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A molybdenum sulfo-oxide/cobalt oxysulfide Zscheme heterojunction catalyst for efficient photocatalytic hydrogen production and pollutant reduction†

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The existence of a heterojunction in the catalyst structure improves the efficiency of photocatalytic hydrogen production. The two phases of the heterojunction can be adjusted simultaneously by doping sulfur and oxygen to achieve a suitable energy band structure. Herein, we demonstrated a Mo(S,O)/ Co(O,S) Z-scheme heterojunction catalyst system (abbreviated as MoCoOS) with different Co(O,S) doping amounts by a simple one-step method. Doping of different atoms in the crystal changes the energy band structure which tunes the optical, electrical, and electrochemical properties of MoCoOS. MoCoOS-2 prepared at a Co precursor content of 20 mmol performed the best at a rate of 153.4 μ mol h^{-1} in photocatalytic hydrogen evolution, and it also showed excellent catalytic activities and good stability in reducing MO, MB, RhB, 4-NP and Cr^{6+} . This efficient MoCoOS-2 catalyst can have practical applications in photocatalytic hydrogen production and catalytic reduction of chemical pollutants.

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1. Introduction

For decades, the consumption of conventional sources of energy like fossil energy in industrial production and human life has been increasing while their reserves are decreasing, and the problems of energy depletion and environmental pollution are becoming more and more obvious. Carbon dioxide emissions from the combustion of oil and coal have aggravated global warming to a certain extent, and the emission of sulfur dioxide causes acid rain and other problems. Floating fine particles have polluted the air and cause respiratory diseases. In the process of fossil energy exploitation, the damage to the local geological environment and surface environment is deep and difficult to recover. These problems have forced people to have a strong demand for the development of new energy. Since the new century, researchers' exploration of new energy has entered a new stage. Clean, environment-friendly and efficient energy is the focus of researchers. Solar energy, hydropower, geothermal

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energy, wind energy, biomass energy and hydrogen energy, as new industries, have become supplementary means of energy supply sources and important measures of environmental protection and governance. Among them, hydrogen, as the most common element in nature, mainly exists as water. Hydrogen is one of the best energy carriers because of its high energy density. The calorific value of hydrogen is 142 351 kJ kg⁻¹, which is three times that of gasoline. Thus, it has very high conversion efficiency for both heat energy and electric energy. Hydrogen can also meet the needs of storage and application environments in the form of gas, liquid and solid hydrides. Therefore, as an excellent clean and renewable energy, hydrogen has become one of the most widely used energy sources. Methods such as steam reforming, thermo catalytic cracking, electro catalysis, and photocatalysis were developed to produce hydrogen.¹⁻⁴ But, photocatalysis has attracted much attention because of its simple hydrogen production process and low cost. Different catalytic materials such as metal sulfur oxides,5,6 solid solutions of sulfur oxides7 and phosphides,8 semiconductor polymers,9 g-C3N4,10 graphene,11 and the like were commonly used for photo catalytic hydrogen production. However, the two metals in metallic sulfur oxide catalysts often have unique synergistic effects, such as doping of the crystal lattice or the formation of a porous and large specific surface area leads to high efficiency for photo-catalytic hydrogen production. Poornaprakash et al. 12 synthesized CdS, CdS: Er (2 at%), and CdS: Er (4 at%) catalysts through a simple reflux

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route, and found that CDs: Er (2 at%) has excellent photocatalytic hydrogen production performance. Niu et al.13 synthesized and studied catalysts loaded with ZnS, Bi₂S₃ and CoS by a hydrothermal method, and found that CoS has excellent photocatalytic performance. Ghiat synthesized a high activity and durability SiO₂@NiCuPS photocatalyst by a hydrothermal method, which was a highly stable phyllosilicate material with a core@shell structure and high catalytic activity and stability.14 Chen et al.15 prepared a low-cost and high-efficiency catalyst for hydrogen production by photocatalytic decomposition of water with CuO/TiO2.

The existence of a heterojunction in the heterogeneous catalyst structure greatly improves the efficiency of catalytic hydrogen production. Electrons are excited from the valence band to jump into the conduction band under the illumination of radiation with the appropriate wavelength, in which different types of heterojunction structures play different roles and have different characteristics. 16-19 Z-scheme heterojunctions are widely investigated for their enhanced photocatalytic properties. By combining two semiconductors with suitable energy band structures, reducing holes and electron-hole recombination by transferring electrons of another semiconductor to a hydrogen-producing semiconductor, the hydrogen-producing ability is improved. This kind of heterojunction often forms between two phases that load sulfide or oxide of metal on CdS. Molybdenum is one kind of metal with rich mineral resources and low cost, and its sulfur oxides are often used in the study of heterojunction photocatalytic decomposition of water.²⁰⁻²³ The research of molybdenum-based metallic doping catalysts is one of the focuses of the hydrogen energy research field. The ultrasmall MoO₃/amine-CdS catalyst was obtained by electrostaticassisted ultrasonic exfoliation of its bulk counterpart with the help of dissolved amine-based polymers in Wang's work.24 Sharma et al.25 fabricated multifunctional composite photocatalysts by coupling sea urchin shaped ZnO with MoS2 and polyaniline sheets and a significant improvement of photocatalysis was obtained. Han et al.26 synthesized CdS@MoS2 with high photocatalytic performance. Goud et al.27 prepared an oxygen-vacancy rich, Bi₂O₃ based MoS₂/Bi₂O₃ Z-scheme heterojunction catalyst by an autoclave hydrothermal method using ethanol and water.

Likewise, the fast growth of industrialization has brought about environmental pollution which has currently become more serious. Factories such as leather factories, pharmaceutical factories, petroleum processing factories and metal smelting factories discharge various environmental pollutants into water bodies, causing damage to the water environment and having an adverse impact on human beings and animals. Organic dyes and nitro-aromatic compounds have high stability, low biodegradability, and hydrophilic nature that lead to their persistence in environmental compartments for a long time, cause the change of color and affect the normal function of water, affect human perception, hinder the photosynthesis of aquatic plants, and also lead to the destruction of the aquatic ecosystem. Pollution caused by nitroaromatic compounds is the most typical and extremely harmful organic pollution, not only because of their toxicity and mutagenicity, but also because of their chemical stability and wide use in various industries such as polymers, drugs, dves, rubber chemicals, pesticides, etc.28-32 Some heavy metal ions have extremely high toxicity, such as typical metal mercury, cadmium, lead, chromium and so on. Among them, the hexavalent chromium ion is extremely toxic to swallow/inhale, is easily absorbed by the human body, and enters the human body through the digestive tract, respiratory tract, skin and mucous membrane, causing damage to stomach, kidney and liver function, damaging eyes, causing retinal hemorrhage and optic atrophy, and causing cancer risk after long-term contact.33-37

