outlined how the overall conversion efficiency and the IPCE of the $p$-DSCs employing non colored NiO samples that have been previously treated with CDCA decrease heavily upon increase of CDCA concentration in the sensitization solution (Figures ESI5 and ESI6). This effect of CDCA has been ascribed to the photoelectrochemical passivation of the active Ni(III) centres by such an anti-aggregating agent. In doing so, we believe that CDCA behaves as a blinding agent for the photoactive Ni(III) centers on the oxide surface. If the main photoexcitation process at the basis of the origin of the observed cathodic photocurrent in NiO-based $p$-DSCs was the activation of triiodide, the phenomenon of CDCA passivation of the NiO surface would have not produced such a strong effects of photoinhibition. Moreover, the comparison of the IPCE profiles obtained from back and front illumination (Figure 6 and Figure ESI7), would have not been characterized by the most profound differences in correspondence of NiO characteristic absorption (Figure 4), even in absence of CDCA treatment.

3.3 Electrochemical characteristics of SQ-sensitized $p$-DSCs

Dark $JV$ curves measurements were conducted on the most efficient photoelectrochemical cells sensitized with SQs at the molar ratio of sensitization [CDCA]:[SQ] = 10:1 (Table 3), on the ERY sensitized $p$-DSC and on the $p$-DSC with bare NiO cathode (Figure ESI9 and Table ESI3). The dark current of the $p$-DSCs originated from a reduction process being the current values with negative sign, and refer to the dark reduction of triiodide to iodide $I_3^- + 2e^- \rightarrow 3I^-$. This occurs at the dye-NiO/electrolyte interface and, eventually, at the FTO/electrolyte interface. The FTO/electrolyte interface intervenes in the electrochemical process if there is no compact layer of
NiO or any insulating layer like Al₂O₃, which would prevent the phenomenon of shunting between FTO substrate and the electrolyte. In the present work, we have not considered the deposition of a compact layer of NiO for the prevention of such a phenomenon and it is planned to consider this type of modified NiO photocathodes in a successive study. Under these circumstances shunting can take place at the FTO/electrolyte interface if the first layer of nanoporous NiO in contact with FTO does not cover uniformly the transparent conductive substrate.

Except VG11, in dark conditions all dyes led an increase of the dark open circuit voltage in the corresponding p-DSC with respect to cell with bare NiO cathode. This implies that part of the surface charge present in the sensitized cathode originates either from a process of dark charge transfer between the dye and NiO substrate, or from a process of anion adsorption the extent of which depends on the nature of the anchored sensitizer, its surface concentration and the nature of the adsorbed anion. The analysis of the dependence of dark $J_{SC}$ on the nature of the sensitizer (Table ESI3) revealed that only the sensitization with ERY produces cells with a larger value of dark $J_{SC}$ with respect to the p-DSC with bare NiO cathode (Figure ESI9). This implicates that within the C8 series here investigated none of the three SQs actually mediates the process of dark electron transfer from NiO to triiodide species. Among SQs only VG11 behaved as an agent of passivation for NiO towards the dark reduction process $I_3^- + 2 e^- \rightarrow 3 I^-$ as evinced from the net decrease of the dark $J_{SC}$ in passing from bare NiO to VG11-sensitized p-DSC (Table ESI3). This indicated that VG11 does not intervene directly as a mediator during such a cathodic process in dark conditions, i.e. in absence of any photoexcitation, and prevents the dark electrochemical reduction of triiodide by NiO. Therefore, triiodide reduction in dark conditions involves solely the intervention of NiO (and eventually the one of FTO exposed to the electrolyte). Conversely, the doubling of dark $J_{SC}$ in passing from the electrochemical cell with bare NiO cathode to its ERY-sensitized version evidences the participation of ERY sensitizer as surface-immobilized redox mediator during the process of dark reduction of the triiodide species. 


