Chemical Science



EDGE ARTICLE

View Article Online
View Journal | View Issue



Cite this: Chem. Sci., 2017, 8, 6111

Received 20th April 2017 Accepted 24th June 2017

DOI: 10.1039/c7sc01756a

rsc.li/chemical-science

CoO_x nanoparticle anchored on sulfonatedgraphite as efficient water oxidation catalyst[†]

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Development of efficient, robust and earth-abundant water oxidation catalysts (WOCs) is extremely desirable for water splitting by electrolysis or photocatalysis. Herein, we report cobalt oxide nanoparticles anchored on the surface of sulfonated graphite (denoted as "CoO_x@G-Ph-SN") to exhibit unexpectedly efficient water oxidation activity with a turnover frequency (TOF) of $1.2~{\rm s}^{-1}$; two or three orders of magnitude higher than most cobalt-based oxide WOCs reported so far. The CoO_x@G-Ph-SN nanocomposite can be easily prepared by a soft hydrothermal route to have an average CoO_x size below 2 nm. Additionally, the loading of CoO_x@G-Ph-SN catalyst on the surface of a BiVO₄ or Fe₂O₃ photoanode can boost remarkably the photoanode currents for robust photocatalytic water oxidation under visible light irradiation. Its excellent activity and photochemical stability for water oxidation suggest that this ultrasmall cobalt-based composite is a promising candidate for solar fuel production.

Introduction

Current energy resources are derived mainly from fossil fuels (oil, coal and gas), and the utilization of these has caused numerous environmental problems, such as the increasing emission of greenhouse gas, most notably carbon dioxide (CO₂).¹ To address this issue, an increased amount of clean, renewable and sustainable energy resources should be presented for energy resource distribution.² Solar energy is the largest clean and renewable energy source on the Earth.³ Solar-driven water splitting can store energy in the form of hydrogen fuel,² and water oxidation is considered to be the key reaction for overall water splitting.⁴,⁵ Therefore, the development of highly efficient and robust water oxidation catalysts (WOCs) is crucial to the implementation of solar fuel production.

In the past decades, continuous efforts have been devoted to developing efficient and robust WOCs. Mn-,⁶⁻⁸ Fe-,⁹ Co-,¹⁰⁻¹³ Ni-,¹⁴ Ru-,¹⁵⁻¹⁷ and Ir-based¹⁸ complexes or metal oxides have been examined for water oxidation by chemical, electrochemical or photocatalytic approaches. Some homogeneous molecular complexes, such as [Ru(bda)(isoq)₂] (H₂bda = 2,2'-bipyridine-6,6'-dicarboxylic acid; isoq = isoquinoline)¹⁵ and halogen substituted [Ru(bda)(isoq)₂]¹⁷ exhibited comparable catalytic activity for water oxidation to the CaMn₄O₅ complex in PSII of natural photosynthesis. In addition, cobalt-based polyoxometalates [Co₄(H₂O)₂ (PW₉O₃₄)₂]¹⁰⁻ and Na₁₀[Co₄(H₂O)₂(VW₉O₃₄)₂]·35H₂O showed

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† Electronic supplementary information (ESI) available. See DOI: 10.1039/c7sc01756a

very high catalytic activity ($TOF = 5 \text{ s}^{-1}$) for water oxidation in the $[Ru(bpy)_3]^{2^+}$ -sulfate system under light irradiation.^{11,13} Although the homogeneous complexes have exhibited satisfactory water oxidation activity, they are seldom applied in real solar water oxidation systems, mostly due to their poor photochemical stability. Comparatively, heterogeneous WOCs are robust and thus frequently supported on the surface of semiconductors for solar water oxidation.^{19,20}

Nocera's group reported an *in situ* electrochemical synthesis of cobalt phosphate films on an indium tin oxide anode, which can oxidize water well under a neutral pH environment.21 Since then, the development of cobalt-based WOCs as well as their application in the fabrication of artificial photosynthesis devices has gained considerable attention. For example, Frei et al. synthesized ~ 25 nm Co_3O_4 nanoclusters supported inside mesoporous silica (SBA-15 and KIT-6), and found that smaller Co₃O₄ clusters showed higher water oxidation activity.²² The surface of g-C₃N₄ modified with layered Co(OH)₂ can not only accelerate the interface transfer rate of charge carriers, but also reduce the excessive energy barrier for O-O formation, thus leading to enhanced reaction kinetics for photocatalytic water oxidation.²³ CoO_x as O₂-evolving cocatalysts supported on the surface of a LaTiO2N photocatalyst also showed remarkable promotion of water oxidation performance under visible light irradiation. 19,24 However, the size of CoO_x nanoparticles reported in most of the previous literature is usually larger than 5 nm, with a catalytic activity that is at least 2 orders of magnitude lower than those of homogeneous Co-based catalysts.10

