COMPARATIVE STUDY ON THE PHOTOVOLTAIC PROPERTIES OF DYE-SENSITIZED SOLAR CELLS (DSCs) BASED ON DIFFERENT COUNTER ELECTRODE CONFIGURATIONS

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5 Original Research Article

6 ABSTRACT

7 Previously, we reported an investigation on Delonix regia dye extract as a natural sensitizer for TiO_2/DSC assembled with platinum counter electrode and low power conversion efficiency was 8 recorded. This necessitated the current investigation on *Delonix regia* dye extract as a natural 9 10 sensitizer for *TiO*₂/*DSCs* assembled with different counter electrodes. Platinum counter electrode was used for one of the DSCs while polyaniline (PANI) was used to replace platinum in the 11 other DSC. The vitriol treated PANI thin film consisted of aniline mixed with potassium 12 dichromate directly reacted on circular graphite foam. The conductivity and Hall coefficient 13 were measured to be $4.894 \times 10^{-1} \Omega^{-1} cm^{-1}$ and $2.061 \times 10^{1} cm^{3} C^{-1}$ respectively using ECOPIA 14 Hall Effect Measurement System (HMS-3000 Version 3.52). Sequel to this, the DSCs were 15 assembled and characterized using a standard overhead Veeco viewpoint solar simulator 16 equipped with AM 1.5 filter to give a solar radiation of 1000 W/m² and coupled to a Keithlev 17 source meter (model 4200SCS) which was connected to the computer via GPIB interface for data 18 acquisition. The overall solar power conversion efficiencies of 0.02% and 0.04% were obtained 19 for TiO₂-DSC//Delonix regia dye//platinum electrode and TiO₂-DSC//Delonix regia dye//PANI 20 *electrode* respectively. *Delonix regia dve extract* proved to be rather a poor sensitizer as can be 21 seen by the low spectral absorption at lower energies with short circuit current density of 22 0.10mAcm⁻² and 0.11mAcm⁻² respectively. Nevertheless, a 10% decrease in the electron 23 recombination via redox electrolyte and collection at the photoelectrode was observed for TiO₂-24 DSC//Delonix regia dye//PANI electrode and a 20% increase in the open circuit voltage (V_{ac}) 25 was also observed. Finally, about 37% increase in the fill factor was observed for the TiO₂-26 DSC//Delonix regia dye//PANI electrode over TiO₂-DSC//Delonix regia dye//platinum electrode. 27 This necessitated approximately 50% increase in the power conversion efficiency for the TiO₂-28 DSC//Delonix regia dye//PANI electrode over TiO₂-DSC//Delonix regia dye//platinum electrode. 29

Keywords: Delonix regia dye extract, PANI counter electrode, TiO₂-DSC, short circuit current
 density, open circuit voltage, fill factor, power conversion efficiency.

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34 1. INTRODUCTION

Dye-sensitized Solar Cells (DSCs) are fast becoming promising alternatives to the conventional silicon based solar cells because of cheap fabrication cost coupled with easy fabrication steps that could lead to a myriad of shapes using flexible substrates to meet the need of various