3.4 *Electrochemical impedance spectra of SQ-sensitized p-DSCs*

The electrochemical impedance spectra of the most performing SQ-sensitized *p*-DSC and the photoelectrochemical cell utilizing ERY colorant have been recorded. These are shown in Figure 7. In the determination of the EIS profiles we have not considered the impedance analysis of the *p*-DSC with VG1-C8 sensitizer due to its poorer performance in comparison to the photoelectrochemical cells employing the other two SQs and the benchmark dye ERY.

*Figure 7.* Nyquist plots of the electrochemical impedance spectra of the *p*-DSC sensitized with ERY (red dots), and the best performing SQs VG10 (light green dots) and VG11 (dark green dots) at the respective values of open circuit potential. $V_{OC} = 90, 100$ and $90$ mV for ERY-, VG10- and VG11-sensitized photoelectrochemical cells, respectively.

For the analysis of the EIS data in Figure 7 we used an equivalent circuit that represents an adaptation of the one developed by Bisquert *et al.* for the EIS analysis of TiO$_2$-based *n*-DSCs (Figure 8).
Figure 8. Equivalent circuit employed for the interpolation of the EIS data in Figure 7.

The equivalent circuit here proposed consists of three main elements which are connected in series: i) the series resistance \( R_s \) that collects all resistive contributes arising from the electrical contacts, the external circuit and the resistance of the FTO substrate of the photoelectrode; ii) the interfacial element referring to the electron transfer process at the counterelectrode/electrolyte interface and defined as the parallel combination of the charge transfer resistance \( R_{CE} \) through the interface and a capacitance \( C_{CE} \) of the electrochemical double layer; iii) the element of transmission line (TL) which comprises diffusion-recombination effects within the dye-sensitized photoelectrode and the capacitance associated to the sensitized NiO/electrolyte interface under illumination. The TL element takes into account the mesoporous nature of the semiconducting NiO layer, its thickness and its hole transport properties. The latter properties are limited by the phenomenon of recombination and are controlled by the phenomenon of diffusion. Diffusion, in turn, originates from the existence of a chemical capacitance, i.e. a charge concentration gradient, within the mesoporous layer of NiO as customarily assumed for the photoactive electrodes of DSCs with typically discontinuous electronic structure. The EI spectra have been determined within a range of applied potential centred at \( V_{OC} \) and were varied ± 20 mV around \( V_{OC} \) (vide infra). This procedure was followed with the aim of evaluating the response of the devices under changing operative conditions. The fitting curves obtained with the equivalent circuit of Figure 8 are shown in Figure ESI10. In the TL element of Figure 8, i.e. the circuital element referring to the electrochemical processes localised on the photoactive NiO cathode, the parameters of hole transport resistance \( R_t \) through the NiO layer, the photocharge transfer resistance across the NiO/electrolyte interface
(R_{rec}), and the chemical capacitance $C_{\mu}$ are enclosed. The latter is related to the capacitance $C_{\text{CPE}}$ of the CPE (Constant Phase Element) via the relationship $C_{\mu} = (C_{\text{CPE}}*R_{\text{rec}})^{1/n}/R_{\text{rec}}$ the capacitance $C_{\text{CPE}}$ and the dimensionless parameter $n$ being two fitting parameters. Fitting curves were evaluated with the $\chi^2$-test that gave a value smaller than $2*10^{-4}$.

<table>
<thead>
<tr>
<th>DYE</th>
<th>$R_t / \Omega$</th>
<th>$R_{\text{rec}} / \Omega$</th>
<th>$C_{\mu} / \mu\text{F}$</th>
</tr>
</thead>
<tbody>
<tr>
<td>ERY</td>
<td>29.2 ± 0.7</td>
<td>124.2 ± 1.1</td>
<td>220 ± 4</td>
</tr>
<tr>
<td>VG10-C8</td>
<td>48.8 ± 2.5</td>
<td>264.9 ± 8.5</td>
<td>66 ± 3</td>
</tr>
<tr>
<td>VG 11-C8</td>
<td>28.6 ± 2.4</td>
<td>85.8 ± 6.8</td>
<td>60 ± 8</td>
</tr>
</tbody>
</table>

**Table 2.** Values of the fitting parameters $R_t$, $R_{\text{rec}}$ and $C_{\mu}$ contained in the TL element of Figure 8 for the simulation of the electrochemical impedance spectra determined at open circuit potential (Figure 7).