Recently, many heterogeneous catalysts with single atoms or nanocluster structures have exhibited significantly enhanced catalytic activities with respect to conventional bulk Chemical Science Edge Article

catalysts.25,26 Inspired by this, one possible strategy to obtain highly active heterogeneous Co-based WOC is to reduce further the size of the cobalt oxide. However, the reduction of particle size is generally accompanied by an enhancement of surface energy, causing aggregation and instability of the WOC. Thus, further efforts on how to stabilize the ultrasmall nanoparticles should be considered. To address this, loading of ultrasmall nanoparticles onto the surface of a solid support has been known as effective way to stabilize nanoparticles.10 In addition, as for the application of WOC in photo(electro)catalytic water splitting, another design consideration is that the support should possess good conductivity for efficient carrier transfer. Based on these findings, the synthesis of ultrasmall Co-based WOCs anchored on the surface of a conductor is highly desirable. Graphite not only possesses good mechanical stability, but also exhibits excellent conductivity, and has been used widely in the field of solar cell and electrochemical water splitting. 27-29 Accordingly, graphite is expected to be a good support to stabilize ultrasmall Co-based nanoparticles for further solar water splitting.

Herein, we report functionalized graphite-anchored CoO_x nanoparticles with an average size of sub-2 nm to exhibit unexpectedly efficient heterogeneous water oxidation activity. The sub-2 nm CoO_x nanoparticles were hydrothermally synthesized by anchoring them on the surface of phenylsulfonic acid-functionalized graphite. The TOF value of water oxidation on the optimized CoO_x @G-Ph-SN sample can reach as high as 1.2 s⁻¹, over two orders of magnitude higher than most heterogeneous transition metal oxides. In addition, the direct coupling of CoO_x @G-Ph-SN with an $\mathrm{Fe}_2\mathrm{O}_3$ photoanode demonstrates its good photochemical stability under an artificial photosynthesis environment.

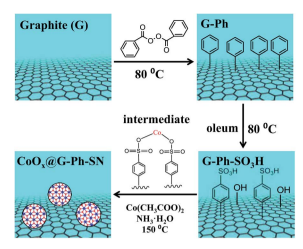
Experimental

Materials and reagents

All chemicals were analytical grade and used as purchased without further purification. Solutions were prepared using high purity water (Millipore Milli-Q purification system, resistivity > 18 M Ω cm). The fluorine-doped tin oxide (FTO) conductive glass was purchased from Nippon Sheet Glass Company (Japan) and was ultrasonically cleaned with acetone, ethanol and deionized water for 20 min each in sequence prior to use.

Synthesis of samples

As described in Scheme 1, the synthesis of the $CoO_x@G-Ph-SN$ sample mainly involves three steps: (i) immobilization of phenyl on the surface of graphite (G-Ph); (ii) sulfonation of phenyl (G-Ph-SO₃H); and (iii) hydrothermal synthesis of CoO_x anchored on the surface of phenylsulfonic acid functionalized graphite ($CoO_x@G-Ph-SN$). The first two steps were achieved by referring to previous work.³⁰ As for the first step on immobilization of phenyl on the surface of graphite, typically, the dark gray graphite powder (0.5 g, 41.6 mmol of carbon) was dispersed in benzene (400 mL) in a 500 mL three necked round-bottom flask



Scheme 1 Schematic illustration of the synthesis of CoO_x nanoparticles anchored on sulfonated graphite (CoO_x @G-Ph-SN).

equipped with a magnetic stir bar. The contents were then stirred vigorously after benzoyl peroxide (10.1 g, 41.6 mmol) was added. The mixture was then heated at 80 $^{\circ}$ C for 12 h with continuous vigorous stirring. After cooling down, the contents of the flask were centrifuged and washed with ethanol for four times. The black solid was dried at 60 $^{\circ}$ C overnight, which was nominated as G-Ph.

