Stability, reusability, and equivalent circuit of TiO₂-treated metakaolinite-based dye-sensitized solar cell: effect of illumination intensity on $V_{oc}$ and $I_{sc}$ values

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Abstract
In this research, treated metakaolinite (TMK) was introduced into the TiO₂ photoelectrode to fabricated dye-sensitized solar cells (DSSCs). The photovoltaic cells have four main natural components, i.e., a photosensitizer (carotenoid bixin), photoelectrode (TiO₂/kaolinite), electrolyte (glycerine carbonate derivative), and counter-electrode (carbon). Their stability, reusability, and equivalent circuit were studied. The presence of 5% of TMK in anatase TiO₂ paste decreased the TiO₂ band gap from 3.21 to 3.16 eV. The result showed that the presence of 5% of TMK in TiO₂ paste was more favorable to obtain higher energy conversion efficiency. Under a light intensity of 200 W/m², it produced an energy conversion yield of 0.086%. The combination of the electrolyte and the TMK demonstrated a synergistic effect to improve the electrical properties of the DSSC. The energy storage function worked well until the third day of analysis. The DSSC based on TiO₂/TMK photoelectrode exhibited 16 times better stability than pure TiO₂-based photovoltaic cell. The Faraday charge transfer processes showed that the TiO₂/TMK photoelectrode is not in direct contact with the carbon counter-electrode.

Keywords Bixin · Dye-sensitized solar cell · Equivalent circuit · Metakaolinite · Photoelectrode

Introduction
Dye-sensitized solar cells (DSSCs) are a third-generation photovoltaic cell that converts any visible light into electrical energy. However, ruthenium and several materials used in these cells are toxic and relatively expensive because they have a low natural abundance. They offer a promising alternative to conventional and expensive silicon-based solar cells [1–3]. Electrical energy is generated when they are exposed to light. Electrons in dye molecules are excited and then injected into a semiconductor photoelectrode (TiO₂, ZnO, WO₃, etc.) conduction band, where dye molecules adsorb. These electrons migrate through the host semiconductor particles until they reach the collector, then the holes simultaneously generated are reduced by a redox electrolyte or hole carrier at the counter-electrode (Pt, carbon, etc.). This cycle is well regenerative since no substance has been consumed or produced during the process. However, some reactions may occur and cause a significant decrease in the effectiveness of the DSSCs. They are the recombination of the electrons injected either with the oxidized sensitizer or with the oxidized redox couple on the surface of photoelectrode [2, 4].

Several studies have reported that SiO₂ and Al₂O₃ are an energy barrier for suppressing charge recombination due to their insulation properties [5–7]. They can decrease the interaction between the excited electrons in the photoelectrode and the electrolyte ions [8]. Synthetic nanoclay has been used to solidify a liquid electrolyte and induce a light scattering, increasing the overall light absorption, especially
in the red region [9–11]. Many clays have been also developed as supporting materials for photocatalytic application [12–14]. The application of natural clay for the photoelectrode in DSSCs has been scarcely reported. Saelim et al. [15] applied TiO2-modified natural bentonite clay photoelectrode for DSSCs. However, TiO2/clay photoelectrode provided a lower DSSC efficiency than the pure TiO2 photoelectrode, probably due to the high composition of clay used.

We previously reported the characteristic of treated kaolinite and metakaolinite compared with the natural kaolinite (NK). Their X-ray diffraction (XRD), Fourier-transform infrared spectroscopy (FTIR), scanning electron microscopy–energy-dispersive spectroscopy (SEM–EDS), and Brunauer–Emmett–Teller (BET) analysis data have been reported [16]. Based on that previous study, NH4OH-treated metakaolinite (TMK) has certain advantages over NK: (1) TMK has a BET specific surface area of 124.33 m2/g, 16 times greater than NK (7.65 m2/g). (2) It also has the highest Si/Al ratio (2.06), 2 times greater than NK (1.26). Their interaction with bixin has also been investigated [16, 17]. This paper presents the role of TMK in small concentrations in the TiO2 photoelectrode for DSSCs.