The transport resistance $R_t$ follows approximately a trend opposite to the one of the overall efficiency (see Table 1). This is equivalent to say that the $p$-DSC with the largest value of $R_t$ presents the lowest value of $\eta$ and viceversa. This correlation combines with the direct proportionality which has been observed between $\eta$ and $J_{\text{SC}}$, the latter parameter being inversely proportional to $R_t$. At the microscopic level the lower the transport resistance through NiO, the higher the product $n' \cdot u = \rho / e$ where $n'$, $u$, $\rho$ and $e$ represent the charge carrier concentration (in cm$^{-3}$), the charge carrier mobility (in cm$^2$ V$^{-1}$ s$^{-1}$), the NiO photoconductivity (in S cm$^{-1}$), and the elementary charge ($1.6 \cdot 10^{-19}$ C), respectively. For NiO based $p$-DSCs the increase of $J_{\text{SC}}$ accompanying the augmentation of $\eta$ is generally determined by the increase of the photoinjected holes concentration ($n'$ term) rather than by the increase of the holes mobility ($u$ term).

The other charge transport parameter $R_{\text{rec}}$ results inversely proportional to $\eta$ (confront Tables 1 and 2). Different to $R_t$, the resistive term $R_{\text{rec}}$ depends on the rate of the bimolecular process of recombination between the photoinjected charges localized at the NiO electrode surface and iodide, i.e. the product of photoreduction. The lowest $R_{\text{rec}}$ value determined for the cell displaying the most efficacious photoconversion process, i.e. the VG11-sensitized $p$-DSC, is a straightforward
consequence of the more efficient hole injection in VG11-sensitized NiO with respect to the other sensitized systems. Upon illumination VG11 sensitizer injects a larger number of holes with respect to the other sensitizers here examined as evinced from efficiency data (Table 1). The presence of a relatively large surface concentration of holes $h^+$ will favour their recombination with iodide since the latter process is a bimolecular event the rate of which is directly proportional to the product $[I^-]'[h^+]_{inter}$ at the NiO/electrolyte interface. Therefore, the faster holes-iodide recombination occurring in VG11-C8 sensitized $p$-DSC will correspond to the lowest $R_{rec}$ with respect to the less efficient colorants VG10 and ERY. The $p$-DSC sensitized with VG10 presented the highest value of $R_{rec}$ coherently with the finding of the largest $V_{OC}$ for this cell (Table 1). The $p$-DSC sensitized with ERY showed a much higher value of $C_\mu$ (220 vs 66 $\mu$F, Table 2) compared to the SQ-sensitized counterparts. Such a finding could be associated to the spontaneous process of charge transfer between ERY dye and metal oxide even in absence of polarization (or illumination) with consequent formation of larger gradients of charge carriers concentrations in screen printed NiO with respect to the other SQ colorants (Figure ESI9). As stated in the precedent section, the phenomenon of charge transfer between sensitizer and oxide in the unbiased state of the $p$-DSC could be also at the basis of the observed behavior of ERY as a redox mediator in the electron transfer process between NiO and triiodide in dark conditions (Figure ESI9 and Table ESI3). The parameters related to the properties of the positive charge carriers in NiO, i.e. hole diffusion time $\tau_d$, hole lifetime $\tau_h$, diffusion length $L_h$ and hole diffusion coefficient $D_h$ can be derived from the fitting parameters $R_{rec}$, $R_t$ and $C_\mu$ reported in Table 2 through the relationships:

\[
\tau_d = R_t \times C_\mu \tag{1}
\]

\[
\tau_h = R_{rec} \times C_\mu \tag{2}
\]

\[
D_h = L_h^2 / \tau_d = L_h^2 / (R_t \times C_\mu) \tag{3}
\]

\[
L_h = l \times (R_{rec} / R_t)^{1/2} \tag{4}
\]