As for the second step on phenylsulfonic acid functionalized graphite, a typical experimental process was as follows: phenylated graphite (G-Ph) (200 mg) was dispersed in oleum (70 mL, $\rm H_2SO_4$, 25% as free $\rm SO_3$), and heated at 80 °C for 5 h to yield phenyl sulfonated graphite. After cooling down, 300 g of ice block was then carefully added into the suspension. The mixture was then centrifuged and washed with water several times until the pH value of the filtrate reached \sim 7. The obtained solid was dried at 60 °C overnight, which was nominated as G-Ph-SO₃H. For comparison, the pristine graphite was also treated with oleum in a similar process with the synthesis of G-Ph-SO₃H, which was nominated as G-O.

As for the third step, the hydrothermal anchoring of CoO_x on the functionalized graphite (G-Ph-SO₃H), typically 13.2 mg of $Co(CH_3COO)_2 \cdot 4H_2O$ and 150 mg of G-Ph-SO₃H were mixed in 20.0 mL of ethanol solution containing 50 μ L of deionized water. After about 10 min stirring, 75 μ L of 28% ammonia were added and mixed for another 10 min. Afterwards, the mixture was transferred into a 30 mL Teflon autoclave and heated at 150 °C for 2 h. After heat treatment, the autoclave was cooled to room temperature, and the product was washed with deionized water for more than three times. The final product was dried at 60 °C overnight, which was nominated as $CoO_x@G-Ph-SN$.

In this work, the immobilization of phenyl and sulfonation of phenyl were confirmed by FTIR (ESI Fig. S1†), UV-Vis (ESI Fig. S2†) and XPS (ESI Table S1†), and the possible formation of cobalt benzenesulfonate intermediate can be verified by UV/Vis diffuse reflectance spectra (ESI Fig. S2 and S3†). As for discussion and comparison, Co_3O_4 nanoparticles (Co_3O_4 -nano), CoO_x G-O and CoO_x graphite nanocomposites were similarly synthesized by following the above hydrothermal process

(the third step) except that the former was free of G-Ph-SO $_3$ H, the middle employed G-O to substitute G-Ph-SO $_3$ H, and the latter employed graphite to substitute G-Ph-SO $_3$ H. Commercial Co $_3$ O $_4$ was also employed for comparison.

Preparation of photoanodes

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The preparation of the nanoporous BiVO₄ photoanode was as described in previous work.³² Typically, F-doped SnO₂ (FTO) substrates were first electrochemically deposited with BiOI, and then a dimethylsulfoxide (DMSO) solution of vanadyl acetylacetonate [VO(acac)₂] was dropped onto their surface and heated in air at 450 °C for 2 hours to obtain BiVO₄ photoanodes. Afterwards, a calculated amount of pre-dispersed CoO_x@G-Ph-SN ethanol solution was dropped onto the BiVO₄ photoanode, which was then heated in air at 80 °C for 1 h to obtain the CoO_x@G-Ph-SN/BiVO₄ photoanode.

As for preparation of Fe_2O_3 photoanodes, Fe_2O_3 was deposited on an FTO substrate electrode using a modified chemical bath deposition method reported elsewhere.³³ Afterwards, a calculated amount of pre-dispersed $CoO_x@G$ -Ph-SN ethanol solution was dropped onto the surface of the Fe_2O_3 photoanode, which was further heated in air at 65 °C for 10 min to produce the $CoO_x@G$ -Ph-SN/ Fe_2O_3 photoanode.

