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Earth-abundant $Fe_{1-x}S$ @S-doped graphene oxide nano–micro composites as high-performance cathode catalysts for green solar energy utilization: fast interfacial electron exchange

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In the process of conversion of solar energy into electricity and fuel, efficient electrocatalysts are indispensable. Rieske iron–sulfur protein and FeS catalysts play an important role in natural photosynthesis (NPS), and in artificial photoelectrochemical cells, respectively. Nano–micro composite catalysts (NMCCs) possess not only high catalytic activity but also fast electron transport. Herein, we prepared a nano–micro composite (NMC) of $Fe_{1-x}S$ nanoparticles decorated on sulfur-doped graphene oxide (S-GO) sheets (namely, $Fe_{1-x}S$ @S-GO–NMC) to be used as a cathode in dye-sensitized solar cells (DSCs). The GO effectively inhibit the aggregation of $Fe_{1-x}S$ nanoparticles. Notably, DSCs based on an $Fe_{1-x}SGS-GO–NMC$ cathode achieved a high solar-to-electrical conversion efficiency up to 7.23%. The conversion efficiency is, to our knowledge, one of the highest efficiencies for DSCs based on an FeS or FeS₂ cathode. Although the Fe_{1-x}S@S-GO–NMC exhibited a low thermodynamic possibility for redox reactions, it showed a higher kinetic rate than that of Pt for the charge transfer between the reaction medium and the cathode. This indicates that a fast electron exchange process occurs at the interface between the reaction and the cathode. The value of the time constant (r) corresponding to the charge exchange resistance based on Fe_{1-x}S@S-GO–NMC (0.0215 ms) was smaller than that obtained with Pt (0.261 ms). Therefore, we ascribed the superior performance of the photoelectrochemical device based on $Fe_{1-x}S@S-GO-NMC$ to its good electrocatalytic performance. The results are of great interest for fundamental research and for practical applications of FeS and FeS₂ and their composites in the solar splitting of water, artificial photoelectrochemical cells, and electrocatalysts. PAPER
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Introduction

With the continuous increase in consumption of fossil energy, environmental pollution¹ and the energy crisis² are becoming more and more stringent. As an alternative, considerable efforts have been devoted to developing new renewable and clean energy resources.³ The effective development and use of these energy sources can greatly reduce the dependence on fossil fuels, which will consequently reduce environmental pollution. The use of renewable energy sources, such as wind energy, hydro energy, or geothermal energy, is restricted by specific

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regional conditions. Solar energy offers the advantage of being available everywhere and at a lower cost. Therefore, the effective transformation and utilization of solar energy is a very promising direction in prospect. Solar energy can be converted into heat,⁴ electrical energy,⁵ or chemical energy (oxygen and carbohydrates are produced from water and carbon dioxide using sunlight).⁶ In the process of conversion of solar energy into electricity and fuel, efficient electrocatalysts are essential.⁷ For example, in a dye-sensitized solar cell and a quantum dot solar cell, redox mediators (sulfur, iodine, cobalt, ferrocene) are recycled by efficient electrocatalysts.⁸ In the process of watersplitting, the role of an efficient electrocatalyst is also indispensable.⁹

The Rieske iron–sulfur protein plays an important role in natural photosynthesis (NPS).¹⁰ In addition, FeS has great potential in dye-sensitized solar cell applications and quantum dots-sensitized solar cells (QDSCs). Quan et al. reported the application of three-dimensional (3D) FeS nanospheres as

