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rGO Sheets/ZnFe$_2$O$_4$ Nanocomposites as an Efficient Electro Catalyst Material for I$_3^−$/I$^−$ Reaction for High Performance DSSCs

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Abstract
In this paper, Reduced Graphene Oxide (rGO)/ZnFe$_2$O$_4$ (rZnF) nanocomposite is synthesized by a simple hydrothermal method and employed as a counter electrode (CE) material for tri-iodide redox reactions in Dye sensitized solar cells (DSSC) to replace the traditional high cost platinum (Pt) CE. X-ray diffraction analysis and High resolution Transmission electron microscopy, clearly indicated the formation of rZnF nanocomposite and also amorphous rGO sheets were smoothly distributed on the surface of ZnFe$_2$O$_4$ (ZnF) nanostructure. The rZnF-50 CE shows excellent electro catalytic activity toward I$_3^−$ reduction, which has simultaneously been confirmed by cyclic voltammetry, electrochemical impedance spectroscopy and Tafel polarization measurements. A DSSC developed by rZnF-50 CE (η = 8.71%) obtained quite higher than the Pt (η = 8.53%) based CE under the same condition. The superior performances of rZnF-50 CE due to addition of graphene in to Spinel (ZnF) nanostructure results in creation of highly active electrochemical sites, fast electron transport linkage between CE and electrolyte. Thus it’s a promising low cost CE material for DSSCs.

Keywords DSSCs · Counter electrode · rGO/ZnFe$_2$O$_4$nanocomposite · Pt-free CEs

1 Introduction

In recent years, fossil fuels becoming increasingly scarce and global warming gained momentum from rising carbon dioxide emission that imply the need to go for alternates which can replace the fossil fuel and mitigate the pace of global warming. Natural resources such as solar energy, wind energy, hydroelectric energy, geothermal and thermal energy can be harnessed as alternate energy reserves [1, 2]. The incident radiation or light energy of the sun is converted to electrical energy or electricity. The Photo Voltaic systems are employed for this conversion process. Dye Sensitized Solar Cells (DSSC) are identified as significant options due to low manufacturing cost, environmental friendly, non-pollution and high energy conversion under low light conditions [3, 4]. The major components of DSSC are photo anode, dye, electrolyte and catalytic Counter Electrode (CE). CE is an indispensable key component in the working of a DSSC to improve conversion efficiency, act as a catalyst for the tri-iodide reduction and also collects the electron from the external circuit and injecting it to the electrolyte. Traditionally Platinum (Pt) is widely used as standard CE material for the DSSC due to its good conductivity, catalytic capacity, low charge transfer resistance and higher activity in visible and infrared regions. However Pt it’s costlier, possesses highly corrosive nature in the liquid electrolyte which may lead to serious drawbacks in the large scale production [5]. Therefore, developing materials which reduce the use of Pt in DSSC is the need of the hour. So far, other alternative materials like carbon allotropes (activated carbon, CNTs and graphene), conducting polymers (CP) and transition metals have also been introduced to substitute Pt CEs. However the conversion efficiency of those electrodes is still low compared to Pt as a result of low electron conductivity and poor catalytic activity. Recently, it’s reported that graphene-based nanocomposites designed to own high chemical stability, significant electro catalytic activity towards iodine triiodide electrolyte, large surface area and more activated electrochemical functionalized sites. Also, there are reports on spinel type rGO based hybrid materials (MFe$_2$O$_4$ and MCo$_2$O$_4$ (M=Ni, Co, Zn, Mg)) examined as a CE material for DSSCs.
owing to their chemical properties, low cost, high stability against corrosion, interactions between metal ions via chemical bonds and multi metal composition provide more active sites improving the electrochemical activity [6, 7]. In this case, a novel spinel type (MFe₂O₄) nanocomposites consisting of reduced graphene oxide (rGO) are suitable with excellent electrocatalytic activity [8, 9]. Moreover, graphene possesses π-π conjugation structure, which endowed them with excellent electrical property and superior chemical stability in electrochemical environment. Properties. In this favor, we have prepared rGO/ZnFe₂O₄ nanocomposites by facile one-pot hydrothermal method with a different weight ratio of GO sheets. The influence of structural and electrochemical performances of the nanocomposites is investigated. Finally, Dye Synthesized Solar Cells are made by as synthesized CEs and the results are compared to that of the cells fabricated with Pt CE.