Researchers have developed various methods like membrane flocculation, catalytic degradation, desorption and filtration to degrade or reduce pollutants in water.38-41 But, detoxifying or removing toxic chemical pollutants in water by using bimetallic catalysts is an efficient and comparatively cheap method. Kebede et al. 42 prepared a Sb-doped Mo(S,O)_{3-x} catalyst at 95 °C in a facile way, which showed high reduction efficiency towards MB. Ecer prepared a magnetic pumice (Fe₃O₄@PMC) composite catalyst with high reduction efficiency towards MO by a chemical co-precipitation method.43 Li et al.44 prepared MoS2@-TiO₂@poly(methyl methacrylate) by freeze hydrothermally treated electrospun PMMA nanofibers containing titanium n-butoxide and MoS2 nanosheets and degraded MO (10 mg L⁻¹, 100 mL) completely in 40 min under UV light irradiation. Li et al. 45 reported a general approach for the synthesis of ultrafine NiMoO_x NPs confined in mesoporous carbon with different morphologies and compositions using the replication method with SBA-15 as a hard template and studied their reduction efficiency towards 4-NP. Our research group synthesized Cu-based oxysulfide catalysts (CuBiOS, CuVOS, CuSbOS, etc.) via a feasible method, and they exhibited excellent catalytic reduction activity towards MB, MO, RhB, 4-NP and Cr(v1).46-51 For those bimetal oxysulfide compounds, they are mainly single phase compounds in the form of a solid solution of (M1, M2)(O,S) or together with a second non-oxysulfide to form the composite. 52-54 Composites of sulfur-doped oxide (oxysulfide)/oxygen-doped sulfide (sulfo-oxide) with a heterogeneous interface are a novel topic in terms of physical properties and the way of synthesis.

Because each semiconductor has its advantages and disadvantages when used as a hydrogen-producing catalyst, the energy band arrangement is particularly important. In order to improve the preparation of heterojunction photocatalysts, it is of great significance to design a heterojunction photocatalyst which can adjust the heterojunction two-phase energy band structure. On the basis of research by the same research group, 55,56 MoCoOS, a Z-scheme heterojunction catalyst prepared by simple hydrolysis of molybdenum and cobalt, has better hydrogen production capacity and excellent pollutant reduction capacity compared with other catalysts.

In this study we designed an oxysulfide heterojunction catalyst that can adjust the heterojunction dual-phase energy band structure. This MoCoOS catalyst is a Z-scheme catalyst in a composite form with sulfur-doped Mo₄O₁₁ and oxygen-doped Co₉S₈. The hydrogen production capacity and the reduction capacities of MO, MB, RhB, 4-NP, and Cr(vi) were studied. Meanwhile, the possible mechanisms of oxysulfide heterojunction catalysts with excellent activity were demonstrated.

2. Experiments

2.1. Preparation of Mo(S,O)/Co(O,S) catalysts

The synthesis of Mo(S,O)/Co(O,S) catalysts is as follows: 20 ammonium molybdate tetrahvdrate $((NH_4)_6$ Mo₇O₂₄·4H₂O, A.R., Tianjin Fuchen Chemical Reagents Factory) was dissolved in 800 mL of deionized water. After stirring for 20 min, 20 mmol cobalt chloride hexahydrate (CoCl₂·6H₂O, A.R., Aladdin Industrial Corporation) was added into the mixture solution and continuously stirred for 20 min. Then, 20 mmol thioacetamide (CH3CSNH2, A.R., Aladdin Industrial Corporation) was added into the mixture solution. After stirring for 30 min, the mixture solution was heated at 95 °C and kept for 2 h. Finally, the obtained solid was washed with deionized water and ethanol and dried, and it was labeled MoCoOS-3. In the same manner 5, 10, and 30 mmol CoCl₂-·6H₂O were used to synthesize MoCoOS-1, MoCoOS-2 and MoCoOS-4 catalysts, respectively. For comparison, samples without adding (NH₄)₆Mo₇O₂₄·4H₂O and CoCl₂·6H₂O were also prepared and labeled Co(O,S) and Mo(S,O), respectively.

2.2. Characterization of Mo(S,O)/Co(O,S) catalysts

Photoelectron spectroscopy (XPS) was conducted with a VG Scientific ESCALAB 250 XPS under Al Ka X-rays radiation, and carbon C 1s ($E_a = 284.6 \text{ eV}$) was used for calibration. The X-ray diffraction spectrum was obtained using a Rigaku X-ray diffractometer with a Cu K α radiation ($\lambda = 1.5406^{\circ}$) source. Field emission scanning electron microscopy (HITACHI SU-8010 microscope) and transmission electron microscopy (TECNAI G2 F20) were used to obtain morphologies. Ultraviolet-visible absorption spectra were obtained on a TU1901 spectrophotometer. The infrared spectrum was obtained by the KBr tableting method on a Fourier infrared spectrometer (Bruker VERTEX 70). An ASAP 2020 porosity and specific surface area analyzer was used to carry out N2 adsorption-desorption experiments. UPS and VB-XPS data were obtained by Ultraviolet Photoelectron Spectroscopy (RESOLVE120MCD5). A fluorescence spectrophotometer (F-380) was used to compare the photoluminescence effect of the catalyst. A Bruker A300 spectrometer was used to collect EPR signals from DMPO. The electron transfer pathway in the heterostructure was studied using an electronic paramagnetic spectrometer (Bruker A300). Solar power was measured using a full spectrum optical power meter (CEL-NP2000).

2.3. Electrochemical measurements

Electrochemical impedance spectra (EIS), transient photocurrent, cyclic voltammetry curves (CV), Mott–Schottky curves (MS) and linear sweep voltammetry (LSV) curves were obtained on an electrochemical workstation (Squid Stat Plus). An Ag/AgCl electrode, platinum sheet electrode and glassy carbon electrode were used as the reference electrode, counter electrode and working electrode to construct a three-electrode system. MoCoOS catalyst,

acetylene black and polytetrafluoroethylene were mixed in a mass ratio of 8:1:1 and a small amount of absolute ethyl alcohol was added, followed by grinding for 5 min to mix them fully. Then they were uniformly loaded onto a titanium mesh to obtain a sample with an area of $1.0~\rm cm^2$, which was kept in a drying oven at $85~\rm ^{\circ}C$ for $4~\rm h$ to be fully dried. Then, the prepared sample was connected with the working electrode and the measurement was carried out in $1.0~\rm mol~L^{-1}~KCl$ electrolyte at pH = 6.8. The frequency range of AC impedance measurements was set at $0.001~\rm kHz$ to $1000~\rm kHz$ with a potential amplitude of $10~\rm mV$. The Mott–Schottky test was carried out in the frequency range of $0.001~\rm kHz$ to $200~\rm kHz$. In the photocurrent response experiment, chronoamperometry was used to measure the photocurrent under the condition of stable current, and a $150~\rm W$ xenon lamp was used as the light source.