applications [1, 2, 3]. The salient features of a DSC include photoelectrode, photosensitizer, 38 electrolyte (redox couple) and counter electrode [4, 5]. However, the highest efficiency recorded 39 to date is still well below that for the silicon based solar cells [6, 7, 8]. The major factor 40 responsible for low energy conversion efficiency is the competition between generation and 41 42 recombination of photo-excited carriers in DSCs [1]. As such, most of the efforts made so far are targeted toward the synthesis of new nanostructured working and counter electrodes to 43 ameliorate this setback [9, 10, 11, 12, 13]. Sequel to this, surface modification of TiO₂ was 44 studied by depositing SrTiO₃ on its surface to form a core-shell structure in order to shift its 45 conduction band upward closer to the excited state of the coated dye causing enhancement in the 46 open-circuit voltage [11]. As for the counter electrode, the research on the 3-dimensional 47 nanostructure is currently ongoing but the increased surface area offers more locations for I³⁻ 48 reduction and also shortens the redox couple diffusion length. As a follow-up to this, a vertically 49 aligned carbon nanotube counter electrode was fabricated for use in DSC and this led to an 50 51 increased short-circuit current compared to that obtained using the conventional platinum counter electrode [12]. With platinum being a costly noble metal, reasonable efforts have been 52 made to find cheaper alternatives [14]. Such efforts include the use of porous polyaniline 53 nanotube, graphene/polyaniline nanocomposite and microporous polyaniline [15, 16, 17, 18]. 54 These concerted efforts are tied to the fact that polyaniline showed lower charge transfer 55 resistance and higher electrocatalytic activity for reduction of I_3^- into I⁻ than platinum [15, 18]. 56 Herein we report a carefully structured polyaniline (PANI) thin film as counter electrode for use 57 in DSC so as to improve its energy conversion efficiency. The film consisted of aniline mixed 58 with potassium dichromate and reacted on circular graphite foam directly to preserve the 59 60 stoichiometry and prevent over oxidation of the aniline which would have reduced the conductivity. The vitriol treated PANI is a p-type semiconducting polymer with low mobility and 61 conductivity values. The sign and value of the Hall coefficient also validated the nature of the 62 carriers with $3.029 \times 10^{17} cm^{-3}$ as the measured bulk concentration and thus can function as 63 efficient counter electrode. In our previous study, we developed and characterized a DSC based 64 on TiO₂ nanoparticles coated with delonix regia and the overall solar power conversion 65 efficiency of 0.02% and a maximum current density of 0.10mAcm⁻² were obtained. Typically, 66 low peak absorption coefficient, small spectra width and very low power conversion efficiency 67 of this DSC boosted additional studies oriented; on one hand, to the use of modified 68 photoelectrode and on the other hand, we hope to improve the power conversion efficiency with 69 use of a semiconducting polymeric counter electrode. Sequel to this, two (2) DSCs; one with 70 platinum counter electrode and the other with PANI counter electrode, were assembled and 71 characterized using a standard overhead Veeco viewpoint solar simulator equipped with AM 1.5 72 filter to give a solar radiation of 1000 W/m^2 and coupled to a Keithley source meter (model 73 4200SCS) which was connected to the computer via GPIB interface for data acquisition. 74