We used bixin (methyl hydrogen 9′-cis-6,6′-diapocarotene-6,6′-dioate) as presented in Fig. 1 as a natural sensitizer. It is a pigment in the apocarotenoids group obtained from annatto seeds (Bixa orellana L.) [18]. It carries a carboxylic group at one end and a carboxyester group at the second end of the hydrocarbon chain. This functional group allows bixin to be easily bonded to the photoelectrode surface and facilitates the injection of electrons into the photoelectrode conduction band. Bixin has absorption coefficients in aprotic polar solvents of more than 104 M−1 cm−1 which is an advantage of dyes as sensitizers in the DSSCs [19–21]. The bio-sourced carbonate solvents suitable for the redox I−/I3− pair have also been developed. Besides that, in this study, we use a mesoporous carbon-based counter-electrode which will allow the DSSCs generated to be environmentally friendly. As it has been known that the existing DSSC common component consists of toxic materials and relatively expensive because they have a low natural abundance. These materials are ruthenium as a sensitizing dye, an electrolyte containing a redox couple in acetonitrile, and a platinum-based counter-electrode [1–3].

Two of the keys parameters which govern the attainable power from a DSSC are the open-circuit voltage (Voc) and circuit current (Isc). The Voc is the maximum voltage available from a solar cell that occurs at zero current, while Isc is the current through the solar cell when the voltage across the solar cell is zero. For long-term usage, both physical and chemical stability and reusability should be considered. The model of an equivalent circuit of DSSC allows for obtaining the implied electric processes analysis. For these reasons, the results presented in this paper are related to the studies carried out on these parameters, which are influenced by direct illumination intensity.

Materials and methods

Materials

Bixin crystals containing 88.11% cis-bixin and 11.75% di-cis-bixin and an unknown compound (0.14%) were obtained from the extraction and purification processes Rahmalia et al. [22]. Glycerol carbonate acetate (GCA) were obtained by acyl transfer reaction by acylation of glycerol carbonate by acetic anhydride catalyzed by cation-exchange resins. GCA were purified by the fine film separator. The properties of GCA: N°CAS 1607-31-4; density at 25 °C 1.296; dipole moment (D) 6.19; dielectric constant (e) 47.8; boiling point, mm Hg (°C) 330; melting point (°C) –62.5; flashpoint (°C) 176 [23].

Natural kaolinite (CAS Number 1318-74-7), anatase TiO2 nanopowder (CAS Number 1317-70-0), carbon mesoporous (CAS Number 1333-86-4), and all analytical reagent (ammonium hydroxide ≥ 25% CAS Number 1336-21-6, isopropanol 99.5% CAS Number 67-63-0, acetylacetone ≥ 99% CAS Number 105-58-8) were supplied by Sigma-Aldrich, Germany. Fluorine-doped tin oxide (FTO)-coated TEC-7 conductive glasses (the material: SnO2:F, the typical layer thickness: 600 nm, the square resistance: 6–8 Ω, the total transmission at 400–800 nm: 70–81%) were provided by SOLEM.

Methods

Preparation of treated metakaolinite

The natural kaolinite (NK) was calcined at 600 °C in a furnace for 6 h to form metakaolinite (MK). An amount of 10 g of MK was added to 100 mL 5 M NH4OH. The mixtures
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were incubated at room temperature for 6 h, with constant shaking (300 rpm). The suspension was filtered. The residue was washed with distilled water until neutral and dried in an oven at 103 °C for 24 h. This process was repeated three times to optimize activation. The final product obtained was called TMK [17].

Preparation of photoelectrode

The photoelectrode was prepared on a space of 0.5 cm² by the method of doctor blading as follows [19]. The FTO substrates were sonicated in a mixture of deionized water and isopropanol (1:1 v/v) for 1 h. Clean FTO substrates were heated at 450 °C for 30 min before film deposition.

TiO₂ paste was first prepared by adding 2.5 g of TiO₂, 16 drops of acetylacetone, and 12 drops of triton™ X-100 to 20 mL of ethanol absolute, and was stirred at 200 rpm in a magnetic stirrer for 72 h until a paste formed. To make TMK-modified TiO₂ photoelectrodes, TMK was applied with different concentrations concerning the mass of TiO₂ (0, 3, 5, 7, and 10% w/w) in the paste. The TiO₂ paste was then applied on FTO by the doctor blading method using magic tape as a masking material, was air dried, and heated at 500 °C for 1 h. After cooling to 50 °C, the electrode was taken out of the oven. It was immersed in a solution of bixin diluted in dimethyl carbonate for 24 h. The concentration of the bixin used was 1.33 g/L.