2.4. Catalyst activity test

2.4.1. Hydrogen evolution activity measurement. PHER was conducted on a Perfect Light Co. Labsolar-6A reaction system under a 150 W Xe lamp with a filter cut-off wavelength less than 420 nm. 20 mg MoCoOS catalyst, 10 mmol Na_2S , 10 mmol Na_2SO_3 and 50 mL deionized water were added into a 400 mL reactor for reaction. During the reaction process, the reactants were uniformly mixed by continuous stirring with a magnetic stirrer, and the temperature of the reaction system is maintained at room temperature by circulating cooling water. The amount of hydrogen was measured with a gas chromatograph (PANNA A91). The apparent quantum efficiency (AQE) was measured under the same photocatalytic reaction conditions with irradiation light through a band-pass filter of 420 nm and calculated according to the following equation:

$$\begin{split} AQE &= \frac{number\ of\ reacted\ electrons}{number\ of\ incident\ electrons} \times 100\% \\ &= \frac{number\ of\ evolved\ H_2\ molecules \times 2}{number\ of\ incident\ electrons} \times 100\% \end{split}$$

2.4.2. Reduction pollutant activity measurement. The catalytic activity of the MoCoOS catalysts was tested for the reduction of chemical pollutants like MB, MO, RhB, 4-NP and Cr(vI). For the MB test, 10 mg NaBH₄ and 5 mg of MoCoOS were added in to 100 mL of 50 ppm MB solution taken in a 100 mL beaker. Subsequently, 3 mL sample solution was taken from the reactor at a regular interval of time and the absorption was measured using a TU-1901 UV-vis spectrophotometer at room temperature in order to monitor the reduction progress of the resulting MB, and the concentration was calculated *via* the Lambert–Beer law. In the same procedure, the catalytic reduction of MO, RhB, 4-NP and Cr(vI) was also measured.

Results and discussion

3.1. Structure analysis

The surface chemistry and chemical state of each atom in the MoCoOS-2 catalyst and their atomic fraction were studied and analyzed by XPS. Fig. 1 shows the XPS fittings of Mo 3d, Co 2p, O 1s, and S 2p spectra of the MoCoOS-2 catalyst. Fig. 1a shows the

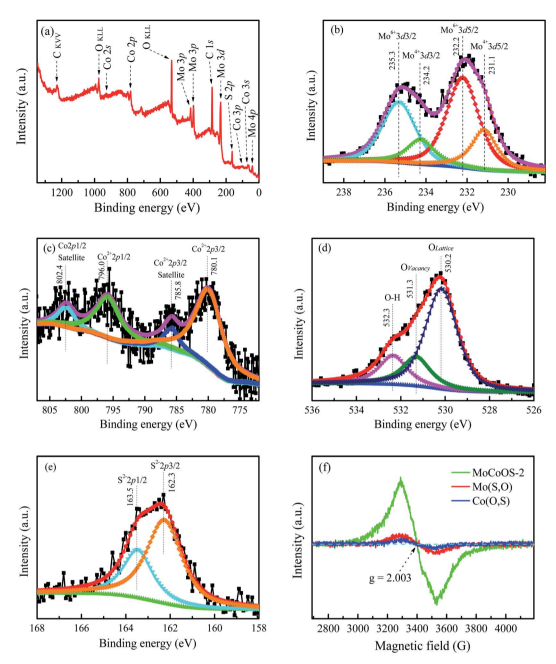


Fig. 1 (a) Survey spectrum and high resolution (b) Mo 3d, (c) Co 2p, (d) O 1s, and (e) S 2p XPS spectra of the MoCoOS-2 catalyst. (f) EPR spectra of the MoCoOS-2 catalyst, Mo(S,O), and Co(O,S).

survey XPS spectrum of MoCoOS-2. The Mo, Co, O, S, and C elements were observed in the spectrum with a trace amount of C 1s from foreign carbon. Fig. 1b shows the Mo 3d XPS spectra of MoCoOS-2 and the peaks located at binding energies of 232.2 eV and 235.3 eV correspond to the Mo $3d_{5/2}$ and Mo $3d_{3/2}$ of Mo(vi), respectively, with a spin-orbit splitting of 3.1 eV. 57,58 Similarly, the peaks located at 231.1 eV and 234.2 eV are assigned to Mo $3d_{5/2}$ and Mo $3d_{3/2}$ of Mo(IV), which indicates that Mo exists in the form of Mo(vI) and Mo(IV) in the catalyst. 59,60 For Co 2p of MoCoOS-2 shown in Fig. 1c, the peaks positioned at 780.1 eV and 796.0 eV belong to Co 2p_{3/2} and Co 2p_{1/2} of Co(II), respectively, whereas the peaks at 785.8 eV and

802.4 eV are attributed to Co 2p_{3/2} and Co 2p_{1/2} satellite peaks.⁶¹ Fig. 1d shows the O 1s XPS spectra of MoCoOS-2 and the asymmetric shape of the O 1s peak indicates the existence of different chemical states of oxygen in the catalyst. The peaks located at 530.3 eV, 531.3 eV, and 532.2 eV are attributed to lattice oxygen, vacancy oxygen, and hydroxyl oxygen, respectively.52,62-64 According to the fitting peak area, the ratios of lattice oxygen, oxygen vacancies, and hydroxyl oxygen were calculated and the results are given in Table S1.† As observed from Fig. 1e, the S 2p peaks at binding energies of 162.4 eV and 163.5 eV correspond to S $2p_{1/2}$ and S $2p_{3/2}$ of S²⁻ states, 65-67 respectively, without other sulfur forms. The proportion of each

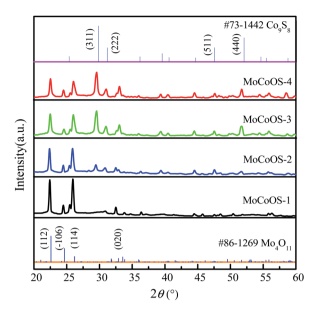


Fig. 2 XRD diffraction patterns of MoCoOS, the Mo_4O_{11} standard of PDF 86-1269, and the Co_9S_8 standard of PDF 73-1442.