The meaning of $\tau_d$ is the time the photoinjected hole spends to reach the current collector by diffusion, the term $\tau_h$ is the lifetime of the photoinjected hole prior it undergoes recombination or
charge trapping phenomena, the term $D_h$ is the diffusion coefficient of the photoinjected hole within the NiO layer, and $L_h$ is the diffusion distance covered by the photohole before incurring recombination phenomena, $l$ being the NiO film thickness in Eq. 4.

<table>
<thead>
<tr>
<th>DYE</th>
<th>$\tau_d$ / ms</th>
<th>$\tau_h$ / ms</th>
<th>$D_h$ / $10^{-6}$ cm$^2$ s$^{-1}$</th>
<th>$L_h$ / μm</th>
</tr>
</thead>
<tbody>
<tr>
<td>ERY</td>
<td>6.40 ± 0.27</td>
<td>27.30 ± 0.74</td>
<td>26.52 ± 3.22</td>
<td>4.12 ± 0.04</td>
</tr>
<tr>
<td>VG10-C8</td>
<td>2.73 ± 0.29</td>
<td>14.83 ± 0.56</td>
<td>79.54 ± 2.56</td>
<td>4.66 ± 0.04</td>
</tr>
<tr>
<td>VG11-C8</td>
<td>1.71 ± 0.37</td>
<td>5.15 ± 0.55</td>
<td>70.00 ± 5.49</td>
<td>3.46 ± 0.01</td>
</tr>
</tbody>
</table>

Table 3. Values of the parameters related to charge carriers features in $p$-DSCs. The listed data were calculated from Eqs. 1-4 utilizing the fitting data reported in Table 2.

The most efficient photoelectrochemical cell with VG11 as colorant presented the smaller temporal parameters $\tau_d$ and $\tau_h$ as a consequence of the lowest values of the two resistive terms $R_t$ and $R_{rec}$ among the cells here tested (Table 2). The VG11-sensitized cell presented also the largest diffusion coefficient for the photoinjected holes (Table 3), since in this case $D_h$ value was principally controlled by the short time of transport diffusion $\tau_d$ (Eq. 3), but less by $L_h$ through the ratio $R_{rec} / R_t$ (Eq. 4). The latter parameter was not strongly dependent on the nature of the dye (see first column on the right of Table 3).

The ERY-based device displayed the higher values of $\tau_d$ and $\tau_h$ for the hole carriers due to the largest value of chemical capacitance (first column on the right of Table 2, Eqs. 1 and 2). For the $p$-DSCs sensitized with VG10, VG11 and ERY the respective diffusion lengths of the hole carriers were always larger than the nominal thickness of the NiO film ($l = 2$ μm). The latter condition is necessary but not sufficient to realize an efficient process of charge collection at the FTO/NiO interface since in a nanoporous film like the ones of NiO here considered the hole pathways are generally not linear. As a consequence of that, the distance the hole actually travels before it reaches the collection surface could result much longer than the value of NiO film thickness determined with the profilometer.
In the EIS study here presented we have considered also the determination of the impedance spectra for the three $p$-DSCs analyzed in Figure 7 when the applied potential of steady state was varied within a range of 40 mV around the open circuit photopotential (see caption of Figure 7). The trends of the three fitting parameters $R_{\text{rec}}$, $R_t$ and $C_h$ with the variation of the applied potential of steady-state are shown in Figure 9 whereas the correlated properties of $\tau_d$, $\tau_h$, $L_h$ and $D_h$ calculated through Eqs. 1-4 for the positive charge carriers of NiO are shown in Figure 10.
Figure 9. Variation of the fitting parameters (top) $R_t$, (middle) $R_{rec}$ and (bottom) $C_μ$ defining the TL element in the model of equivalent circuit here proposed (Figure 8) with the variation of the potential of steady-state applied on the three $p$-DSCs presented in Figure 7. It is here recalled that $V_{OC}$ values of the three $p$-DSCs in question were 100, 90 and 90 mV for VG10-, VG11- and ERY-sensitized $p$-DSC, respectively.
Figure 10. Potential dependence of the hole transport properties (top) $\tau_d$, (second from top) $\tau_h$, (third from top) $D_h$ and (bottom) $L_h$ calculated with Eqs. 1-4 in the main text from the values of $R_t$, $R_{rec}$ and $C_{\mu}$ reported in Figure 9.