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77 2. MATERIALS AND METHODS

Titanium isopropoxide, Titanium nanoxide, acetylacetonate, ethanol, isopropanol, fluorine doped 78 tin-oxide (FTO) conducting glass [11.40 ohm/m², $(1.00 \times 1.00)cm^2$], electrolyte (iodolyte-AN-79 50), sealing gasket (surlyn-SX1170-25PF), and screen-printable platinum catalyst, (Pt-catalyst 80 T/SP) all were obtained from SOLARONIX. Dye extract has been obtained from the natural 81 product (Delonix regia). A mixture of 0.3M of titanium isopropoxide, 1.2M acetylacetonate and 82 isopropanol was spin coated three (3) times with different concentrations sequentially as 83 blocking layer on the pre-cleaned fluorine doped tin-oxide (FTO) conducting glasses and 84 sintered at 150°C for four minutes each time the deposition has been made. Subsequently, a paste 85 of titanium nanoxide in propanol in the ratio 1:3 has been screen printed on the three (3) fluorine 86 87 doped tin-oxide (FTO) conducting glasses and allowed to dry at 125°C in open air for 6 minutes. The FTO/TiO₂ glass electrodes have been sintered in a furnace at 450°C for 40 minutes and 88 allowed to cool to room temperature to melt together the TiO₂ nanoparticles and to ensure good 89 mechanical cohesion on the glass surface. Fresh leaves of Delonix regia have been crushed into 90 tinv bits and boiled in 75ml of deionized water for 15 minutes. The residue has been removed by 91 adopting simple physical filtering technique using muslin cloth and the resulting extract has been 92 centrifuged to further remove any solid residue. The dye extract has been used directly as 93 prepared for the construction of the DSCs at room temperature. A scattering layer of TiO₂ has 94 95 been also deposited on the TiO₂ electrodes before the electrodes have been immersed (face-up) in the natural dye extract for 18h at room temperature for complete sensitizer uptake. This turned 96 97 the TiO₂ film from pale white to sensitizer colour. The excess dye has been washed away with anhydrous ethanol and dried in moisture free air. The thickness of TiO₂ electrodes and the 98 99 deposited scattering layers was determined using Dekker Profilometer. Surface morphology of the screen-printed TiO₂ nanoparticles has been observed using EVOI MA10 (ZEISS) 100 multipurpose scanning electron microscope operating at 20kV employing secondary electron 101 signals while the corresponding Energy Dispersive Spectra (EDS) have been obtained using 102 103 characteristic x-rays emitted by TiO₂ nanoparticles. The X-ray diffraction (XRD) pattern of the screen-printed TiO₂ nanoparticles at room temperature has been recorded using X-ray 104 Diffractometer; Panalytical Xpert-Pro, PW3050/60, operating at 30mA and 40kV, with 105 monochromatic Cu-Ka radiation, of wavelength $\lambda = 1.54060$ Å. A scanned range 3–80.00553° 2 θ , 106 with a step width of 0.001° has been used. The pattern has been analyzed and the peaks have 107

been identified using ICDD data file (01-075-8897). The UV-Visible (UV-Vis) absorption 108 measurements of the dve extract and the dve extract on the screen printed TiO₂ electrodes have 109 110 been carried out with Avante UV-VIS spectrophotometer (model-LD80K). From these measurements, plots for the absorbance, Light Harvesting Efficiency (LHE) and molar extinction 111 coefficient versus the wavelengths of interest have been obtained using the relevant expressions 112 from [20]. Few drops each of aniline and $K_2Cr_2O_7$ have been coated on graphite foam by gently 113 turning the graphite foam by hand to fabricate alternative counter electrode. The mixture has 114 been grown directly on graphite foam to preserve the stoichiometry. After the process, a greenish 115 thin film of polyaniline (PANI) has been formed atop the graphite foam signifying that there was 116 no over oxidation of the aniline which would have reduced the conductivity. After drying, the 117 surface of the counter electrode has been thereafter rinsed using vitriol (H₂SO₄). Subsequently, 118 the electrical characteristics of the semiconducting PANI deposited on soda lime glass following 119 the above process have been determined using ECOPIA HALL EFFECT MEASUREMENT 120 SYSTEM (HMS-3000 VERSION 3.52). A DSC of $0.52cm^2$ active area has been assembled by 121 sandwiching a surlyn polymer foil of 25µm thickness as spacer between the photoelectrode and 122 the platinum counter electrode and then hot-pressed at 80°C for 15s. A few drops of electrolyte 123 have been introduced into the cell assembly via a pre-drilled hole on the counter-electrode and 124 sealed using amosil sealant. In order to have good electrical contacts, a strip of wire has been 125 attached to both sides of the FTO electrodes. Similarly, in assembling the modified DSC, the 126 127 same process as above has been adopted but instead of platinum counter electrode PANI coated on circular graphite foam has been clamped onto the photoelectrode to form a monolithic cell of 128 $0.78cm^2$ active area. Finally, the DSCs have been subjected to current-voltage characterization 129 using a standard overhead Veeco viewpoint solar simulator equipped with Air Mass 1.5 (AM 130 1.5) filter to give a solar radiation of 1000 W/m^2 and coupled to *Keithley* source meter (model 131 4200SCS) which has been connected to the computer via GPIB interface for data acquisition. 132 Subsequently, the working electrode and counter electrode of the DSC have been connected in 133 turn to the positive and negative terminals of the digital *Keithley* source meter respectively. The 134 bias was from short circuit to open circuit and has been obtained automatically using LabVIEW 135 software from National Instruments Inc, USA. From the data, *I-V* curves have been plotted in 136 real time for the DSCs under illuminated condition. Following this, the photovoltaic parameters 137 viz; the open circuit voltage (V_{oc}) and short circuit current (I_{sc}) were obtained from the *I-V* curves 138

for the cells. The fill factor (*FF*) and the power conversion efficiency for the cells have beenobtained using the following relations:

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$$FF = \frac{P_m}{V_{oc} \cdot I_{sc}} \tag{1}$$

$$\eta = \frac{FF \cdot V_{oc} \cdot J_{sc}}{I_{in}} \tag{2}$$

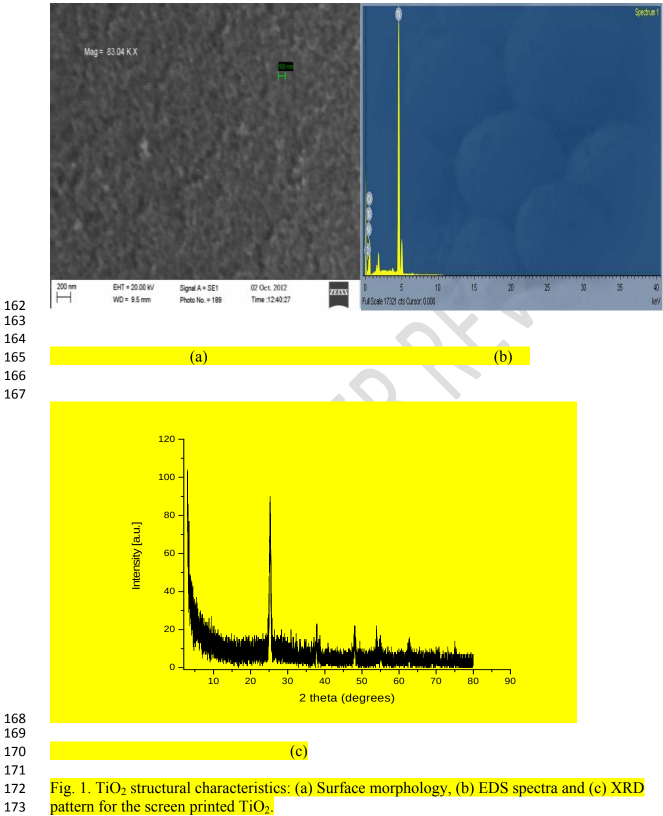
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145 **3. RESULTS AND DISCUSSION**

The image presented in Figure 1 obtained using characteristic x-rays emitted by TiO₂ 146 nanoparticles has been observed at a magnification of 83.04kX. The uniform contrast in the 147 image revealed TiO₂ to be practically isomorphic with titanium and oxygen being the dominant 148 elements with concentration of about 99.9% as depicted in the EDS spectra (Figure 1b). The 149 morphology of TiO₂ nanoparticles is such that the particles are closely parked and spherical in 150 shape. The average diameter of the particles is in the range of 25-40nm reflecting that TiO_2 151 nanoparticles are transparent and suitable for DSC application. The thickness of TiO₂ on the 152 FTO conducting glass determined using Dekker Profilometer has been found to be 5.2µm for 153 each photoelectrode and that of the deposited scattering layers has been found to be 1 µm. The 154 XRD pattern revealed the compound name for the TiO₂ electrode to be anatase syn., and the 155 structure type is tetragonal with 3.53217Å as the *d-spacing* for the most prominent peak, 156 $2\theta = 25.2139^{\circ}$ (ICDD data file: 01-075-8897). Other prominent peaks occur at $2\theta = 37.7883^{\circ}$, 157 48.0463°, 53.9110°, 55.0481°, 62.7104° and 75.1376° with d-spacing d= 2.38075 Å, 1.89370 Å, 158 1.70073 Å, 1.66826 Å, 1.48160 Å and 1.26338 Å. 159