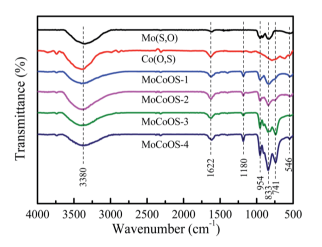


Fig. 3 FTIR spectra of Mo(S,O), Co(O,S), and MoCoOS prepared with different amounts of CoCl $_2\cdot$ 6H $_2$ O.

element in the MoCoOS-2 catalyst is calculated from their peak area and listed in Table S1.† As shown in Table S1,† the MoCoOS-2 catalyst has the highest proportion of oxygen vacancies, which may be related to the activity of the catalyst. For comparison, the XPS spectra of Mo(S,O) and Co(O,S) are displayed in Fig. S1 and S2,† respectively. All the XPS results after peak deconvolution for composition analysis are shown in Table S1.† In order to further verify the existence of oxygen vacancy defects for the MoCoOS-2 catalyst, the EPR was employed and the result is presented in Fig. 1f. It is clear that MoCoOS-2 shows a strong characteristic signal at g=2.003, which confirms the existence of oxygen vacancy defects in the MoCoOS-2 catalyst. However, the EPR intensities of the characteristic signals of Mo(S,O) and Co(O,S) catalysts are quiet lower than that of the MoCoOS-2 catalyst. From previous

studies, it was found that oxygen vacancies can reduce the band gap by forming a defect state below the conduction band or act as photogenerated electron capture centers, thereby inhibiting electron-hole recombination, promoting the transfer of electrons trapped at the defect sites, and improving the catalytic efficiency.^{53,72-74}

Fig. 2 shows the X-ray diffraction pattern of MoCoOS. The main 2θ peak positions at 22.387°, 24.419°, and 26.048° correspond to the (112), (-106), and (104) crystal planes of Mo_4O_{11} (#86-1269) while the peaks at 29.532°, 30.968°, 57.624° and 52.145° correspond to Co_9S_8 (#73-1442). The peaks of Co_9S_8 were not observed in MoCoOS-1 when the smallest amount of CoCl₂·6H₂O was added. When the added amount of CoCl₂-·6H₂O increased, the diffraction peaks appeared at 29.532°, 30.968°, 57.624° and 52.145° and their intensity also gradually increased. However, the peaks at 22.387°, 24.419°, and 26.048° have lower intensity due to the decreased amount of the Mo₄O₁₁ phase in the MoCoOS catalyst. The average crystallite size of the synthesized MoCoOS catalysts is calculated using the Scherrer equation with JADE software and is given in Table S1.† Using the peak at 30.968°, the crystallinity was calculated to be 78.93%, 82.41%, 90.68% and 87.26% for MoCoOS-1 to MoCoOS-4. Fig. S3† shows the XRD diffraction patterns of (a) Mo(S,O) and the Mo₄O₁₁ standard of PDF 86-1269 and (b) Co(O,S) and the Co₉S₈ standard of PDF 73-1442.

Fig. 3 shows the Fourier infrared spectra of MoCoOS. The higher intensity peak located at 833 cm⁻¹ was attributed to the stretching of Mo-O.75 The peak at 958 cm⁻¹ could be assigned to Mo=O.76 The peaks at 833 and 958 cm⁻¹ were from the crystalline structure of Mo₄O₁₁, whereas 546 and 741 cm⁻¹ were obviously the metal sulfide peaks from the Co₉S₈ phase.⁷⁷ The broad absorption band situated at 3380 cm⁻¹ and the peak at 1622 cm⁻¹ belong to the stretching vibration and bending vibration of the hydroxyl group, respectively.78,79 The higher intensity peak located at 1180 cm $^{-1}$ was attributed to S=0. The peaks of Mo(S,O) are consistent with the peaks of MoCoOS in the spectrum. The Co(O,S) phase is expected to be dissolved in Mo(S,O) to form the (Mo,Co)(S,O) solid solution. It can be clearly seen from the pattern that the overall intensity of the peaks tends to increase with the increase of CoCl₂·6H₂O dosage.

3.2. Morphology and BET analyses

The microstructure and morphology of the MoCoOS catalyst were examined by SEM. As observed from the SEM images of MoCoOS-2 shown in Fig. 4a and b, the rod-shaped Mo_4O_{11} phase formed by crystal stacking has the decorated Co(O,S) covered on its surface. Fig. 4b shows one end of the Mo_4O_{11} -phase nanorods to demonstrate the stacking of uniform rectangular sheet structures layer by layer into rectangular nanorods. The thickness of the nanosheets is about 15–20 nm and the thickness of the nanorods after stacking varies from 500 nm to 1000 nm. To further study the microstructure of the catalyst, the MoCoOS-2 catalyst was analyzed by transmission electron microscopy (TEM). Fig. 4c and d present the microstructural morphology of Mo(S,O) and Co(O,S), respectively. It is obvious

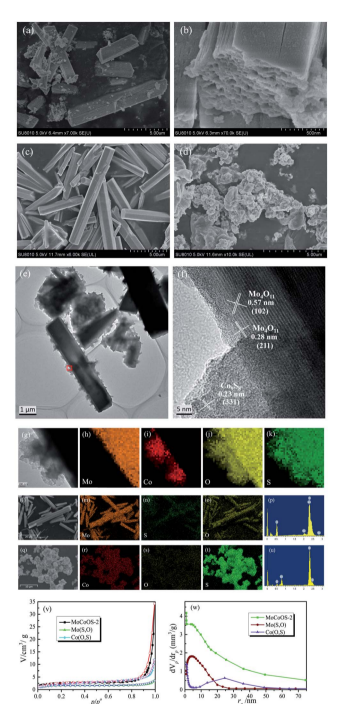


Fig. 4 (a and b) SEM images of MoCoOS-2. SEM images of (c) Mo(S,O) and (d) Co(O,S). (e) TEM and (f) HR-TEM images of MoCoOS-2. (g-k) EDX element distribution mapping images of Mo, Co, O, and S for MoCoOS-2. (1-p) EDX element distribution mapping images of Mo, O, and S for Mo(S,O). (q-u) EDX element distribution mapping images of Co, O, and S for Co(O,S). (v) Nitrogen adsorption-desorption isotherm and (w) the pore size distribution curve of the catalysts.

that the rod-shaped Mo(S,O) and nanoparticle-type Co(O,S) are consistent with the SEM image of MoCoOS. Fig. 4e shows the low magnification transmission electron microscopy image of the MoCoOS-2 catalyst, which is consistent with the SEM image. Under high magnification, many stacked nanosheets can be

displayed with a stripe dimension of about 0.8 nm, which is not the lattice spacing but the size of nanosheets. As shown in the HR-TEM image in Fig. 4f, 0.57 nm and 0.28 nm of lattice fringe values correspond to the (102) and (211) crystal orientations of the Mo₄O₁₁ phase, respectively. But, the lattice space of 0.23 nm corresponds to the (331) crystal orientation of the Co_9S_8 phase. These results confirm the existence of interfacial contact between Mo(S,O) and Co(O,S). In addition, the EDX elemental maps of the MoCoOS-2 catalyst demonstrate the homogeneous dispersion of Mo, Co, O, and S elements (Fig. 4g-k), which further confirms the incorporation of S into the Mo₄O₁₁ phase to form Mo₄(S,O)₁₁ or Mo(S,O) in an abbreviated form, and O into the Co_9S_8 phase to form $Co_9(O,S)_8$ symbolized as Co(O,S).