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electrocatalysts for the S^{2-}/S_n^2 redox reaction in quantum dots-sensitized solar cells: a power conversion efficiency (PCE) of 3.34% was achieved.¹¹ FeS was deposited on 3D carbon scaffolds to develop electrocatalysts for the S^{2-}/S_n^2 redox reaction in quantum dots-sensitized solar cells, demonstrating a power conversion efficiency of 4.58%.¹² QDSCs based on an FeS/nickel foam counter electrode achieved a PCE of 4.39%.¹³ Recently, Wu^{14} and Batmunkh¹⁵ demonstrated the application of a hybrid electrocatalyst $SGN\text{-}FeS_2$ (sulfur-doped graphene with $FeS₂$ microspheres) and polypyrrole/FeS in DSCs. An SGN-FeS₂ and polypyrrole/FeS-based device displayed PCEs of 7.48% and 8.1%, respectively. In addition, the performance of carbon (such as graphene)-based cathodes in DSCs was also good.¹⁶–²³

Nano–micro composite catalysts (NMCCs) not only give higher catalytic activity but also fast electron transport.¹⁸ Herein, we report the one-pot hydrothermal synthesis of a nano-micro composite (NMC) composed of $Fe_{1-x}S$ nanoparticles decorated on sulfur-doped graphene oxide (S-GO) sheets (namely, $Fe_{1-x}S$ @S-GO–NMC). The graphene oxide substrate in NMC inhibited the aggregation of $Fe_{1-x}S$ nanoparticles. Notably, DSCs based on an $Fe_{1-x}S@S-GO-NMC$ cathode achieved a high solar-to-electrical conversion efficiency up to 7.23% with an open circuit voltage (V_{oc}) of 0.72 V, a short circuit current density (J_{sc}) of 14.08 mA cm⁻², and a fill factor (FF) of 0.72. We ascribe the superior performance of an Fe_{1-x} S@S-GO–NMC based photoelectrochemical device to its excellent electrocatalytic performance.

Experimental section

Preparation of $Fe_{1-x}S$ agglomerated particles

Typically, 0.003 mol of $FeCl₃·6H₂O$ were dissolved in 30 mL of deionized water by vigorous agitation. 0.045 mol of thioacetamide (TAA) were dissolved in 30 mL of deionized water by vigorous agitation. The above two solutions were mixed. The mixture was stirred for 0.5 h at room temperature and then transferred into a Teflon-lined autoclave. After being heated at 200 \degree C for 15 h, the product was naturally cooled to room temperature. The product was washed three times with water and ethanol.

Preparation of $Fe_{1-x}S$ nanoparticles fixed on S-doped graphene oxide (GO) sheets (namely, $Fe_{1-x}S$ @S-GO)

Graphene oxide (GO) was prepared using a modified Hummers' method.¹⁸ The specific synthesis method can be seen in our previous paper.¹⁸ The obtained GO precipitates were dispersed in deionized water to obtain a 1 wt% GO dispersion. 2 g of 1 wt% GO dispersion were dispersed in 10 mL of deionized water by ultrasound. 0.81 g of $FeCl₃·6H₂O$ were dissolved in 30 mL of deionized water by vigorous agitation. Different quantities (0.2340, 1.1348, 2.2546, 3.3847, 4.5045 g) of TAA were dissolved in 20 mL of deionized water by vigorous agitation. The above three solutions were mixed. The mixture was stirred for 0.5 h at room temperature and then transferred into a Teflon-lined autoclave. After being heated at 200 \degree C for 15 h, the product was naturally cooled to room temperature. The product was washed three times with water and ethanol.

Photoanode preparation and cell fabrication

A 12 μ m thick layer was deposited on FTO glass by printing 20 nm-sized TiO₂ particles (P25, Degussa, Germany).²⁴ The obtained film was sintered at 500 \degree C. After cooling to 90 \degree C, the TiO₂ films were immersed in a solution of N719 dye (5 \times 10^{-4} M) in acetonitrile/tert-butyl alcohol (1 : 1 volume ration) for 20 h. For the TiO₂ photoanode film treated with TiCl₄, the films were immersed in 40 mM TiCl₄ solution at 70 °C for 30 min and then sintered at 500 $^{\circ}$ C for 30 min. The triiodide/ iodide electrolyte for cell testing was composed of LiI (0.03 M), 1-butyl-3-methylimidazolium iodide (0.6 M), I_2 (0.03 M), 4-tert-butyl pyridine (0.5 M), guanidinium thiocyanate in acetonitrile (0.1 M). DSCs were assembled with a $TiO₂$ photoanode with the corresponding counter electrode sandwiching the redox couple in the electrolyte. Symmetrical cells with an effective area of 0.64 cm^2 were analyzed in a Tafel-polarization test and through EIS experiments. Paper