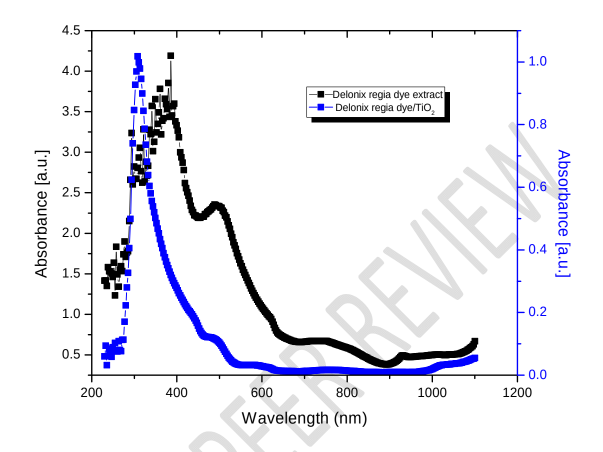
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175 In figure 2, the dye extract exhibits absorption maxima slightly above 400nm and the most prominent shoulder occur slightly above 500nm. But upon sensitization on TiO₂, there was a 176 decrease in the absorption maxima and shoulder with a cut off slightly above 600nm. It has been 177 reported that chemisorptions of anthocyanins on TiO_2 was due to alcoholic bound protons which 178 condense with the hydroxyl groups present at the surface of nanostructured TiO_2 [19]. Such 179 attachment to the TiO_2 surface stabilizes the excited state, thus shifting the absorption maximum 180 towards the lower energy of the spectrum. In our study, a shift in the absorption maximum 181 towards higher energy of the spectrum has been observed for the dye extracts adsorbed on TiO_2 . 182 This observation suggests that there was weak adsorption of the dye extract onto TiO₂ surface 183 which could be attributed to the high pH value and the long bond length of the OH groups 184 present in the dye extract. The shift may also be attributed to the changing of the anthocyanin 185 molecule from the more stable flavilium state to the unstable quinoidal state upon chelation. 186

It is an established fact that the light absorption by a dye monolayer is small since the cross section for photon absorption of most photosensitizers is much smaller than the geometric area occupied on the semiconductor surface, but with thin film semiconductor the obtainable LHE is usually close to unity [21]. In this work, we have used TiO_2 thin film of thickness 5.2µm and the

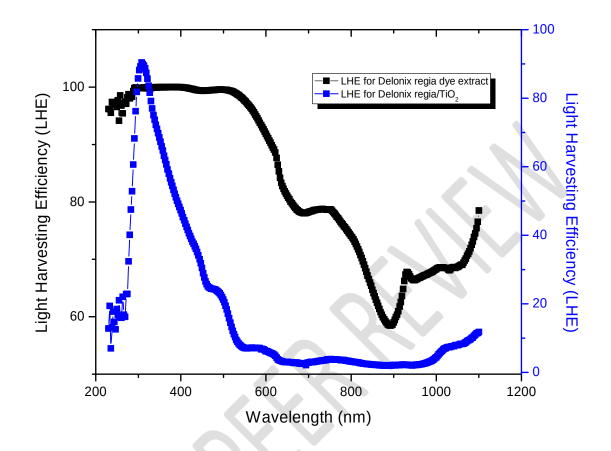
191 LHE of the dye extracts and the dye mixture adsorbed onto TiO_2 surface is close to unity.



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Fig. 2. Absorption versus wavelength (nm) for Delonix regia dye extract and Delonix regia/TiO₂
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The light harvesting efficiency values (usually obtained in percentages) are plotted against wavelengths as shown in figure 3. The absorption band of the dye extract on TiO_2 becomes a bit discrete after sensitization but quite broad for the dye extract. Whilst the molar extinction coefficients are very high for the dye extract on TiO_2 but it turned out that only small area is being covered by the solar irradiance spectrum. Most notably, the spectra bandwidth is within the range of *150nm* to *200nm* and this could be significantly enhanced if the pH is lowered using organic solvent.