Cell assembly

DSSC assembled as a sandwich structure. The carbon paste was utilized as a counter-electrode. It was prepared by adding 640 mg carbon mesoporous, 1 mL of photoelectrode paste, and 1 mL of triton x-100 into 2 mL of demineralized water. The mixture was mixed under ultrasound 20 kHz for 5 min. The carbon paste was also coated on TCO by the doctor blading method using magic tape as a masking material, was air dried, and heated at 250 °C for 1 h. A drop of electrolyte composed of 404 mg of KI, 261 mg of I₂, and 551 mg of glycerol carbonate acetate, was introduced onto the cells. Finally, the cell was assembled using epoxy glue (Fig. 2). The cells produced are called TiO₂/Bx/KI-I₂ + GCA/C, TiO₂ + 3%TMK/Bx/KI-I₂ + GCA/C, TiO₂ + 5%TMK/Bx/KI-I₂ + GCA/C, TiO₂ + 7%TMK/Bx/KI-I₂ + GCA/C, and TiO₂ + 10%TMK/Bx/KI-I₂ + GCA/C for the use of 0, 3, 5, 7 and 10% of TMK, respectively, in the TiO₂ paste. We prepared different redox couples (KI-I₂ and LiI-I₂ separately) in GCA to study electrolyte type effects.

Measurement of DSSCs performance

DSSCs performance was measured using a 50-W xenon lamp (PRO-LITE), and the intensity was measured using a multimetric SPM72 solar meter. The distance between the light and the DSSC was 20 cm, and the intensity of the lamp was adjusted using a lighting control dimmer. The $V_{oc}$ and $I_{sc}$ generated by DSSC under varying intensities (0–1000 W/m²) were determined using an Agilent 34461A multimeter equipped with Keysight BenchVue software. Measurement of $I_{sc}$ and $V_{oc}$ produced by DSSCs at each light intensity was carried out for 3 min. The equivalent circuits of DSSCs were determined using an AMETEX solartron equipped with Modulab MTS software.

Result and discussion

Characterization results of kaolinites and prepared photoelectrode

This research used the anatase crystalline phase of TiO₂ because it is an indirect band gap semiconductor and can favor electron transport and diffusion of electrolyte species, directly impacting solar cell efficiency [24, 25]. The absorption spectra (Fig. 3a) were used to determine the optical properties of the pure TiO₂ and TMK-modified TiO₂-based photoelectrodes. The absorption spectrum fitting method using Tauc model was applied to estimate their optical band gap [26]. The calculated band gap of pure TiO₂-based photoelectrode is 3.21 eV (Fig. 3b), corresponding to their absorbance at 387 nm in the ultraviolet region of the solar spectrum and consistent with the reported results [24, 26]. The band gap of TiO₂ + 3%TMK, TiO₂ + 5%TMK, TiO₂ + 7%TMK, and TiO₂ + 10%TMK photoelectrode are approximately 3.17 (392 nm), 3.16 (393 nm), 3.20 (388 nm), and 3.21 (387 nm), respectively (Fig. 3). All of the photoelectrodes yielded band gap values that were very close together, around 3.21–3.16 eV. However, there was an effect of concentration on the band gap values, and it is evident that there is a slight shift in the adsorption towards the visible region for TiO₂ + 3%TMK-, TiO₂ + 5%TMK-based photoelectrodes. The observed shift is due to the transfer of charge carriers between the TMK and the TiO₂ conduction or valence band.
Functional groups in DSSC photoelectrodes due to the addition of TMK were analyzed using FTIR. Figure 4 shows that FTIR spectrum of TiO$_2$ photoelectrode characteristic bands of: at 3398 cm$^{-1}$ corresponds to stretching of the hydroxyl group of water, 1626 cm$^{-1}$ corresponds to the O–H deformation, and 1365 cm$^{-1}$ corresponds to stretching vibration of Ti–O–Ti and O–Ti–O. For TMK-modified TiO$_2$ photoelectrodes, the peak at 1058 cm$^{-1}$ is assigned to the skeleton Si–O stretching vibration of the network (Si–O–Si and O–Si–O). This peak is not observed in the FTIR spectrum of pure TiO$_2$-based photoelectrode. The detection of O–Ti–O and O–Si–O bonds in the TiO$_2$/TMK-based photoelectrodes can support the hypothesis that the increase in energy conversion yield was influenced by these functional groups, which can facilitate the chemical adsorption of the bixin on the photoelectrode surface.