Characterizations of samples

The as-prepared samples were characterized by X-ray powder diffraction (XRD) on a Rigaku D/Max-2500/PC powder diffractometer. The sample powder was scanned using Cu Ka radiation with an operating voltage of 40 kV and current of 200 mA. A scan rate of 5° min⁻¹ was applied to record the patterns in the range of 10-80°. Transmission electron microscope (TEM) images were observed by a Hitachi HT7700. High resolution TEM (HRTEM) images were recorded on a JEM-2100 transmission electron microscope (Tokyo, Japan) at 200 kV. The loading amount of cobalt oxide in the catalyst was determined using an inductively coupled plasma atomic emission spectrometer (ICP-AES) on a Shimadzu ICPS-8100. Prior to ICP-AES measurement, the supported cobalt oxide was dissolved in aqua regia. FT-IR spectra were obtained using a Varian 3100 FTIR spectrophotometer in DRIFT mode (diffuse reflectance infrared Fourier transform). The spectra were collected in the wavenumber range from 3900-400 cm⁻¹ with 2 cm⁻¹ resolution (average of 32 scans). The valence state of the cobalt oxide cluster was determined using XPS recorded on a Thermo ESCALAB 250Xi. The X-ray source selected was a monochromatized Al Ka source (15 kV, 10.8 mA). Region scans were collected using a 20 eV pass energy. Peak positions were calibrated relative to the C 1s peak position at 284.6 eV.

Tests of chemical water oxidation

Typically, a calculated amount of $0.03-0.3~g~L^{-1}~CoO_x@G-Ph-SN$ was added to a borate buffer solution (pH 9.0, 3.0 mL) containing Na₂S₂O₈ (5.0 mM) and [Ru(bpy)₃](ClO₄)₂ (1.0 mM). After 5 min stirring, the mixture was then irradiated with a halogen lamp through a coloured filter glass transmitting $\lambda > 420$ nm at room temperature. The photon flux of the incident light was

determined to be 1.28×10^{-7} mol cm⁻² s⁻¹ using an EKO LS-100 spectroradiometer. Evolved oxygen gas was monitored by using a Clark-type electrode (Strathkelvin SI130 UK). The turnover frequency (TOF) was calculated by using the initial constant O_2 evolution rate and assuming all the cobalt atoms were active sites. The contents of cobalt on typical samples were analyzed by ICP-AES measurement, which are 1.2 wt% and 1.29 wt% for CO_2 graphite and CO_2 (a)G-Ph-SN, respectively.

Tests of electrochemical water oxidation

The electrochemical water oxidation performances of all the cobalt-based electrodes were tested in a conventional three-electrode electrochemical cell with a platinum plate as the auxiliary electrode and a saturated calomel electrode (SCE, saturated KCl) as the reference electrode. A 1 M NaOH aqueous solution was used as the electrolyte with a pH measured at *ca.* 13.6. The scanning rate was 10 mV s⁻¹. All potentials measured were calibrated to RHE using the following equation: E(RHE) = E(SCE) + 0.241V + 0.0591pH.

Photoelectrochemical (PEC) measurements

The PEC tests were conducted in a three electrode system under simulated AM 1.5G solar light irradiation (100 mW cm $^{-2}$, Newport Sol3A, Class AAA Solar simulator) or a 300 W xenon lamp (PLS-SXE300C, Perfectlight Company). The fabricated electrode, a platinum electrode, and a SCE were used as the working, counter and reference electrodes, respectively. 0.5 M lithium borate buffered solution (pH 9) or 1 M NaOH aqueous solution (pH = 13.6) was used as an electrolyte after saturation with Ar gas for 30 min. The photocurrent was measured by linear sweep voltammetry with a scan rate of 10 mV s $^{-1}$. The light irradiation came from the front side of the electrodes in all cases

Kelvin probe force microscopy measurements

Kelvin Probe Force Microscopy (KPFM), which measures the contact potential difference between the probe and sample, was employed for potential imaging.³⁴ An amplitude-modulated mode of the KPFM in lift mode was used. For each scan, a SCM-PIT Pt/Ir-coated conductive probe passes over the surface twice. On the first pass, a feedback loop controls the sample height to acquire the topography and phase signal of the sample. On the second pass, the cantilever is held at 100 nm of tip-sample distance to record the surface potential. The scan rate is 0.5 Hz.

Results and discussion

As seen in the TEM images of Fig. 1a and b, the CoO_x nanoparticles on the surface of $CoO_x@G$ -Ph-SN (Fig. 1a) are more homogeneously dispersed compared to the $CoO_x@g$ -papite (Fig. 1b), $CoO_x@G$ -O (Fig. S4†) and Co_3O_4 -nano (Fig. S4†); based on which their average sizes are calculated to be 1.6, 3.2, 2.5 and 5.3 nm, respectively. This demonstrates that compared to the synthesis of Co_3O_4 -nano, the addition of graphite can inhibit the growth of CoO_x nanoparticles ($CoO_x@g$ -graphite) to a certain extent, but its aggregation (Fig. 1b) is clearly observed