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Characterization

To analyze the as-synthesized composite electrocatalyst, X-ray diffraction (XRD) patterns were acquired using a PANalytical X'Pert diffractometer (Cu K α radiation at $\lambda = 1.5406 \text{ Å}$) sampling at 5° min⁻¹, 36 kV and 20 mA. As-prepared micro or nanostructures were characterized and analyzed by scanning electron microscopy (SEM, Nova Nano SEM 450). The photocurrent– voltage performance of DSCs with 0.16 cm² of photoanode film was measured without a metal mask by a Keithley digital source meter (Keithley 2400, USA) equipped with a solar simulator (IV5, PV Measurements, Inc., USA). EIS experiments were carried out with symmetrical electrodes in the dark using an electrochemical workstation (CHI760 Chenhua, China). Cyclic voltammetry (CV) was performed in a three-electrode configuration. The triiodide/iodide electrolyte for CV testing was composed of LiI (2 mM), LiClO₄ (20 mM) and I_2 (0.2 mM).

Results and discussion

In our experiments, $Fe_{1-x}S@S-GO-NMC$ was prepared using a simple one-pot solvothermal approach, as shown in Fig. 1. The amounts of $FeCl₃$ and TAA in solution were 0.003 and 0.045 mol. Meanwhile, $Fe_{1-x}S$ particles were prepared as a reference. Details of the synthesis process are reported in the experimental section. The SEM images shown in Fig. 2a illustrate that $Fe_{1-x}S$ agglomerated particles were seriously agglomerated. Fig. 2b shows that the agglomeration of $Fe_{1-x}S$ particles resulted in a non-uniform size (within the range of 500 nm to 2 microns) of the block or caking. The agglomerated $Fe_{1-x}S$ blocks or cakings cover many catalytically active sites on the surface of each $Fe_{1-x}S$ nanoparticle. In addition, the massive grain boundaries in the $Fe_{1-x}S$ blocks could possibly inhibit electron transport. These two factors may be the reason for the poor performance of previously reported devices based on pure FeS. We tried to inhibit the aggregation of $Fe_{1-x}S$ nanoparticles with GO. Fig. 2c

Fig. 1 Schematic diagrams illustrating the simple one-pot hydrothermal approach for preparing Fe_{1-x}S nanoparticles fixed on sulfur-doped graphene oxide (S-GO) sheets (namely, $Fe_{1-x}S@S-GO-NMC$).

shows the 2D structure of GO prepared using the modified Hummers' method.²⁴ The surface of GO flakes up to ten microns is very smooth with only few wrinkles. In the case of a conductive film or a composite substrate, the large size of GO is beneficial to reducing the boundaries in the thin film and helps to limit the recombination of the electrons at the

boundaries, and thus increases the conductivity. The TEM images of Fe_{1-x}S@S-GO–NMC (Fig. 2d and e) show that Fe_{1-x}S nanoparticles were dispersed on the GO surface. Fig. 2e shows the sizes of $Fe_{1-x}S$ nanoparticles dispersed on the RGO surface within the range of 40 nm to 350 nm. Thus, the GO effectively inhibits the aggregation of $Fe_{1-x}S$ nanoparticles. XRD patterns

Fig. 2 (a) SEM images of as-prepared Fe_{1-x}S agglomerated nanoparticles; (b) Size distribution of as-prepared Fe_{1-x}S agglomerated particles; (c) SEM images of as-prepared 2D GO sheets; (d) TEM images of as-prepared Fe_{1-x}S nanoparticles fixed on sulfur-doped graphene oxide (S-GO) sheets (Fe_{1-x}S@S-GO–NMC); (e) size distribution of Fe_{1-x}S nanoparticles fixed on sulfur-doped graphene oxide (S-GO) sheets; (f) XRD patterns of as-prepared $Fe_{1-x}S$ @S-GO–NMC powder.