The elemental percentages of the MoCoOS catalysts were measured by XPS, XRF and EDX, and the results are listed in Tables S1-S7.† The results of XPS element ratio analysis are not accurate due to its small sampling area. So the EDX element ratio analysis data with a larger sampling area are used as the supporting data of element proportion. The proportion of Co in the sample increased significantly with the increased amount of CoCl₂·6H₂O precursor, and the values of Mo/Co in MoCoOS-1, MoCoOS-2, MoCoOS-3, and MoCoOS-4 are about 1.53, 1.37, 1.24 and 0.93, respectively. Therefore, when the amount of Co is excessive, it is easy to phase separate it from the composite state. The EDX elemental mapping images of Mo(S,O) and Co(O,S) are displayed in Fig. 4l-u. The morphological images of rod-like Mo(S,O) and nanoparticle-type Co(O,S) are consistent with that of the MoCoOS-2 catalyst. In order to further prove the influence of oxygen vacancies on photocatalysis, the element contents of Mo(S,O) and Co(O,S) in MoCoOS-1, MoCoOS-2, MoCoOS-3, and MoCoOS-4 were measured by EDX, as shown in Fig. S4 and Tables S4, S5, S6, and S7,† respectively. Five locations were checked on Mo(S,O) and Co(O,S) to measure the element content and the average values at each location were calculated with A, B, C, D, and E data points from Mo(S,O) and F, G, H, I, and J data points from Co(O,S). The molar ratios of S doping in Mo(S,O) for MoCoOS-1, 2, 3 and 4 are 11.75%, 9.58%, 11.38% and 12.39%. The molar ratios of O doping in Co(O,S) for MoCoOS-1, 2, 3 and 4 are 4.77%, 5.94%, 4.40%, and 4.94%. The molar ratios of Co(O,S) or $n[Co(O,S)]/\{n[Co(O,S)] + n[Mo(S,O)]\}$ in MoCoOS-1, 2, 3 and 4 are about 0.281, 0.394, 0.448 and 0.585, respectively. EPR analysis in Fig. 1f shows that MoCoOS exhibits abundant oxygen vacancies. The existence of sulfur on molybdenum oxide rods might promote the generation of O vacancies. The oxygen vacancies present on the surface of the photocatalyst can activate water molecules and exist as active sites.52,53

The nitrogen adsorption-desorption isotherm of the MoCoOS-2 catalyst is displayed in Fig. 4l and the curve is consistent with the type IV isotherm of the hysteresis loop at relative pressure (p/p^{o}) between 0.7 and 1.0. Fig. 4m displays the BJH pore size distribution of MoCoOS-2, Mo(S,O) and Co(O,S). The S_{BET} , total pore volume, and average pore diameter of MoCoOS-2 are 8.5 m² g⁻¹, 0.053 cm³ g⁻¹, and 24.98 nm, respectively. S_{BET} values of MoCoOS, Mo(S,O) and Co(O,S) are listed in Table S1.† Both the average particle size and specific surface area decreased for samples prepared with a high

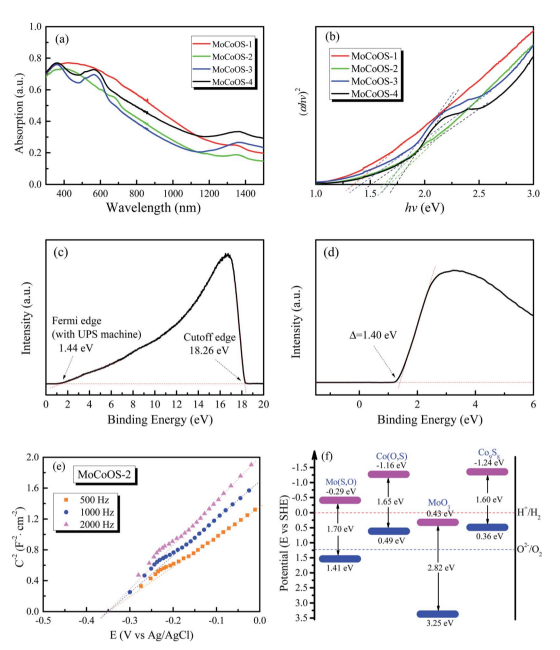


Fig. 5 (a) UV-vis absorption spectra of MoCoOS. (b) The $(\alpha hv)^2 - hv$ plots from the ultraviolet absorption spectra. (c) UPS spectrum of MoCoOS-2. (d) XPS-VB spectrum of MoCoOS-2. (e) Mott–Schottky curve of MoCoOS-2. (f) Band structure alignment of catalysts.

 $CoCl_2 \cdot 6H_2O$ content. Therefore, MoCoOS-1 has shown a larger specific surface area.

3.3. Bandgap structure analysis

The absorption and optical properties of the MoCoOS catalysts were studied using the UV-vis spectroscopic technique in the wavelength range of 200–2000 nm. Fig. 5a shows the ultraviolet and near infrared absorption spectra of MoCoOS-1, 2, 3, and 4. The bandgaps of the samples were calculated with the formula of $(\alpha h \nu)^{\frac{1}{n}} = k(h \nu - E_g)$ in the classic Tauc approach, where α , h, k, $h \nu$, and E_g are the absorption coefficient, Planck constant, absorption constant of direct transition, absorption energy, and bandgap energy, respectively. α can be replaced by absorbance

here. When the indirect band gap formula with n=2 is used, the bandgap of Co(O,S) in MoCoOS is 0.48 eV (Fig. S5†), which is quite different from the bandgaps in previous studies and is not in line with Co₉S₈. S2-84 Therefore, n=1/2 was adopted in the equation.

It can be seen from Fig. 5b that the band gap of MoCoOS-2 is 1.68 eV. Two bandgap values observed in the UV-vis absorption curves of MoCoOS-3 and MoCoOS-4 support the co-existence of MoCoOS and Co(O,S), which were formed due to partial phase separation from the precipitates leading to the appearance of absorption peaks. The bandgap values of Co(O,S) in MoCoOS-3 and MoCoOS-4 are 1.66 and 1.73 eV, respectively. The change of bandgap width may be caused by the doping amount of oxygen,

as shown in Fig. 4j. For comparison, the UV-vis absorption spectra of Mo(S,O) and Co(O,S) and the $(\alpha hv)^2$ -hv plots from the UV-vis absorption measurements are displayed in Fig. S5.† The bandgap values of Co(O,S) are close to those in MoCoOS-3 and MoCoOS-4. The slightly different bandgap values of Co(O,S) from the Co₉S₈ phase in MoCoOS can be related to the doping of O atoms into the Co₉S₈ lattice, as evidenced by the graph in Fig. 4i. The excess Co(O,S) second phase in MoCoOS can't play a good role in photocatalytic hydrogen production because it alone is not an effective catalyst for hydrogen evolution, which is also in accordance with our experimental data. The hydrogen production of the catalyst begins to decrease when superfluous CoCl₂·6H₂O is added during the catalyst synthesis. When the amount of CoCl2·6H2O added in the synthesis process is appropriate, the best combination of MoCoOS composite catalysts can be reached with the best match in the band level position of two compounds, which is beneficial for the electron transfer and its lifetime. The Co(O,S) in the composite catalyst also extends the light absorption to a wide wavelength range to benefit photocatalysis (Fig. 5a).