2 Experimental

2.1 Synthesis of rZnF Nanocomposites

The Preparation of graphene and reduced graphene oxide (rGO) procedure were followed as reported by Muthu et al. [10]. For rGO/ZnFe₂O₄ nanocomposites, 25 mg of GO were dispersed in 40 ml of DI water by ultrasonication for 30 min, on the side 1:2 ratio of Zinc (III) nitrate hexahydrate (Zn(NO₃)₆H₂O) and iron III nitrate nonohydrate (Fe(NO₃)₃0.9H₂O) were mixed 40 ml of DI water mixed by stirring. Then 5 M of NaOH solution added until the pH value exceeds above 10. The above solution mixture was slowly added to as previously prepare GO suspension. Then the solution was taken in autoclave and heated at 180 °C for 12 Hrs. After that the precipitate was collected by centrifuge several times and dried at 80 °C for 12 Hrs. The obtained sample named as rZnF-25. For rZnF-50 prepare by adapting the same procedure additionally double amount of adding GO. For comparison pure ZnFe₂O₄ (ZnF) also prepared by without adding GO.

2.2 Fabrication of DSSCs

The CEs paste is prepared by 100 mg as prepared sample was ground well in mortar, then adding (N, N-dimethylformamidel solution) NMP and PVDF to form a slurry sticky paste. Then the slurry was coated in FTO subtract by using doctor blade method with a contact area of 1.5 cm² and sintered 500 °C for 30 min. The Pt CE was prepared by drop coating method using H₂PtCl₆ solution with a contact area of 1.5 cm² and sintered 500 °C for 30 min. After natural cooling to room temperature all prepared CE was ready for both electrochemical and photovoltaic studies. simultaneously, the same procedure followed by the preparation of photo anode using commercially available TiO₂ paste. Then the prepared photo anode soaked into 0.5 M of N719 dye solution for 12 h and dried at 60°C. Liquid electrolyte (I₃⁻/I⁻) mediator prepared by 0.5 M of KI and 0.05 M of I were dissolved in acetonitrile solution. The gap between photo anode and CEs were filled with the above electrolyte solution.

The Photo conversion Efficiency and Fill Factor were calculated by the following equations

\[
\text{Efficiency(\%)} = \frac{J_{sc} \times V_{oc} \times FF}{P_{in}}
\]

\[
\text{Fill Factor} = \frac{J_{mp} \times V_{mp}}{J_{sc} \times V_{oc}}
\]

where the J_{mp} and V_{mp} are the current density and voltage at the maximum power point.

2.3 Characterization Measurements

Electrocatalytic activities of the CEs, Cyclic Voltammetry (CV) were examined 0.5 M of KI and 0.05 M of I iodine/triiodide electrolyte in the potential range 0.1–0.6 V at the Scan rate of 50mVs⁻¹. Tafel polarization curves of the symmetrical cells composing of two identical electrodes in the same electrolyte were recorded between (0.1–0.6 V) on the same workstation. Electrochemical impedance spectroscopy (EIS) of the symmetric cells were recorded at zero bias potential and 10 mV amplitude over 0.01 ≈ 10⁵ Hz. The photovoltaic (J–V) measurements of the device were at room temperature using a Keithley 2400 high current source power meter under the light illumination of 500 W Xenon lamp (AM1.5G).

3 Results and Discussions

3.1 X-Ray Diffraction

XRD pattern of GO, rGO, ZnF, rZnF-25 and rZnF-50 are Shown in Fig. 1a, b. Graphene oxide characteristic peak at 9.8° following to the plane (001) indicates the successful preparation of graphene oxide from graphite power. For rGO, small broad peak observed at 24.4°corresponds to (002) plane has confirmed the conversion of rGO from GO. The peaks at 29.78° (220), 35.15° (311), 37.90° (222), 42.73° (400) 53.28° (422), 56.71° (511) and 62.36° (440) confirm the formation of high crystallinity cubic spinel structure of ZnF, which were very well synchronized with the JCPDS (card No.01-077-0011).For nanocomposites, there is no disordered graphene oxide peaks were seen, which is due to the considerable amount of adding graphene in the composites.
and also the diffraction of disordered graphene planes is weak as compared to fine crystalline ZnF planes [11]. This confirms the graphene oxide is successfully dispersed on the surface of ZnF nanoparticles and does not make any structural changes in the planes. The average particle size is calculated using Scherer’s equations and the value is found to be 26.1, 17.8 and 15.3 nm for ZnF, rZnF-25 and rZnF-50 respectively. Also, in this study, addition of graphene plays a key factor for adjusting the size of the particles. Lattice parameter of cubic crystal lattice (a = b = c) was calculated using the formula, the obtained lattice parameter is listed in Table 1.