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h CoO @G-Ph-SN satellite 2p. CoO_@G-O Intensity (a.u.) CoO @G-Ph-Intensity (a.u.) CoO_@graphite CoO @graphite Commercial Co₃O Co,O,-nano 400 800 810 780 200 600 800 Raman shift (cm⁻¹) Binding Energy (eV)

Fig. 1 Representative TEM images of typical samples: (a) $CoO_x@G$ -Ph-SN, (b) $CoO_x@g$ raphite; and their Raman (c) and XPS (d) spectra.

because of the hydrophobic surface feature of graphite itself. However, the surface of graphite after the functionalization of phenylsulfonic acid is changed from hydrophobic to hydrophilic according to the measurement of its contact angle (Fig. S5†). A hydrophilic surface is generally favorable for the homogeneous dispersion of CoO_x nanoparticles.¹⁹ In addition, the phenylsulfonic group is expected to complex with Co²⁺ according to comparison of the UV-Vis absorption results on typical samples (Fig. S2†), which should be another factor inhibiting its growth kinetics, causing the reduction of particle size. As an extended discussion, we also synthesized the oleum modified graphite for the loading of CoO_r nanoparticles. Similarly, the surface of graphite after treatment by oleum changes from hydrophobic to hydrophilic according to the measurement of its contact angle (Fig. S5†), and after loading the same amount of CoO_x on the oleum treated graphite, the dispersion of CoO_x can be improved and the particle size can be reduced compared with those on the pristine graphite (Fig. S4†). However, the average particle size of CoO_x on the oleum treated graphite is 2.5 nm, which is a little larger than that (ca. 1.6 nm) loaded on the surface of GE-Ph-SO₃H. This means that surface wettability modification caused by oleum treatment can improve the dispersion and decrease the size of CoO_r to a certain extent, but its influence is not as obvious as that caused by the surface phenyl-sulfonation modification. This difference may lie in the complex effect of Co²⁺ ions with phenylsulfonic groups, as this may change the number of crystal nuclei as well as the concentration of Co²⁺ ions remaining in the solution, leading to different growth kinetics.

The states of the cobalt species existing on the samples were analyzed by Raman and XPS spectra. As seen in Fig. 1c, three prominent Raman peaks, located at 678, 512, and 471 cm⁻¹ can be observed for the $CoO_x@G-Ph-SN$, $CoO_x@G-O$ and CoO_x

phase, Co_3O_4 phase, or nano- Co_3O_4 phase.³⁵ This demonstrates that the grafted cobalt species differ from the single phase of CoO or Co_3O_4 . The valence state of CoO_x in CoO_x @G-Ph-SN is further investigated by Co 2p XPS (Fig. 1d), in which two prominent shake-up satellite peaks, indicative of Co^{2+} ions,³⁶ are clearly observed, and the Co 2p_{1/2}–Co 2p_{3/2} energy separation is approximately 16.0 eV. Based on our fitted curves of Co 2p_{3/2} peak, the surface atomic ratios of $\text{Co}^{3+}/\text{Co}^{2+}$ are calculated to be ca. 0.35, 0.38 and 2.0 for CoO_x @G-Ph-SN, CoO_x @graphite and CoO_x @G-Ph-SN can be further verified by EPR spectra (Fig. S6†).

To characterize further the CoO_x particles on graphite, we carried out XRD and HRTEM measurements. No obvious XRD peaks assigned to the CoO_x can be observed even for the sample with a CoO_x loading content up to ca. 10.0 wt% (Fig. S7†). Based on the HRTEM image (Fig. S8†), the space distance of the CoO_x can be calculated to be 0.251 nm, which is not in accordance with those of CoO , $\mathrm{Co}(\mathrm{OH})_2$, CoOOH and $\mathrm{Co}_3\mathrm{O}_4$. Thus, the CoO_x phase in this work should be different from each of them. Based on the above analysis, we thus label the cobalt species in this work as CoO_x for simplicity. The surface element contents of CoO_x @graphite, CoO_x @G-O and CoO_x @G-Ph-SN samples were analyzed and are given Table S1.†