The main content of Mo(S,O) has a Mo₄O₁₁ phase as described in the previous XRD pattern. As the band gap of 1.70 eV for Mo(S,O) is much lower than that of \sim 2.8 eV for the MoO₃ structure,82 it is speculated that the doping of sulfur atoms into the Mo₄O₁₁ lattice leads to the reduction of the band gap. The band gap of Co(O,S) in MoCoOS-2 is \sim 1.65 eV, which is slightly higher than the reported value of 1.49-1.6 eV in the literature.83,84 The slightly higher value can also be related to the limited doping of oxygen atoms in Co₉S₈. To establish the band diagram of the composite catalyst systems, the MS tests were conducted for each catalyst, as shown in Fig. S7.† From the MS test, the flat band potential (V_{FB}) values adding to the gap value between the SHE and Ag/AgCl electrode are -1.16 eV for Co(O,S) and -0.29 eV for Mo(S,O), respectively. Due to the fact that the gap value between the SHE and Ag/AgCl electrode is around 0.20 eV: $V(\nu s. \text{ SHE}) = V(\nu s. \text{ Ag/AgCl}) + 0.197 \text{ eV} + 0.0591$ pH. This value is similar to the value (0.20 eV) of the gap between $V_{\rm FB}$ and the conduction band minimum potential $(E_{\rm CB})$ for the n-type semiconductor, so the equation is written as: $V_{\rm CB} = V_{\rm FB} - 0.20$ eV + 0.20 eV. Therefore, the CBM values of Mo(S,O) and Co(O,S) are around -1.16 and -0.29 eV. Co(O,S)not only shows a value of −1.16 eV close to −1.24 eV for Co₉S₈ but also has a similar band gap width. Co(O,S) is expected to be similar to Co₉S₈ with a limited amount of substituted oxygen.83,84 The Mott-Schottky curve of MoCoOS-2 is also presented in Fig. 5e, and the $E_{\rm CB}$ is obtained as -0.35 eV. Furthermore, UPS was used to determine the work functions of MoCoOS, Mo(S,O), and Co(O,S), and helium I α was applied as the ultraviolet source (hv = 21.22 eV). As shown in Fig. 5c and d, the binding energies of the secondary electron cutoff edge (E_C) and Fermi level edge $(E_{\rm F})$ are 18.26 eV and 1.44 eV. The equation used to calculate the work function is expressed as $\Phi = hv$ $-E_{\rm C}+E_{\rm F}$. The work function of MoCoOS-2 was determined to be 4.40 eV. The XPS-VB spectrum of MoCoOS-2 was conducted to further verify the valence band maximum (V_{BM}) . As presented in Fig. 5d, the gap between the Fermi level and $V_{\rm BM}$ is 4.80 eV. Therefore, the energy level of $V_{\rm BM}$ in MoCoOS-2 is 5.80

eV. As shown in Fig. S7,† the V_{BM} of Mo(S,O) and Co(O,S) is calculated using the same approach and the values are 5.87 eV and 4.89 eV, respectively. The E_{VB} values of MoCoOS, Mo(S,O) and Co(O,S) in a vacuum are converted into the standard hydrogen electrode (SHE) scale according to the standard: 0 eV vs. -4.44 eV. Therefore, the E_{CB} values of MoCoOS, Mo(S,O) and Co(O,S) are 1.36, 1.43, and 0.45 eV. The results are consistent with those of UV and MS tests. The aligned band structure of the catalysts is shown in Fig. 5f.

3.4. Electrochemical analysis

In order to further study the electrochemical characteristics of the MoCoOS catalyst, electrical impedance spectroscopy (EIS), cyclic voltammetry (CV), and multi-frequency Mott-Schottky (MS) and transient photocurrent (TPC) tests were used to study the charge transfer kinetics in the three-electrode system. The diameter of the semicircle in the Nyquist diagram is a useful feature for evaluating charge transfer kinetics in EIS analysis. After installing a simple Randle circuit composed of charge transfer resistance (R_{ct}) , electrolyte resistance (R_s) , constant phase element (CPE) and diffusion impedance (W), Z-View was used to fit EIS data, and the fitted graph is shown in the pattern. As observed from Fig. 6a with the amount of Co₉S₈ phase increased with the Co precursor, the semicircle in the Nyquist diagram shows an overall increasing trend which indicates that MoCoOS with a higher Co content leads to higher charge transfer resistance except for MoCoOS-1 due to an undetectable amount of Co(O,S). The resistance values of Mo(S,O), Co(O,S), MoCoOS-1, MoCoOS-2, MoCoOS-3 and MoCoOS-4 are 96.3 Ω , 584.6 Ω , 2.30 Ω , 1.62 Ω , 12.57 Ω and 21.46 Ω , respectively. MoCoOS-2 has the lowest resistance value, which can be beneficial to photocatalysis. The huge difference in resistance between MoCoOS and the control group proves that the formation of the heterostructure not only reduces the electronhole recombination rate, but also greatly reduces the electron transfer resistance, which is beneficial for the electrons to reach the surface. Fitting parameters of MoCoOS, Mo(S,O), and Co(O,S) are listed in Table S2.†

Fig. 6b shows the current densities of MoCoOS catalysts, Mo(S,O) and Co(O,S) at a scan rate of 50 mV s⁻¹. It is obvious that the MoCoOS-2 catalyst has a highest current density. Current densities of MoCoOS are much higher than those of Mo(S,O) and Co(O,S). Fig. 6c shows the current density of the MoCoOS-2 catalyst at scan rates of 10–50 mV s⁻¹, and the peak specific current increases slightly with the increase of scan rate, which indicates that the MoCoOS nanoparticle electrode shows good rate performance.85 The current density plots of Mo(S,O) and Co(O,S) at scan rates of 10-50 mV s⁻¹ are shown in Fig. S8.† Fig. 6d shows the specific capacitance of the MoCoOS catalyst with the value calculated by the following equation:

$$C_s = \frac{\int I \mathrm{d}V}{AS\Delta V}$$

In this equation, C_s is the specific capacitance, $\int I dV$ is determined by the integral area under the CV curve, I is the current, Vis the voltage, S is the scan rate, and A is the active area of the