\[ a = \frac{d}{\sqrt{h^2 + k^2 + l^2}} \]  \hspace{1cm} (1)

### 3.2 HR-TEM

Additionally, Fig. 2a, b shows the HR-TEM images of rGO and rZnF-50 samples respectively. The sheet-like rGO sheets are found to exhibit wrinkled morphology with high transparency representing single layer graphene. In Fig. 2b it can be seen that the amorphous rGO sheets were smoothly distributed on the surface of ZnF nanostructure, which is expected to provide low charge transfer resistance and efficient channel path for conductivity of the material [10]. Figure 2c shows the SAED pattern of rGO, which confirms the amorphous nature of the graphene sheets. In Fig. 2d SAED pattern of rZnF-50, consist of several concentric diffraction rings and spots and thus indicating high crystallinity nature of the sample. The well-resolved lattice fringes are observed in the magnified HR-TEM image Fig. 2e. The fringes with interplanar spacing of 0.25 nm and 0.24 nm correspond to the (002) plane of rGO and (311) plane of ZnF, respectively. From HR-TEM analysis, the particle size is found to be 29.7, 20.2 and 18.8 nm for ZnF, rZnF-25 and rZnF-50 respectively. The decreasing in particle size benefitted the quick spread of electrolyte and fast charge transport. The obtained particle size from TEM results are in good agreement with XRD results.

### 3.3 FT-Raman

FT-Raman spectra of graphene oxide, rZnF-25 and rZnF-50 nanocomposites are shown in Fig. 3. Generally, Ferrite cubic spinel structure which gives rise to 39 normal Raman modes in which 2A1g, 1Eg, 3F2g are Raman active modes. The Raman spectra of rZnF-25 and rZnF-50 samples reveals four Raman active modes. The band at 648, 371, 297 and 198 cm\(^{-1}\) were belonging to the vibration modes of A1g, F2g, Eg and T2g respectively. The vibration mode A1g and F2g represent Fe–O and M–O stretching at the tetrahedral sites, additionally Eg and T2g modes present corresponds to the asymmetric stretching of oxygen atoms with respect to the metal ion at the octahedral site respectively [12]. The corresponding graphene oxide peaks such as 1340 cm\(^{-1}\) and 1590 cm\(^{-1}\) were represents D-Band, breathing point of Phono A1g symmetry and G-band, E2g Phonon of Sp2 C-atoms respectively. Generally, the value of the intensity ratio of the D band to the G band (I_D/I_G) is used to estimate the degree of disorder and the defects of carbon material. The obtained ratio is 0.98, 1.17 and 1.26 for rGO, rZnF-25 and

![Figure 1](image1.png)  
(a) XRD pattern of GO and rGO, (b) ZnF, rZnF-25 and rZnF-50 nanoparticles

| Table 1 Obtained parameters from X-ray diffraction |
|---------------------------------|----------|----------|----------|
| Parameter                       | ZnF      | rZnF-25  | rZnF-50  |
| Average particle size (D) nm    | 26.1     | 17.8     | 15.3     |
| Lattice parameter (a ± 0.001) Å | 8.373    | 8.362    | 8.357    |
rZnF-50 respectively. $I_D/I_G$ values were higher than that rGO confirms a low graphitic degree, more defects or edges, and disordered structures were introduced in rZnF-25 and rZnF-50 samples which could be beneficial for triiodide reduction process [8].