To evaluate the potential of the as-obtained $CoO_x@G-Ph-SN$ sample as a WOC, a visible-light-driven water oxidation system containing $[Ru(bpy)_3]Cl_2$ and $Na_2S_2O_8$ in the presence of a borate-buffered solution was examined with oxygen detected by a Clark electrode. Fig. 2a gives their typical activity curves as a function of reaction time, based on which their TOF values of water oxidation are calculated. As a comparison, Co_3O_4 -nano

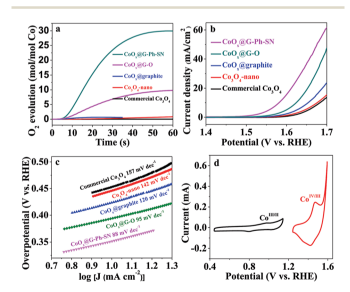


Fig. 2 Photochemical (a) and electrochemical (b–d) water oxidation performances of typical samples. (a) Time profiles of light-driven O_2 evolution catalyzed by various catalysts; (b) LSV curves of various catalysts in 1.0 M NaOH; (c) Tafel plots derived from (b). (d) Cyclic voltammograms (CVs) of $CoO_x@G-Ph-SN$ in 1.0 M NaOH at a scan rate of 10 mV s⁻¹. Reaction conditions: 0.5 mg catalyst, 3 mL H_2O , $[Ru^{II}(bpy)_3]Cl_2$ (1.0 mM), and $Na_2S_2O_8$ (5.0 mM) in borate buffer (80 mM, pH 9).

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(TOF of 0.012 s⁻¹) shows an obvious promotion of water oxidation activity compared to the commercial Co₃O₄ (TOF of 0.0013 s^{-1}). The photocatalytic water oxidation activity can be enhanced by the CoO_r@graphite sample, showing a TOF of 0.058 s^{-1} . The photocatalytic water oxidation activity can be further enhanced by improving the dispersion of CoO_r and decreasing the particle size of CoO_x . A TOF of 0.31 s⁻¹ can be achieved over CoO_r@G-O. Comparatively, the CoO_r@G-Ph-SN sample shows the best TOF value of 1.2 s⁻¹, an unexpectedly efficient water oxidation activity with respect to previously reported cobalt-based oxides (Table S2†). It should be pointed out that the TOF value is related to the light intensity (Fig. S9†). In addition, the water oxidation performances are normally affected by the surface reaction and charge separation processes, so factors beyond the size of CoO_r should have an effect on the activity; these probably include the mass transfer at the interface of the catalyst and aqueous solution, the structures of the active species and the the good conductivity of graphite etc.

The water oxidation performance was also evaluated by electrochemical water oxidation. Their linear sweep voltammetry (LSV) curves are depicted in Fig. 2b. The glassy carbon electrode (GCE) or graphite itself shows a negligible current and very high onset potential. Similarly, the water oxidation activity trend on typical electrodes can be described as follows: $CoO_x@G-Ph-SN > CoO_x@G-O > CoO_x@graphite > Co_3O_4-nano >$ commercial Co₃O₄, whose overpotential values at a current density of 10 mA cm⁻² are 350 mV, 395 mV, 450 mV, 470 mV, and 475 mV, respectively. The excellent O2-evolving activity of the CoO_x@G-Ph-SN composite was further confirmed by its much smaller Tafel slope (88 mV per decade) at lower overpotentials than that measured for CoO_x@G-O (95 mV per decade), CoO_x@graphite (120 mV per decade), Co₃O₄-nano (142 mV per decade), and commercial Co₃O₄ (157 mV per decade) (Fig. 2c). The cyclic voltammogram (CV) of CoO_x@G-Ph-SN in 1.0 M NaOH solution at a scan rate of 10 mV s⁻¹ (Fig. 2d) revealed two reversible reduction waves at $E_{1/2} = 0.89$ V and 1.44 V versus RHE, assigned to two sequentially occurring one-electron redox reactions involving Co^{II}/Co^{III} and Co^{III}/Co^{IV} couples, respectively.37-39