**Effect of illumination intensity on $V_{oc}$ and $I_{sc}$ values of DSSC**

Figure 5 shows the parameters affecting DSSCs performance using TiO$_2$/TMK-based photoelectrodes with different concentrations of TMK. As shown in Fig. 5, $V_{oc}$ and $I_{sc}$ increase logarithmically with light intensity. All DSSCs have the same curve pattern. TMK played an essential role in the performance of DSSCs. The $V_{oc}$ generated by TiO$_2$/Bx/KI-I$_2$+GCA/C is higher than that of TiO$_2$+3%TMK/Bx/KI-I$_2$+GCA/C, TiO$_2$+7%TMK/Bx/KI-I$_2$+GCA/C, and TiO$_2$+10%TMK/Bx/KI-I$_2$+GCA/C, but lower than that of TiO$_2$+5%TMK/Bx/KI-I$_2$+GCA/C. It could be related to the optical energy gap value ($E_g$) of anodes. Lower energy gap to be responsible the enhancement of electron injection and transport. The TiO$_2$+5%TMK/Bx/KI-I$_2$+GCA/C has the highest $V_{oc}$, and this shows that 5% of TMK added to the photoelectrode paste is the optimum concentration of TMK to apply in these works.

Overall, the intervention TMK in the TiO$_2$ photoelectrode has a significant effect on the increase of the $I_{sc}$. DSSC used TiO$_2$/TMK photoelectrode to show higher $I_{sc}$ relative to DSSC used pure TiO$_2$ photoelectrode. The percentage increase in TMK more than 5% decreases the $I_{sc}$ slightly. Since the insulating properties of SiO$_2$ and Al$_2$O$_3$, they can function as a barrier to energy by removing the recharge. In this case, TMK will reduce the interaction between the excited electrons of bixin in the TiO$_2$ photoelectrode and the electrolyte ions [6, 7, 15].

DSSC used TiO$_2$/TMK photoelectrode show the highest peak efficiency under 200 W/m$^2$ illumination. The energy conversion yield is higher than that of DSSCs with pure TiO$_2$. Under these conditions, TiO$_2$/Bx/KI-I$_2$+GCA/C, and TiO$_2$+3%TK5/Bx/KI-I$_2$+GCA/C, TiO$_2$+5%TK5/Bx/KI-I$_2$+GCA/C, TiO$_2$+7%TK5/Bx/KI-I$_2$+GCA/C, and TiO$_2$+10%TK5/Bx/KI-I$_2$+GCA/C show efficiencies 0.021, 0.014, 0.050, 0.015, and 0.013%, respectively.
Under 100 W/m² of light intensity, the efficiency obtained by TiO₂ + 5%TK5/Bx/KI-I₂ + GCA/C is also higher than that of TiO₂/Bx/KI-I₂ + GCA/C.

A redox couple is a critical element in the liquid electrolyte. Its functions concern the regeneration of the dye and the charge transport between the two electrodes, playing a crucial role in determining the photovoltaic performance of DSSCs. The redox couple of I⁻/I₃⁻ was used as a redox couple at the beginning of the DSSCs research work. It has proven to be one of the most versatile redox couples. Overall, I⁻/I₃⁻ pair has good solubility, does not absorb too much light, has an appropriate redox potential, and allows rapid dye regeneration [1, 27, 28].

In this research, we compared LiI- and KI-based redox couples, resulting in TiO₂ + 5%TMK/Bx/KI-I₂ + GCA/C and TiO₂ + 5%TMK/Bx/LiI-I₂ + GCA/C. The performance comparisons of these cells are shown in Fig. 6. Under different light intensities, TiO₂ + 5%TMK/Bx/KI-I₂ + GCA/C and TiO₂ + 5%TMK/Bx/LiI-I₂ + GCA/C produce an almost similar value of Voc, indicated by the coinciding curve. This phenomenon was in agreement with the study results by Lee et al. [29], who reported that Voc was influenced by the amount of dye adsorbed on the surface of the photocathode. Since the same photoelectrode composition was used, the amount of bixin absorbed on the surface of the photocathode of the two cells and facilitating the harvesting of photons was almost the same.

However, TiO₂ + 5%TMK/Bx/LiI-I₂ + GCA/C shows the higher I₃ relative to that of TiO₂ + 5%TMK/Bx/KI-I₂ + GCA/C. Consider the smaller radius of Li⁺ cation that causes its diffusion coefficient higher than K⁺. The Li⁺ cations had better adsorption on the TiO₂/electrolyte surface when compared to K⁺ cations [28]. Tsai et al. [30] reported that LiI improves I₃. The LiI molecule decomposes to Li⁺ when it is adsorbed on the TiO₂ surface by attracting electrons from the conduction band. Our results show that LiI-I₂-based redox couples can improve the performance of DSSC with an efficiency up to 0.062%, which is higher than that of DSSC fabricated with KI-I₂ based redox couple 0.050%.