3.4 Electrochemical Measurements

3.4.1 Cyclic Voltametry (CV)

To evaluate the ion diffusivity and electro catalytic activity towards $I_3^-$ reduction were examined by CV measurements and (oxi/red) peaks of CE shown in Fig. 3a. The pair of negative and positive peaks was observed in all the samples were determined the following redox reactions.

$$I_3^- + 2e^- \leftrightarrow 3I^- \text{(Oxidation)}$$

$$3I^2^- + 2e^- \leftrightarrow I_3^- \text{(Reduction)}$$

The highest cathodic reduction peak current density ($J_{\text{red}}$) and the smallest peak to peak ($E_{\text{pp}}$) separation between pair of negative and positive peaks are the important parameter to determine the electrocatalytic activity towards $I^-$ regeneration [13] It is observed that rZnF-50 obtained highest $J_{\text{red}}$ density and smallest $E_{\text{pp}}$ among the other Counter electrodes. Indicates that rZnF-50 CE provide large edge active sites for electro catalytic activity, provide fast electron conducting channel and shorten diffusion sites to the $I_3^-$ reduction reaction which results faster redox reactions. Moreover, graphene could provide large pore channels which enables efficient
electrolyte diffusion path and spread quickly into the pores leading to fast reaction kinetics. The \( J_{\text{red}} \) values are in the following sequences: \( r\text{ZnF-50} > \text{Pt} > r\text{ZnF-25} > \text{ZnF} > r\text{GO} \). Stability is another significant parameter to determine function and reversibility of the CEs shown in Fig. 3b. No significant loss in after 100 CV cycles which indicates \( r\text{ZnF-50} \) has better reversibility and stability for triiodide reductions.

### 3.4.2 Electrochemical Impedance Spectroscopy (EIS)

EIS measurements were performed to evaluate charge transfer abilities at different CEs interfaces in the presence of \( \text{I}_3^-/\text{I}^- \) redox electrolyte. Figure 4c shows the nyquist plot of symmetric cells with two identical electrodes (CE/electrolyte/CE). The obtained values were fitted to the equivalent circuit shown in inset Fig. 4c. Where \( R_s \) is series resistance, \( R_{ct} \) is charge transfer resistance, \( Z_w \) is resistance of nernst diffusion [14]. Obtained values in the following sequence: \( r\text{ZnF-50} > \text{Pt} > r\text{ZnF-25} > \text{ZnF} > r\text{GO} \). \( r\text{ZnF-50} \) exhibits lower \( R_s \) and \( R_{ct} \) values than other electrodes due to addition of graphene sheets evenly covers ZnF nanoparticles which offer more channels for electron transfer and also improve electrode electrolyte interface area, as well as their bonding in the subtract benefits fast charge transfer improves electrochemical activity.

### 3.4.3 Tafel Polarization

To further confirm the electrocatalytic activity abilities of different CEs, Tafel polarization measurements were performed on same potential window as performed in CV measurements (Fig. 4d). In the tafel studies exchange current density \( J_0 \) and limiting current density \( J_{\text{lim}} \) were obtained from anodic and cathodic slopes which are directly related electro catalytic activity of CEs. \( J_0 \) and \( J_{\text{lim}} \) are obtained from the following Equations:

\[
J_0 = \frac{RT}{nF\cdot R_{ct}}
\]

\[
D_{\text{eff}} = \frac{\delta}{2nFC_{\text{lim}}}
\]

A Tafel polarization result confirms that \( r\text{ZnF-50} \) exhibit higher \( J_0 \) values than other electrodes and displays superior electrocatalytic performance of the CE material [15]. This is mainly credited to adding graphene structure into ZnF which gives large number of tiny holes and pores for \( \text{I}_3^-/\text{I}^- \) electrolyte to linkup inside the CE area and accelerates the reduction reaction. As a result, large \( J_0 \) and \( J_{\text{lim}} \) values demonstrate significant electro catalytic performance of \( r\text{ZnF-50} \) CE material. All the obtained parameters from CV, EIS are in good agreement with Tafel measurements. Additionally ECSA was examined by double layer capacitance \( (C_{dl}) \) of each CE material in different scan rates, 1 to 5 mVs\(^{-1}\) [16]. The graph plotted between \( \Delta j = (J_{\text{anodic current}} - J_{\text{cathodic current}}) \) Vs scan rates depicted in Fig. 4e. The linear fit is corresponding to double layer capacitance \( (C_{dl}) \) of the CEs. Obtained electrochemical parameters are listed in Table 2.