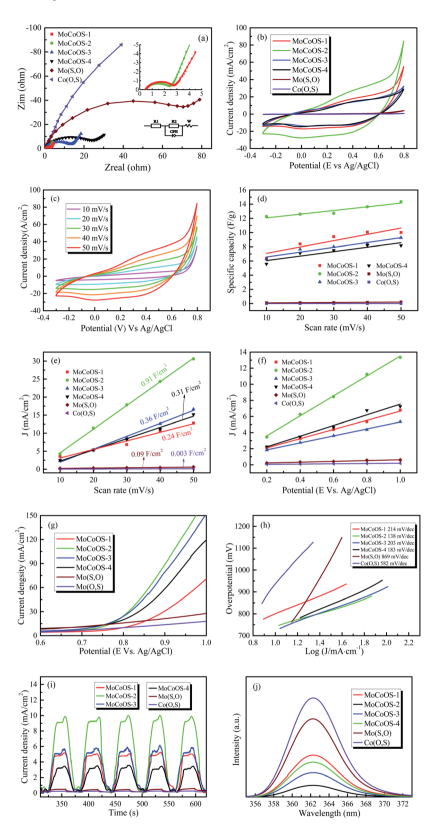


Fig. 6 (a) Electrochemical impedance spectra of MoCoOS with different Co contents. (b) Current density of different MoCoOS under 50 mV s $^{-1}$ scan rate. (c) Current density of MoCoOS-2 under different scan rates. (d) Specific capacity of MoCoOS under different scan rates. The variations of current density with (e) scan rate and (f) different voltages. (g) LSV, (h) Tafel curves, (i) transient photocurrent (TPC), and (j) PL spectra of Mo(S,O), Co(O,S), and MoCoOS catalysts.

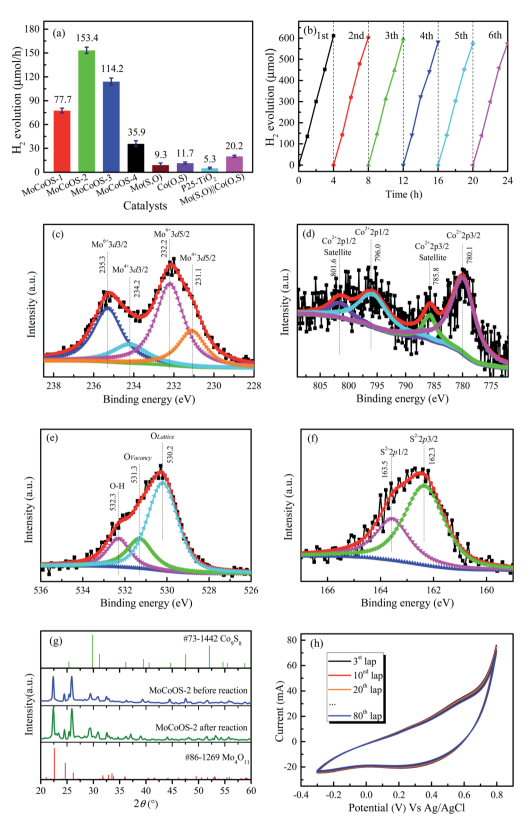


Fig. 7 (a) Hydrogen production of MoCoOS, Mo(S,O), Co(O,S), Mo(S,O)||Co(O,S) and P25-TiO $_2$ catalysts. (b) Recycling hydrogen production reaction performance of MoCoOS-2. (c-f) XPS spectra of MoCoOS-2 after the hydrogen evolution reaction. (g) XRD patterns of MoCoOS-2 before and after the hydrogen evolution reaction. (h) Cyclic voltammetry of MoCoOS-2 for 80 cycles.

MoCoOS electrode. When the scan rate is 50 mV s⁻¹, the specific capacitances of Mo(S,O), Co(O,S), MoCoOS-1, MoCoOS-2, MoCoOS-3 and MoCoOS-4 are 0.366, 0.204, 12.0, 13.3, 10.3 and 9.18 F g⁻¹, respectively. ⁸⁶ It can be seen from the figure that with the increase of scan rate, the capacitance slightly increases due to the fact that the number of ions contained in the electrolyte transmitted to the electrode increases. As shown in Fig. 6e, MoCoOS-2 exhibits a much larger $C_{\rm dl}$ of 0.91 F cm⁻² than the others, indicating the formation of active sites by creating surface oxygen vacancies in the MoCoOS-2 catalyst (Table S1†). Fig. 6f shows the current density of the catalyst under different voltages recorded by chronoamperometry. It can be seen from Fig. 6f that the catalyst has good stability when different voltages are applied.

Linear sweep voltammetry (LSV) is also an effective method to characterize the water decomposition ability of catalysts. The LSV curve of the catalyst is presented in Fig. 6g and it can be clearly seen from the figure that the MoCoOS-2 catalyst shows a higher current density at the same potential. Furthermore, Tafel slopes of MoCoOS-1, MoCoOS-2, MoCoOS-3, MoCoOS-4, Mo(S,O) and Co(O,S) obtained from the LSV diagram (Fig. 6h) are 214, 138, 203, 183, 869, and 582 mV dec^{-1} , respectively. MoCoOS-2 has the lowest overpotential of 138 mV dec^{-1} .

Illumination is the main driving force for the catalyst to generate holes and electrons in photocatalytic reaction. Fig. 6i shows the transient photocurrent (TPC) generated by Mo(S,O), Co(O,S), and MoCoOS catalysts prepared with different amounts of CoCl2·6H2O, under the illumination of incident photons. Chronoamperometry was used to record the response of MoCoOS to visible light. Under the voltage of 0.8 V, MoCoOS-2 showed the highest current density of 9.99 mA cm⁻². MoCoOS-1, MoCoOS-3, and MoCoOS-4 had relatively low current densities at 7.63, 7.33 and 7.24 mA cm⁻², respectively. The TPC values of Mo(S,O) and Co(O,S) are 0.49 and 0.24 mA cm⁻², respectively, which are much lower than those of the MoCoOS catalysts. TPC data are consistent with those with the EIS test, indicating that photoelectron-hole pairs can be effectively separated for the MoCoOS-2 catalyst. Fig. 6j shows the photoluminescence spectrum of the MoCoOS catalysts. The photoluminescence spectrum of the catalyst was obtained under excitation at 362 nm with a laser beam at a wavelength of 310 nm. MoCoOS-2 shows the lowest excitation intensity. The PL test indicates that the electron-hole coincidence efficiency of the MoCoOS-2 catalyst is the worst. Of all MoCoOS catalysts, MoCoOS-2 shows the best properties of carrier migration and photocurrent response, and thus an improved photocatalytic reaction can be expected.