Fig. 3 (a) Full-range XPS spectra of GO (blue), S-GO (black) and Fe_{1-x}S@S-GO–NMC (red). (b) Comparison of S 2p for S-GO (black) and $Fe_{1-x}S$ @S-GO–NMC (red). (c) Fe 2p spectra of $Fe_{1-x}S$ @S-GO–NMC (red).

of $Fe_{1-x}S@S-GO-NMC$ (Fig. 2f) show that $Fe_{1-x}S@S-GO-NMC$ displays.

 $FeS-FeS₂$ hybrid crystalline structure. Diffraction peaks at 30.0°, 30.8°, 33.8°, 43.6° and 53.3° were attributed to the (100), (002), (101), (102) and (110) planes, respectively, of the standard FeS nanocrystals with a hexagonal $(P6_3/mmc)$ structure (JCPDS#75-0602). Diffraction peaks at 28.5° , 33.0° , 37.2° , 40.7° , 47.4 $^{\circ}$ and 56.2 $^{\circ}$ could be indexed to the (111), (200), (210), (211), (220) and (311) planes, respectively, of the standard FeS₂ nanocrystals with a cubic [Pa-3(205)] structure (JCPDS#42-1340).

To further determine the structure and composition of $Fe_{1-x}S$ @S-GO–NMC, we conducted X-ray photoelectron spectroscopy (XPS) tests. The results are shown in Fig. 3. The full range of the XPS spectra of $Fe_{1-x}S@S-GO-NMC$, S-GO and GO is shown in Fig. 3a. The mole ratio of $FeCl₃/TAA = 1/15$, so the excess TAA will react with GO to form S-GO. The obvious peaks at 162.4 and 168.7 eV in the S 2p spectra of $Fe_{1-x}SGOS-GO-NMC$ can be attributed to S in the $FeS-FeS₂$ hybrid crystalline structure. In addition, shoulder peaks at 163.8 and 164.9 eV in the S 2p spectra of $Fe_{1-x}S$ @S-GO–NMC (as shown in Fig. 3b) can be observed. In order to assign shoulder peaks at 163.8 and 164.9 eV in the S 2p spectra of $Fe_{1-x}S$ @S-GO–NMC, we have synthesized S-GO with graphene oxide and TAA under the same experimental conditions as for $Fe_{1-x}S@S-GO-NMC$. The obvious S 2p peaks at 163.8 and 164.9 eV can also be observed in the S 2p spectra of S-GO, as shown in Fig. 3b. Thus, shoulder peaks at 163.8 and 164.9 eV in the S 2p spectra of $Fe_{1-x}S@S-GO$ NMC confirmed the formation of sulfur-doped graphene oxide in Fe_{1-x}S@S-GO-NMC. A complex energy distribution of Fe 2p photoelectrons was obtained and is shown in Fig. 3c. The

spectrum could be fitted using the "XPS peak" software with Gaussian functions after subtraction of a Shirley background. The Fe 2p spectrum could be fitted into four peaks which were associated with different Fe 2p in FeS and FeS_2 .^{26,27} The fitting data for different peaks are listed in Table 1. The main peaks at 712.1 and 725.4 eV were attributed to FeS $2p_{3/2}$ and $2p_{1/2}$. The peaks at 707.3 and 720.6 eV were attributed to FeS_2 2p_{3/2} and $2p_{1/2}$. As can be seen from the areas in Table 1, FeS is the dominant component in the $Fe_{1-x}S@S-GO-NMC$. The results are in agreement with the intensity of the XRD diffraction. Thus, the synthesis mechanism of $Fe_{1-x}S@S-GO-NMC$ can be described as follow:

> $CH_3CSNH_2 + H_2O \rightarrow CH_3COONH_4 + H_2S$ $FeCl₃ + H₂S \rightarrow Fe_{1-x}S + HCl + S$ H_2S + GO \rightarrow S-GO

In order to demonstrate the hypothesis that an $Fe_{1-x}S$ @S-GO–NMC cathode has a good electrocatalytic performance, we

investigated the electrocatalytic performance and the electron transfer of Fe_{1-x}S@S-GO. Fe_{1-x}S@S-GO ink was fabricated into a thin film electrode by spraying it onto a FTO/glass substrate. In our previous report, we optimized the catalyst loading in similar nano-micro composite catalysts (Fe₃O₄@RGO). For $Fe₃O₄(\text{RGGO}, a layer thickness of 16 µm is best. So we control$ the layer thickness of our cathode at 16 μ m. Meanwhile, pyrolytic platinum was prepared for use as a cathode reference. Pyrolytic platinum was fabricated on FTO/glass by pyrolysis of a 0.16 wt% H_2PtCl_6 aqueous solution. The obtained film was sintered at 450 \degree C for 1 h. For EIS and Tafel characterizations, symmetrical cells were assembled using two identical cathodes filled with I_3^-/I^- electrolytes and spaced by 40 μ m tape, as shown in Fig. 4a. The active area of the symmetrical cell was 0.64 cm². The exchange current density (J_0) in the Tafel region (as shown in Fig. 4b) is very closely related to the kinetic rate of charge transfer between the reaction medium and the cathode. Tafel polarization curves of symmetrical cells fabricated with two identical cathodes (mole ratios of FeCl₃/TAA = $1/1$, $1/5$, $1/$

10, $1/15$, $1/20$) are shown in Fig. 4b. J_0 for the cathode material synthesized with a mole ratio of $FeCl₃/TAA = 1/1$ was 1.04 mA cm^{-2} , which is lower than that based on Pt $(1.16 \text{ mA cm}^{-2})$. When the mole ratio of FeCl₃/TAA was increased to $1/5$, $1/10$, $1/15$ and $1/20$, the values of J_0 also increased. The results indicated that the element of S-GO (formed by reaction between excess TAA and GO) in $Fe_{1-x}S$ @S-GO–NMC was beneficial for improving the kinetic rate of charge transfer between the reaction medium and the cathode. For $Fe_{1-x}S$ @S-GO–NMC synthesized with a mole ratio of FeCl₃/TAA = 1/15, the J_0 (2.28 mA cm⁻²) in the Tafel zone which resulted was remarkably higher than that of Pt (1.16 mA cm^{-2}), indicating a faster electron exchange at the interface between the reaction medium and the cathode. The larger J_0 well explains the high photovoltaic activity observed with the $Fe_{1-x}S$ @S-GO–NMC cathode. The $Fe_{1-x}S$ @S-GO–NMC cathode delivered a higher photocurrent density (13.0 \pm 1.0 mA cm⁻²) compared to Pt $(12.1 \pm 0.6 \text{ mA cm}^{-2})$, as shown in Table 2. Thus, the $Fe_{1-x}S$ @S-GO–NMC cathode allows for an accelerated

Fig. 4 (a) Schematic illustration and electrochemical properties of symmetrical cells based on the $Fe_{1-x}SGS-GO-NMC$ cathode (cathode/liquid electrolyte (I $_3$ [–]/I $^-$)/cathode) under dark and Pt-based devices taken as a reference; (b) Tafel polarization curves of symmetrical cells fabricated $\,$ with two identical Fe_{1-x}S@S-GO–NMC cathodes; mole ratios of FeC₃/TAA = 1/1, 1/5, 1/10, 1/15, 1/20 (the inset table gives the values of the exchange current density J_0); (c) CV curves of the Fe_{1-x}S@S-GO–NMC (mole ratio of FeCl₃/TAA = 1/15) cathode measured by a three-electrode system in liquid electrolyte (I3⁻/I⁻); (d) EIS of symmetrical cells fabricated with two identical Fe_{1-x}S@S-GO–NMC cathodes (mole ratios of FeCl₃/
———————————————————— $TAA = 1/1$, $1/15$) under bias voltage with the open voltage corresponding to photovoltaic devices.