Encouraged by the extraordinary water oxidation activity on the CoO_x@G-Ph-SN sample, we thus further evaluated its potential use in a practical artificial photosynthesis system. As an initial attempt, we loaded the CoOx@G-Ph-SN onto the surface of a BiVO₄ electrode for photoelectrochemical water oxidation. Here the CoO_r@G-Ph-SN WOC can be considered as a cocatalyst of the BiVO₄ photoanode for water oxidation.⁴⁰ Fig. 3a gives the typical linear sweep voltammetry (LSV) curves of BiVO₄ photoanodes with and without loading of CoO_x@G-Ph-SN, in which loading of CoO_x@G-Ph-SN not only obviously promotes the current of the BiVO₄ photoanode, but also causes a negative shift of the onset potential. This clearly demonstrates that the CoO_x@G-Ph-SN is not only more active for water oxidation than BiVO₄ itself, but also efficient for the extraction of holes reaching the surface of BiVO4 for efficient transfer as well as promoted charge separation. The efficient transfer of photo-generated carriers between BiVO₄ and CoO_x@G-Ph-SN is

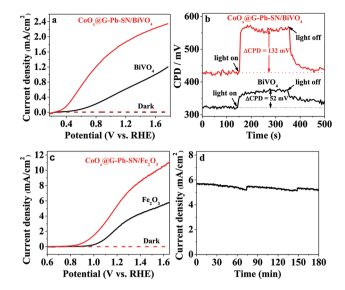


Fig. 3 Comparison of the photoelectrochemical performances and characterization of BiVO₄ and/or Fe₂O₃ photoanodes with and without loading of CoO_x@G-Ph-SN. (a) LSV curves of BiVO₄ photoanodes; 0.5 M lithium borate buffered solution (pH 9); 100 mW cm⁻² AM1.5G; scan rate of 50 mV s⁻¹. (b) Kelvin probe force microscopy of BiVO₄ photoanode. (c) LSV curves of Fe₂O₃ photoanodes; 1.0 M KOH; 300 W xenon lamp; scan rate of 50 mV s⁻¹. (d) Photoelectrochemical stability of CoO_x@G-Ph-SN/Fe₂O₃ photoanode at 1.23 V vs. RHE. Electrolyte: 1.0 M KOH; light source: 300 W xenon lamp.

confirmed by analysis of KPFM. As given in Fig. 3b, the steady state contact potential difference (Δ CPD) of the BiVO₄ photoelectrode with and without light irradiation is more significantly increased after the loading of CoO_x@G-Ph-SN.

The effectiveness of CoO_x@G-Ph-SN as a cocatalyst of water oxidation to promote the performance of artificial photocatalysts can be further revealed by loading it on the surface of another robust Fe₂O₃ photoanode. As revealed by the LSV curves of Fig. 3c, the loading of CoO_r@G-Ph-SN on the surface of Fe₂O₃ photoanode not only negatively shifts the onset potential of the photocurrent by ca. 150 mV, but also promotes the photocurrent at 1.23 eV (vs. RHE) by about 2.1 times. It is worth noting that the photocurrent of the CoO_x@G-Ph-SN/Fe₂O₃ electrode at 1.23 V vs. RHE can be maintained by over 90% after 3 h irradiation, indicating the good photochemical stability of the CoO_x@G-Ph-SN catalyst as a water oxidation cocatalyst in artificial photosynthesis. Together with the promotion of charge separation, we can reasonably deduce that CoO_x@G-Ph-SN, with high activity and stability, is a promising WOC for artificial photosynthesis.

Conclusions

In summary, ultrasmall CoO_x nanoparticles (around 1.6 nm) anchored on the surface of sulfonated graphite have been synthesized by a simple hydrothermal process. The success of the synthesis in reducing particle size mainly originates from the improvement of the surface hydrophilicity of graphite as well as its possible complex effect with Co^{2+} ions, leading to

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homogeneous dispersion and retarded growth. The as-obtained nanocomposite exhibits unexpected photochemical water oxidation activity with an optimal TOF of 1.2 s⁻¹, at least two orders of magnitude higher than that of conventional cobaltbased oxides. Furthermore, the CoO_r@G-Ph-SN can be grafted on the surface of artificial photocatalysts like BiVO₄ and Fe₂O₃ for promoted photoelectrochemical water oxidation, demonstrating its bright future in constructing solar-to-chemical energy conversion systems.

Acknowledgements

This work was supported by the Basic Research Program of China (973 Program: 2014CB239403), and the Natural Science Foundation of China (No. 21633009, 21522306, 21373210). F. Zhang thanks the priority support from the "Hundred Talents Program" of the Chinese Academy of Sciences.

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