Generally, the I⁻/I₃⁻ ratio in the electrolyte of a DSSC is adjusted to achieve the highest PV performance. During electricity generation in a DSSC, I⁻ is oxidized to I₃⁻ by the sensitizer in the photoelectrode, whereas I₃⁻ is reduced to I⁻ on the counter-electrode. Therefore, the I⁻/I₃⁻ ratio

![Diagram](https://via.placeholder.com/150)
remains constant during electricity generation. However, when the forward current continues to flow in the DSSCs, \( \Gamma^- \) is not sufficiently oxidized in the photoelectrode, causing the \( \Gamma^-/I_3^- \) ratio to increase and the current characteristics to change [31]. Because of these reasons, we also optimized LiI and I\(_2\) composition in GCA-based electrolytes. The results are shown in Fig. 7. In the following discussion, we will use the terms TiO\(_2\) + 5\%TMK/Bx/LiI-I\(_2\) + GCA-1/C, TiO\(_2\) + 5\%TMK/Bx/LiI-I\(_2\) + GCA-2/C and TiO\(_2\) + 5\%TMK/Bx5/LiI-I\(_2\) + GCA-3/C for use of 350:1, 250:1, and 150:1 in mol ratio of LiI:I\(_2\) in GCA.

Figure 7 shows that LiI-I\(_2\) ratios in the electrolyte have a great influence on the performance of DSSC. A decrease in \( I_{sc} \) and \( V_{oc} \) was observed using a larger percentage of LiI so that in this study, TiO\(_2\) + 5\%TMK/Bx/LiI-I\(_2\) + GCA-3/C showed the best performance. This observation may be due to the ion-pair formation of ion pairs and cross-linking sites that impede the movement of ions in GCA and reduce ion mobility. Under 200 W/m\(^2\) of light intensity, TiO\(_2\) + 5\%TMK/Bx/LiI-I\(_2\) + GCA-1/C, TiO\(_2\) + TMK/Bx/LiI-I\(_2\) + GCA-2/C and TiO\(_2\) + 5\%TMK/Bx/LiI-I\(_2\) + GCA-3/C efficiencies of 0.039, 0.062, and 0.086\%, respectively.

### Stability and reusability of the DSSCs

To investigate the role of TMK in DSSCs stability and reproducibility, we examined two DSSCs fabricated with the optimal conditions. One of DSSC with TiO\(_2\)/TMK photoelectrode (cell A) and the other one of cell with pure TiO\(_2\) photoelectrode (cell B). Both of cells are illuminated continuously for 7 h under 200 W/m\(^2\) of light intensity. The ability of these cells to recover energy (reusability) was also investigated. For this purpose, after 7 h of illumination, the cells are stored in the dark for 17 h. We repeated this operation for three days. Changes in the performance parameters of the DSSCs are presented in Fig. 8.

Figure 8 shows cell A produces \( V_{oc} \), \( I_{sc} \), and energy conversion yields are higher than those of cell B. This phenomenon is consistent with the results presented previously. On the first and second days, according to this figure, the two cells need 10 min of illumination time to reach the maximum value of \( I_{sc} \). After 10 min, the considerable decrease in \( V_{oc} \) and \( I_{sc} \) results in decreased energy conversion yield for both cells. We assume that the initial increase in energy conversion yield can be attributed to photoinduced annealing, then improving overall...
performance. This can occur, for example, by recovering and/or rearranging some of the dangling or taut bonds between the dye and the photoelectrode or within the dye molecules [31]. Under these circumstances, the decrease in voltage changes much more than the current.

The data also showed that the two cells assembled are reusable. They could recover their energy after being rested for 17 h. Their energy storage and charging function worked well until the third day. However, by the third day, cell B can recover less energy compared to cell A. This is shown by the initial value of $I_{sc}$ and $V_{oc}$, which fall sharply and fall below the second day. We calculated kinetics of degradation of cell A and B efficiencies using kinetic models of first (Eq. 1) and second (Eq. 2) order for quantitative analysis. In these analyses, we consider that there is only one factor affecting the decrease in cell yields.

$$\ln \frac{R_t}{R_0} = -k_1 t,$$

$$\frac{1}{R_t} - \frac{1}{R_0} = k_2 t.$$  

In these equations, $R_0$: initial energy conversion yield of the cell, $R_t$: energy conversion yield of the cell after illumination for $t$ time, $k_1$: first-order degradation rate constant, and $k_2$: second-order degradation rate constant. The $k_1$ and $k_2$ were calculated from the gradient of the corresponding curve of $\ln \frac{R_t}{R_0}$ vs. $t$ and $\frac{1}{R_t} - \frac{1}{R_0}$ vs. $t$. Calculation results are shown in Table 1.