electron transfer and for the regeneration of the redox mediator; this property in turn increases the solar-to-electrical conversion efficiency of the corresponding photovoltaic devices. Fig. 4c shows cyclic voltammograms of $Fe_{1-x}S@S-GO-NMC$ synthesized with a mole ratio of FeCl₃/TAA = 1/15 and Pt for the I^{-}/I_{3}^{-} redox couple. Curves obtained with the $Fe_{1-x}S$ @S-GO–NMC and a Pt cathode exhibited two pairs of redox peaks. The redox peak at low potential was ascribed to the reaction: $I_3^- + 2e^- \rightarrow 3I^{-25}$ The reduction potential and the oxidation potential obtained with Fe_{1-x}S@S-GO-NMC were -0.085 V (vs. Ag/AgCl in saturated potassium chloride) and 0.369 V (vs. Ag/AgCl in saturated potassium chloride), respectively. The reduction potential and oxidation potential obtained under the same conditions with Pt

were 0.078 V and 0.281 V. The separation between anodic and cathodic peaks (ΔE) is inversely proportional to the thermodynamic possibility of the above redox reaction. Interestingly, the ΔE value for Fe_{1-x}S@S-GO–NMC (0.454 V) was higher than that for Pt (0.203 V), which may explain why the photovoltaic devices based on an $Fe_{1-x}S$ @S-GO-NMC cathode delivered a lower fill factor (0.72 \pm 0.02) compared to the Pt cathode (0.78 \pm 0.01), as shown in Table 2. EIS was carried out to reveal the inherent interface resistance, as shown in Fig. 4d. The arc at high frequency depends on the charge transfer resistance (R_{ct}) at the interface between the cathode and the electrolyte. This parameter reflects the performance of the cathode material. The point of intersection between the plot at high frequency and the real

Fig. 5 (a) Schematic illustration and solar-to-electrical conversion properties of DSCs based on the Fe_{1-x}S@S-GO–NMC cathode under AM1.5, 100 mW cm⁻² simulated illumination. Pt-based devices are also shown as a reference (the inset table gives the values of the best photovoltaic parameters); (b) stability of the Fe_{1-x}S@S-GO–NMC cathode measured by a three-electrode system in I₃⁻/I⁻ liquid electrolyte; (c) the values of the redox current density in each cycle of the CV plots.

axis of the Nyquist plot gives the series resistance (R_s) . The values of the EIS parameters were obtained by fitting Nyquist plots with the Z-view software according to an equivalent circuit diagram (Fig. 4d, inset). The tting results of the Nyquist plots are listed in Table 2. The R_{ct} of symmetrical cells based on FeS@S-GO–NMC (synthesized with mole ratios of $FeCl₃/TAA$ = 1/1 and 1/15) and a Pt cathode were 16.17, 2.59 and 9.17 Ω $\text{cm}^2,$ respectively. The reciprocal of the highest frequency (f) of the top of the arc corresponds to the charge exchange resistance representing the reaction time constant (τ) of the reactant at the cathode surface. The smaller the time constant, the faster the reaction. The value (0.0215 ms) of τ obtained with the Fe_{1-x}S@S-GO–NMC cathode (synthesized with a mole ratio of $FeCl₃/TAA$ = 1/15) was smaller than those obtained with Pt and $Fe_{1-x}S@GO-$ NMC (synthesized with a mole ratio of FeCl₃/TAA = $1/1$). Thus, the Fe_{1-x}S@S-GO-NMC (synthesized with a mole ratio of $FeCl₃/TAA = 1/15$) electrode exhibits a fast electron exchange process occurring at the interface between the reaction and the cathode. In addition, the Z_N of symmetrical cells based on $Fe_{1-x}S$ @S-GO–NMC (synthesized with a mole ratio of FeCl₃/TAA = 1/1) is 44.06 Ω cm², which is smaller than that of a Pt cathode (132.80 Ω cm²). This indicates that the channels formed by 2D lamellar graphene are beneficial for the diffusion of the electrolyte, which makes the diffusion resistance (Z_N) smaller. From the analysis of CV, Tafel and EIS results, we can see that the $Fe_{1-x}S@S-GO-NMC$ cathode exhibited excellent electrocatalytic performance, which in turn promoted the performance of the corresponding photovoltaic devices. **BSC** Advances Some Constructer Constraints (Eq.) The CONCIUSIONS Constraints are considered on a magnitude of the sequent plus in considered are commonly are a located in the commonly are commonly are expected unit are c