206	Fig. 3. Light Harvesting Efficiency (LHE) versus wavelength (nm) for Delonix regia extract an
207	Delonix regia/TiO ₂ .

209	The e	lectrical	characteristics	for PANI	determined	using ECOP	IA HMS-3000	(VER 3.52) are
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tabulated in Table 1.

211	Table 1. Electrical Characteristics for PANI
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Bulk concentration	$3.029 \times 10^{17} cm^{-3}$
Mobility	$1.009 \times 10^{1} cm^{2} V^{-1} s^{-1}$
Sheet resistance	$6.050 \times 10^5 \Omega$
Resistivity	2.043Ω <i>cm</i>
Magneto resistance	$9.451 \times 10^4 \Omega$
Conductivity	$4.894 \times 10^{-1} \Omega^{-1} cm^{=1}$
Hall coefficient	$2.061 \times 10^{1} cm^{3} C^{-1}$

213 It is evident from table 1 that the polymeric counter electrode (PANI) is semiconducting and it is a p-type semiconducting polymer with low mobility and conductivity values. The sign and the 214 value of the Hall coefficient also validate the nature of the carrier. The bulk carrier concentration 215 is $3.029 \times 10^{17} cm^{-3}$. Current density and power versus voltage characteristics of the DSCs are 216 plotted and shown in figure 4. The photovoltaic parameters are determined and tabulated in 217 Table 2. The current density for the DSC with platinum counter electrode is $0.10 mAcm^{-2}$ while 218 that for the DSC with PANI counter electrode is $0.11 mAcm^{-2}$. This corresponds to 10% decrease 219 in the electron recombination via redox electrolyte and collection at the photoelectrode. In the 220 221 same light, a 20% increase in the open circuit voltage (V_{oc}) has been observed for the DSC with PANI counter electrode. Since the V_{oc} of an electrochemical cell is determined by the difference 222 between the Fermi level of the semiconductor and the redox potential (E_{fredox}) of the redox 223 electrolyte then, the high V_{oc} observed for the monolithic DSC suggests that this difference in the 224 Fermi levels is large. Generally the fill factor is influenced by the series resistance (R_s) arising 225 from the internal resistance and resistive contacts of the cell and shunt resistance (R_{sh}) arising 226 from the leakage of current. As such, about 37% increase in the fill factor has been observed for 227 the DSC with PANI counter electrode over the DSC with platinum electrode. Approximately, 228 50% increase in the power conversion efficiency was obtained for the DSC with PANI counter 229 electrode over the DSC with platinum electrode. Thus, it is evident from table 2 that high values 230 of J_{sc} , and V_{oc} are responsible for the higher efficiency obtained for the DSC with PANI counter 231 electrode over the DSC with platinum electrode. In our previous studies, we developed and 232 233 characterized DSC based on TiO₂//Hibiscus sabdariffa//platinum electrode and the overall solar power conversion efficiency of 0.033% and a maximum current density of 0.17mAcm⁻² have 234 235 been obtained [5]. This boosted additional studies oriented to the use of anthocyanin dyes with alternative and modified components that would lead to an enhancement in the light harvesting 236 efficiency and hence the photocurrent density which is owed to the high peak absorption 237 coefficient and large spectra bandwidth. In this work, it was discovered that TiO_2 band gap has 238 been reduced upon sensitization with the dye extract. The optical band gap obtained at the point 239 where the absorption spectra showed a strong cut off, when the absorbance value is minimum is 240 2.40 eV. The bands shift could be attributed to molecular transitions that take place when the dye 241 molecules chelate with TiO_2 . Typically, anthocyanin dyes exhibit π - π^* orbital transition which 242 is attributed to the wavelength range between 500 nm to slightly above 650 nm. 243