Every fitting parameter $R_{rec}$, $R_t$ and $C_{\mu}$ showed a similar trend within the range of applied steady-state potential here examined regardless of the nature of the dye (Figure 9). The chemical capacitance resulted the parameter less affected by the variation of applied potential around $V_{OC}$ with the observation of the quasi constancy of the $C_{\mu}$ value for a given dye-sensitizer (Figure 9, bottom plot). The resistive terms $R_{rec}$ and $R_t$ as well as the correlated characteristic times $\tau_d$ and $\tau_h$ (Eqs. 1 and 2, Figure 10) decreased upon increase of the applied potential with respect to the open
circuit value because of the forward polarization of NiO which induces the injection of a larger number of photoholes with respect to the absence of polarization. Since $R_{\text{rec}}$ decreases more steeply than $R_t$ with the applied steady-state potential, the resulting trend of the diffusion length is the slow decrease of $L_h$ with $E_{\text{appl}}$ (Figure 10, bottom plot) with $L_h$ always surpassing $l$ in the examined range of applied potential. In sole case of VG11, the diffusion coefficient increased with the applied potential (Figure 10, third plot from top) mainly as a consequence of the decrease of the diffusion time $\tau_d$ and $R_t$, the latter resistive term representing the main controlling factor of the diffusion coefficient $D_h$ of the photoinjected holes (combination of Eqs.1 and 3, vide supra).

Finally, we recorded the EI spectra (not shown) of the VG10- and VG11-sensitized $p$-DSCs when the sensitization solution contained the concentration ratios [CDCA]:[SQ] = 0:1, 10:1 and 100:1. This set of experiments was carried out to analyze the effect of the anti-aggregating agent CDCA on the transport and charge carrier parameters of VG10- and VG11-sensitized $p$-DSCs. The electrochemical impedance spectra of the $p$-DSCs with the NiO cathodes sensitized in the solutions with [CDCA]:[SQ] = 1000:1 were not taken for the insignificance of their low efficiencies of conversion ($\eta < 0.02 \%$, Table 1). The variations of $R_{\text{rec}}, R_t$ and $C_\mu$ with the applied potential for the $p$-DSCs sensitized in differently concentrated CDCA solutions are presented in Figures ESI11 and ESI12. The corresponding variations of the hole transport properties $\tau_d, \tau_h, D_h$ and $L_h$ (Eqs.1-4) with the applied potential have been shown in Figures ESI13 and ESI14. The fitting parameters of the impedance spectra obtained with such CDCA-conditioned $p$-DSCs showed common trends of $R_t, R_{\text{rec}}$ and $C_\mu$ with the applied potential $E_{\text{appl}}$ (Figures ESI11 and ESI12): all these three parameters decrease with $E_{\text{appl}}$ as expected in a situation of forward bias no matter of the value of sensitizing ratio [CDCA]:[SQ]. The resistive term $R_t$, associated to the photocarriers transport through NiO resulted the lowest for the most efficacious sensitizing ratio [CDCA]:[SQ] (= 10:1). This is because in absence of anti-aggregating agent in the electrode dipping solution, the eventual formation of dye aggregates occurs on the NiO surface this preventing efficacious photoinjection from the excited dye aggregates. At the other extreme, i.e. for a too high concentration of CDCA with respect to SQ,
case of \([\text{CDCA}] : [\text{SQ}] = 100:1\), we would verify insufficient dye loading (Figure 4, black profiles) with poor charge injection and consequent higher transport resistance. In correspondence of the optimized sensitizing ratio \([\text{CDCA}] : [\text{SQ}] = 10:1\) (Table 1 and Figure ESI3) the term \(R_{\text{rec}}\) was the highest for VG10, and the lowest for VG11 among the various compositions of the sensitizing solution. Since the cell sensitized with VG11 generally displayed a higher conversion efficiency with respect to the one with VG10 sensitizer (Table 1), we conclude here that the parameter \(R_{\text{rec}}\) is less crucial than \(R_{\text{i}}\) in determining the overall photocell performance.