The photovoltaic performance was investigated by characterizing DSCs based on an $Fe_{1-x}S$ @S-GO (synthesized with a mole ratio of FeCl₃/TAA = $1/15$) cathode under AM 1.5, 100 mW cm^{-2} simulated illumination (schematic diagrams shown in Fig. 5). For each cathode, we fabricated four DSCs devices and determined the mean photovoltaic performance. The detailed photovoltaic parameters are summarized in Table 2. Fig. 5a shows the best photocurrent density with respect to voltage (J–V curve) for the DSCs in each group. The DSCs based on the $Fe_{1-x}S$ @S-GO–NMC cathode showed a solar-to-electrical conversion efficiency of 7.23% with an open circuit voltage (V_{oc}) of 0.72 V, a short circuit current density $(J_{\rm sc})$ of 14.08 mA cm⁻² and a fill factor (FF) of 0.72. The conversion efficiency based on the $Fe_{1-x}S$ @S-GO–NMC cathode is nearly the same as that obtained from DSCs based on pyrolytic Pt cathodes (7.21%). Although the device is very similar to a tandem DSC, we did not find that the value of V_{oc} in the device based on the Fe_{1-x}S@S-GO-NMC cathode was significantly higher than from the one obtained with the reference Pt. Therefore, the excellent device performance obtained with the $Fe_{1-x}S@S-GO-NMC$ cathode is ascribed to the excellent electrocatalytic activity of the composite material, which could effectively recycle the redox mediator and reduce the over-potential. In addition, an investigation of the stability of $Fe_{1-x}S@S-GO-NMC$ in I_3^-/I^- liquid electrolyte was carried out, as shown in Fig. 5b and c. The changes in the potentials and current density for the redox peaks in each cycle of the CV plots were not obvious. The results showed that the $Fe_{1-x}S@S-GO-NMC$ exhibited good stability in I_3^-/I^- liquid electrolyte.

Conclusions

In conclusion, we synthesized a nano–micro composite composed of $Fe_{1-x}S$ nanoparticles decorated on sulfur-doped graphene oxide sheets by a one-step hydrothermal method. Compared to $Fe_{1-x}S$ agglomerated particles, GO sheets in NMC obviously prevented the aggregation of $Fe_{1-x}S$ nanoparticles. DSCs based on an $Fe_{1-x}S$ @S-GO–NMC cathode achieved the highest reported solar-to-electrical conversion efficiency up to 7.23% for photoelectrochemical cell. The superior performance of the photoelectrochemical device based on $Fe_{1-x}S@S-GO$ NMC was attributed to the good electrocatalytic performance of the material. We hope that our work here will be of great interest for fundamental research and for practical applications of FeS or $FeS₂$ and their composites in the solar splitting of water, artificial photoelectrochemical cells, and electrocatalytic applications.

Conflicts of interest

There are no conflicts to declare.

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