As seen in Table 1, the first- and second-order kinetic equation correlation coefficient values are $r^2 > 0.9$ except the first-order kinetic equation for cell B on the third day. The results confirm that cell B degradation is faster than cell A on the first day, both according to the kinetic model is first order and second order. On the second day, the first-order kinetic model shows that cell A degradation constant is slightly higher than that of cell B, but according to the kinetic model of second order, cell B degradation is always much faster than cell A. Also, for the third day, cell B degradation is 16 times faster than that of cell A. These results

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**Fig. 7** Parameters affecting the performance of the DSSCs using the LiI-I$_2$-based redox couple in the different compositions
indicate that the DSSC with TiO₂/TMK photoelectrode is more stable than pure TiO₂-based DSSC. Reusability is significantly better than the CSSB incorporated in the pure TiO₂ photoelectrode.

Mathematical modeling of solar cells is essential for any performance optimization operation. An electrochemical cell such as DSSC can be represented by a network of resistors and capacitors known as an equivalent circuit. In this work, the technique of impedance spectroscopy was carried out to analyze the electrochemical phenomena in a DSSC according to Faraday’s Law. From an impedance spectrum (drawn in the imaginary plane, or as gain and phase as a function of frequency), it is possible to deduce the equivalent circuit and determine the different elements meaning. This equivalent circuit model of DSSC provides the network of cells and the system simulation and contributes to analyzing the electrical processes involved [32]. Figure 9 shows the impedance spectrum and the corresponding equivalent circuit of the DSSC with TiO₂/TMK photoelectrode (cellA).

Figure 9 schematically illustrates discrete elements of the equivalent circuit of DSSC manufactured in this research conditions. According to this figure, the model system of
DSSC introduces a resistance parallel to the capacitance related to the equivalent circuit of the PN junction [33, 34]. It is proved that the TiO2/TMK photoelectrode is not in direct contact with the counter-electrode of carbon. Since the profile of the DSSC is geometrically symmetrical, we propose some possibilities as follows: towards the TCO, TiO2/TMK forms an interface with the bixin, it gives the parallel combination of a charge transfer resistance (R2) and a capacitance space charge layer (C1). Therefore, there is no interface between TCO and bixin, and it does not need to be considered. The interface between the TiO2/TMK and the electrolyte gives another parallel combination (R3 and C2). The same is assumed for the interface between the carbon counter-electrode and the electrolyte that produces R4 and C3. The Faraday charge transfer processes could be summarized in a localized resistance \( R_1 = R \text{TiO}_2/\text{TMK/bixin} + R \text{R TiO}_2/\text{TMK/electrolyte} + R \text{carbon/electrolyte} \). The DSSC with pure TiO2 photoelectrode (cell B) shows a similar impedance spectrum pattern and the equivalent circuit.

**Conclusions**

The results are acceptable in the DSSCs using natural dyes as a sensitizer. The possibility of replacing a portion of TiO2 with TMK can lead to a considerable reduction in the cost of TiO2 used. The interaction between TiO2 and TMK was confirmed from the results of DR-UV and FTIR analysis. Besides that, an appropriate electrolyte composition, which has increased \( V_{oc} \) and \( I_{sc} \) has been identified. Means for improving the DSSC energy conversion yield of the TiO2/ TMK electrode should be examined in further studies by optimizing Ti:Si molar ratio and setting the electrode semiconductor preparation process. The presence of TMK in the TiO2 photoelectrode contributes to improving the performance, stability and reproducibility of the DSSC compared to the DSSC manufactured with the pure TiO2 photoelectrode. The electrolyte exerts a synergetic effect with the TMK to improve the electrical and conductive parameters of the DSSC.

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**Declarations**

**Ethical statements**

We declare that this paper is original. All authors approve the manuscript and this submission. We have no known competing financial interest or personal relationships that could have appeared to influence the work reported in this paper.

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