DSC	$\overline{J}_{sc}(mAcm^{-2})$	V_{oc} (V)	FF	η (%)
Movable TiO ₂ - DSC with Platinum electrode	0.10	0.45	0.38	0.02
Monolithic TiO2-DSC with PANI electrode	0.11	0.56	0.60	0.04

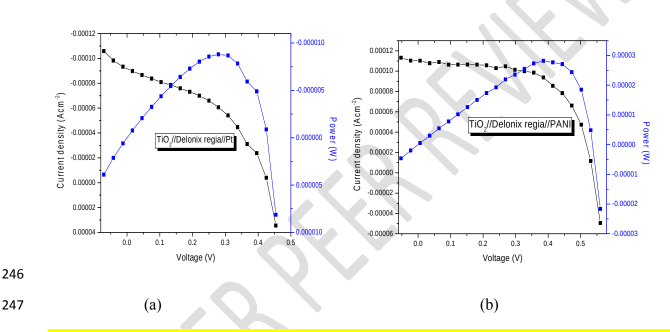


Fig. 4. Current density (J_{SC}) and Power (W) versus Voltage (V) for (a) TiO₂-DSC//*Delonix regia dye*//Platinum electrode and (b) TiO₂-DSC//*Delonix regia dye*//PANI electrode.

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In this work, the cut off wavelength for the spectra is slightly above 600 *nm*. Finally, it is well known that proton adsorption causes a positive shift of the Fermi level of the TiO_2 , thus limiting the maximum photovoltage that could be delivered by the cells [19]. Nevertheless, the TiO_2 -DSC//Delonix regia dye//PANI electrode proved to be a better cell compared to TiO_2 -DSC//Delonix regia dye//Platinum electrode that exhibited lower power conversion efficiency. However, no deviation from this trend has been observed when the duration of continuous stimulated sunlight illumination has been increased for several hours.

4. CONCLUSION

In this work we reported an investigation on *Delonix regia* dye extract as natural sensitizer for 260 261 TiO₂-DSC//Delonix regia dye//platinum electrode and TiO₂-DSC//Delonix regia dye//PANI *electrode* and the overall solar power conversion efficiencies of 0.02% and 0.04% have been 262 obtained respectively under AM 1.5 irradiation. Delonix regia dye extracts proved to be rather a 263 poor sensitizer as can be seen by the low spectral absorption at lower energies with current 264 density of 0.10mAcm⁻² and 0.11mAcm⁻² respectively. Nevertheless, a 10% decrease in the 265 electron recombination via redox electrolyte and collection at the photo-electrode has been 266 observed for TiO₂-DSC//Delonix regia dye//PANI electrode and a 20% increase in the open 267 circuit voltage (V_{oc}) has been also observed. Furthermore, the high V_{oc} observed for the 268 monolithic TiO₂-DSC//Delonix regia dye//PANI electrode suggests that the difference between 269 the Fermi level of the photoelectrode and the redox potential $(E_{f,redox})$ of the redox electrolyte is 270 large. Finally, about 37% increase in the fill factor has been observed for the TiO₂-DSC//Delonix 271 regia dye//PANI electrode over TiO₂-DSC//Delonix regia dye//platinum electrode. This 272 necessitated approximately 50% increase in the power conversion efficiency for the TiO₂-273 274 DSC//Delonix regia dye//PANI electrode over TiO₂-DSC//Delonix regia dye//platinum electrode. Although the efficiencies obtained with this natural dye extract are still below the current 275 requirement for large scale practical application, the results are encouraging and may boost 276 additional studies oriented to the optimization of solar cell components compatible with the dye. 277 278 In view of this, we are currently exploring the possibility of increasing the power-conversion efficiency of the DSCs based on TiO_2 using modified TiO_2 and counter electrodes and Delonix 279 280 regia.

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