At the optimized ratio \([\text{CDCA}] : [\text{SQ}] = 10 : 1\), \(R_{\text{i}}\) is generally lower for the most performing photoelectrochemical cell, i.e. the one employing VG11 sensitizer, whereas \(C_\mu\) shows a very weak dependence on the nature of the SQ-sensitizer (Figures ESI11 and ESI12). Since any capacitive term is directly proportional to the surface charge density and inversely proportional to charge separation, the relatively high values of \(C_\mu\) for the most performing VG10- and VG11-sensitized \(p\)-DSCs have to be ascribed mostly to the amount of injected charge rather than to the spatial separation of charge which is mainly controlled by the size and the orientation of the chemisorbed dyes (Figure 5). The latter consideration stems from the assumption that charge separation at the electrode/electrolyte is controlled by the size of the molecule and not by the capability of the counter-ions of the electrolyte to reach the charged portion of illuminated sensitized NiO surface.88 Therefore, we conclude here that the most efficacious processes of photoinjection will be generally accompanied by a relatively large capacitive term \(C_\mu\) in case of \(p\)-DSCs with SQ-sensitized NiO when \(I_3^-/I^-\) is the redox shuttle. As a confirmation of that, we stress that from the observation of \(C_\mu\) trends (bottom plots of Figures ESI11 and ESI12), the capacitive term displayed the largest values at the optimized sensitizing ratio \([\text{CDCA}] : [\text{SQ}] = 10 : 1\) for both SQs under examination. Therefore, a large value of \(C_\mu\) could constitute a condition that accompanies an ameliorated performance for this type of photoelectrochemical cell. In absence of any anti-aggregating agent, the full dye-sensitization of NiO produces a corresponding \(p\)-DSC with less efficacious injection with respect to the optimized one because of the presence of unwanted aggregates of dyes which do
not allow a sufficiently fast hole injection in the photoexcited state with subsequent incapability of building up a charge distributions with high surface concentration. On the other hand, too diluted surface concentrations of anchored dye, as the ones obtained with [CDCA] : [SQ] ≥ 100 : 1 in the sensitizing solution, do not allow a sufficient injection of charge for the build-up of charge distributions with correspondingly high $C_p$ values. The discussion of the dependence of the microscopic parameters $\tau_d$, $\tau_h$, $D_h$ and $L_h$ of the $p$-DSCs employing VG10 and VG11 sensitizers on the concentration of anti-aggregating agent and applied potential (Figures ESI13 and ESI14) is reported in the ESI.