### 3.5 Photovoltaic Measurements

The photocurrent density (J) vs voltage (V) curves of DSSCs based on as prepared different CEs shown in Fig. 5. The obtained photovoltaic parameters are summarized in Table 3. Generally, High electrocatalytic ability of the CE material leads to fast reduction of \( \text{I}_3^- \) and generation of \( \text{I}^- \) ions, which enables to achieve highest short current (\( J_{\text{sc}} \)) and photo conversion efficiency (PCE). The DSSC fabricated with \( r\text{ZnF-50} \) CE exhibits higher \( J_{\text{sc}} \) of (17.98 mA/cm\(^2\)), FF of 0.62 value and power conversion efficiency PCE of (8.71%) than the PCE of platinum (Pt) (8.53%) CE. The increase in \( J_{\text{sc}} \) and FF is achieved by the intrinsic conductivity created by grafting graphene sheets on \( r\text{ZnF-50} \) CE which results enhanced electrochemical active sites for triiodide reduction \( \text{I}_3^- \) and generation \( \text{I}^- \) reactions. It also reduces the diffusion resistance between CE and electrolyte and smooth surface accelerates fast electron transmission pathways. At the same time ZnF and rGO exhibit lower value of \( J_{\text{sc}} \) and conversion efficiency mainly due to rough surface and low conductivity and electrochemical stability of the CEs.
Fig. 4  a CV Graphs of $I_3^{-}/I^-$ redox electrolyte for various CEs at 50mVs$^{-1}$.  b CV graph of rZnF-50 CE at 100 cycles  c Nyquist plot of GO, rGO, ZnF, rZnF-25 and rZnF-50 CEs, d Tafel polarization curve of GO, rGO, ZnF, rZnF-25 and rZnF-50 CEs, e ECSA fit of GO, rGO, ZnF, rZnF-25 and rZnF-50 CEs.

Table 2  Obtained parameters from electrochemical measurements

| CEs   | $J_{ssl}$ (mA cm$^{-2}$) | $E_{pp}$ (mA cm$^{-2}$) | $R_s$ (Ω) | $R_\alpha$ (Ω) | $Z_{d}$(Ω) | $D_D$ (cm$^2$ s$^{-1}$) | $\log J_o$ (mA cm$^{-2}$) | $\log J_{lim}$ (mA cm$^{-2}$) | ECSA (cm$^2$) |
|-------|---------------------------|--------------------------|-------------|----------------|-------------|--------------------------|---------------------------|---------------------------|-------------|
| rGO   | 0.70                      | 1.45                     | 8.9         | 2.1            | 0.89        | $2.32 \times 10^{-6}$   | 0.433                     | 0.72                       | 0.44         |
| Pt    | 4.95                      | 1.24                     | 7.4         | 1.1            | 0.32        | $4.23 \times 10^{-6}$   | 0.441                     | 0.94                       | 1.62         |
| ZnF   | 2.76                      | 1.34                     | 9.2         | 2.3            | 0.71        | $2.71 \times 10^{-6}$   | 0.434                     | 0.84                       | 0.92         |
| rZnF-25 | 4.8                      | 1.27                     | 7.8         | 1.4            | 0.35        | $3.92 \times 10^{-6}$   | 0.438                     | 0.90                       | 1.71         |
| rZnF-50 | 7.36                     | 1.21                     | 7.2         | 0.7            | 0.27        | $4.54 \times 10^{-6}$   | 0.447                     | 0.98                       | 2.35         |
Conclusion

rGO has been successfully dispersed on the ZnFe$_2$O$_4$ nanoparticle by one pot hydrothermal method. The electrochemical performance of the electrode was examined by CV, EIS and Tafel measurements. Dispersion of graphene on ZnF nanoparticle results in faster electron dispersion, rapid reduction of I$_3^-$ ions and act as a functional CE material for DSSC. The presence of graphene increases the adhesion of catalytic layer on FTO subtracts displays low charge Transfer ($R_c$) resistance and swift transfer of electrons. As a result, rZnF-50 nanocomposite is capable of providing excellent electro catalytic activity and stability towards I$_3^-$/I$^-$ redox couple. The PCE of 8.71% is obtained for rZnF-50 electrode which is better than existing Pt CE (8.53%). Its implies that to build a low cost, Pt-free fabrication of DSSCs and for its commercialization, rZnF-50 can be an alternate CE material.

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