4. Conclusions

The three symmetric squaraines VG1, VG10 and VG11 of the C8 series, i.e. with an octyl group as pending N-substituent of the indolenine moiety, have been used to sensitize screen-printed NiO photocathodes for $p$-DSCs. In particular, VG11 dye has been purposely designed for a $p$-DSC employing $I_3^-/I^-$ redox shuttle taking into account the rationale of introducing the dicyano-vinyl group as electron withdrawing chromophoric unit positioned far away from the –COOH unit which connects the dye to the NiO surface. This design was inspired by the analysis of the structural factors that were associated to the most performing dye-sensitizers of $p$-DSCs with the predefined aim of accelerating electron transfer process between photoexcited dye and oxidized form of the redox shuttle as well with the intention of minimizing unwanted recombination phenomena between the holes photoinjected in NiO and the product of photoreduction $I^-$ when the sensitizer was a SQ.

The molecular design of squaraine VG11-C8 was inspired by the individuation of the function exerted by the different structural units of those dyes that constitute the most known and efficient sensitizers of $p$-DSCs. It has been proven with electrochemical and photoelectrochemical techniques that the electron attracting dicyano-vinyl group at the central core of VG11 and located in a position opposite to site of anchoring imparted a stronger directionality of the net electron transfer from NiO to triiodide mediated by photoexcited VG11 with respect to the other two SQs.
deprived of an electron withdrawing substituent. Therefore, the structure of VG11 produced favourable effects on the kinetics of dye-mediated electron transfer between the semiconductor and the redox mediator as evinced from the comparative analysis of the electrochemical impedance data of the variously sensitized p-DSCs. The benzo-indoleninic unit of VG10 and VG11, i.e. the most extended conjugated spacer here considered, was identified as the structural moiety at the basis of the relatively slower process of recombination of VG10 and VG11 vs VG1 in conditions of open circuit potential and steady state polarization. The latter conclusion was evinced from the modelling of the photoelectrochemical impedance data of the corresponding p-DSCs, which allowed the calculation of the transport parameters $R_{\text{rec}}$ and $R_{t}$, and the derived microscopic parameters $\tau_{d}$, $\tau_{h}$ and $D_{h}$. Therefore, it resulted that the larger size of VG10 and VG11 had a beneficial effect on the overall conversion efficiencies in the corresponding p-DSCs. The photoelectrochemical cells with the largest efficiencies were those displaying the lowest charge transport resistance $R_{t}$ (a parameter correlated to the amount and mobility of photoinjected charge), and the lowest recombination resistance $R_{\text{rec}}$ (a parameter correlated to the amount and lifetime of the photoinjected charge), with $R_{t}$ being more crucial than $R_{\text{rec}}$ for the achievement of the largest efficiencies. VG11 gave p-DSCs with maximum overall efficiency of 0.043 % which represents, to our knowledge the highest value of efficiency ever reported for a symmetric squaraine sensitizing a p-DSCs when the squaraine is the sole cromophoric unit of the sensitizer, and I$_{3}^{-}$/I$^{-}$ is the electrolyte mediator. Sensitization of screen-printed NiO with the three SQs here proposed has been conducted also in presence of CDCA, i.e. an anti aggregating agent preventing intermolecular interactions between of the dyes chemisorbed on electrode surface. CDCA resulted an ameliorant of the photoelectrochemical performances of the SQs-sensitized p-DSCs when the molar ratio [CDCA]:[SQ] in the solution of sensitization had an optimum value of 10:1 with an increase of overall efficiencies of about 25 % with respect to CDCA-free sensitization treatments. This was ascribed to kinetic competition between CDCA and SQ for the occupancy of the allotted sites of anchoring on NiO surface. An hypothesis of geometry of SQ anchoring has been here proposed for
the different systems taking into account how the carboxylic units were relatively oriented in the structure of these three symmetric SQs. Another aim of this work is the attempt of laying the foundation for further improvement of \( p \)-DSC performances through the proper design of SQ sensitizers as the main focus of \( p \)-DSC optimization.

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6. Supporting Informations Available

In the supporting information has been reported the paste synthetic path and dye characterization; in-depth and useful electrochemical impedance analysis has been reported. Most important new hypothesis discussed in the main text has been deeply explained in ESI section.

References