3.5. Photocatalytic hydrogen evolution activity and stability

The photocatalytic performance of the prepared catalysts was evaluated under irradiation by 150 W filtered ultraviolet light. To ensure the reproducibility of the catalysts, each catalyst was prepared three times and tested for hydrogen production under the same conditions. Then the average value of hydrogen production was obtained. Fig. 7a shows the hydrogen evolution

performance of the MoCoOS, MoSO, and CoOS catalysts. As all catalysts have a certain degree of catalytic hydrogen production, MoCoOS-2 shows the highest hydrogen rate of 153.4 $\mu mol~h^{-1}$ which is 16 and 12 times higher than that of MoSO and CoOS, respectively. MoCoOS-2 had the strongest current response and the lowest electric resistance in electrochemical impedance testing. Therefore, MoCoOS with a suitable doping amount with the CoCl $_2\cdot 6H_2O$ precursor can effectively improve the hydrogen production efficiency. As the maximum hydrogen production rate of MoCoOS-2 can reach 153.4 $\mu mol~h^{-1}$, the apparent quantum yield under visible light can reach 14.42%. The solar-to-hydrogen (STH) values were calculated according to the equation:

$$\mathrm{STH} = \frac{\mathrm{Output\ energy\ as\ hydrogen}}{\mathrm{Energy\ as\ solar\ light}} \times 100\%\ \frac{r_{\mathrm{H_2}} \times \varDelta G_{\mathrm{r}}}{P_{\mathrm{sun}} \times S} \times 100\%$$

 P_{sun} of solar light was tested using a full spectrum optical power meter (CEL-NP2000) and calculated to be 920 W m^{-2} . The reaction was performed under irradiation and the area of the reactor was 12 cm². The rate of H₂ evolution is about 42.03 μmol h⁻¹. Based on this, the STH energy conversion efficiency of this system can be calculated as 0.25%. Moreover, the commercial P25-TiO₂ and the physical mixture of Co(O,S) and Mo(S,O) were used for photocatalytic hydrogen production to compare with the Z-scheme Mo(S,O)/Co(O,S). Mo(S,O) and Co(O,S) were ground and mixed in a molar ratio of 2:1 for 30 min, and the obtained mixture powder was abbreviated as Mo(S,O)||Co(O,S). After a 4 h hydrogen production test under the same conditions, the Mo(S,O)||Co(O,S) and commercial P25-TiO2 catalysts produced 20.2 and 5.2 μ mol h⁻¹ of hydrogen gas, respectively. The graph of hydrogen generation with time is also shown in Fig. S10.† Table S8† shows the comparison of the photocatalytic activities of hydrogen evolution for various catalysts containing oxides and sulfides of cobalt and molybdenum, as reported in the literature. The MoCoOS-2 catalyst shows superior activity under visible light. Six test cycles every four hours were performed to prove the excellent stability of hydrogen evolution reaction, as shown in Fig. 7b. After each round of reaction, the activity of the catalyst decreased slightly. To eliminate the influence of sulfate and thiosulfate ions, we centrifuged the mixed solution to retain the precipitated catalyst after the end of each reaction cycle. The catalyst after the reaction was added into a new solution with sacrificial agents (10 mmol Na₂S and 10 mmol Na₂SO₃) for a new test cycle. Fig. 7c-f display the XPS spectra of the MoCoOS-2 catalyst after the hydrogen production reaction. Comparing the spectra of each element in Fig. 1a-d, it can be seen that are no obvious changes. The XRD pattern of MoCoOS-2 after hydrogen evolution tests of 12 h shown in Fig. 7g also explains its stability. The redox reaction was measured by cyclic voltammetry for 80 cycles under the scan rate of 50 mV s⁻¹ and the potential window of -0.3 to +0.8 V. As shown in Fig. 7h the small oxidation and reduction peaks remain basically unchanged after 80 cycles. To sum up, MoCoOS-2 with excellent hydrogen evolution performance has good photocatalytic stability.

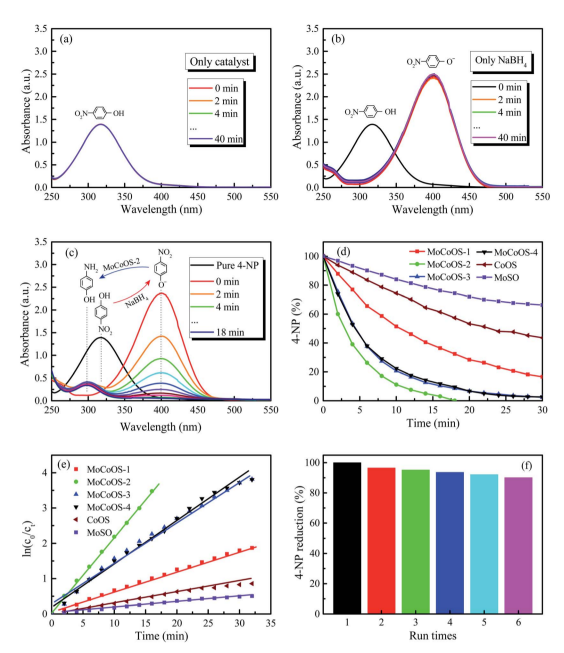


Fig. 8 (a) Reaction of 4-NP with MoCoOS-2. (b) Reaction of 4-NP with NaBH₄. (c) Reduction of 4-NP with MoCoOS-2 and NaBH₄. (d) Reduction of 4-NP with Mo(S,O), Co(O,S), and MoCoOS catalysts in the presence of NaBH₄. (e) The first order reaction rate curves of 4-NP. (f) Catalytic reduction efficiency after cycles of reactions.

3.6. Reduction of 4-NP

Before conducting heterogeneous catalytic reduction of 4-NP with NaBH4, the results of the NaBH4-free and the catalyst-free tests were evaluated. Fig. 8a and b show the catalytic inability of MoCoOS-2 in reducing 4-NP under dark conditions. Fig. 8c shows the reduction of 4-NP in the presence of MoCoOS-2 and NaBH₄ together. As shown in the pattern, 100 mL of 20 ppm 4-NP solution has almost completely reacted within 18 min in the presence of 5 mg of MoCoOS-3 catalyst and 50 mg of NaBH₄. During the experiment, the color of the solution changed from yellow to deep yellow with the NaBH4 dissolved gradually. Meanwhile, the original peak position of the ultraviolet spectrum at 317 nm shifted to 400 nm for the ionic state, and then shifted to 300 nm after conversion to 4-AP. With the hydrogenation proceeding, the intensity of the peak at 300 nm increased. Fig. 8d shows the conversion performance for all the MoCoOS catalysts. The MoCoOS-2 catalyst has the best catalytic effect. The first-order reaction kinetics data of reduction reaction can be calculated by using the data of Fig. 8d and the following formula:

$$\ln C_0/C_t = kt$$

 C_0 represents the initial concentration of 4-NP, C_t is the concentration of 4-NP dye in ppm measured at time (min) and k



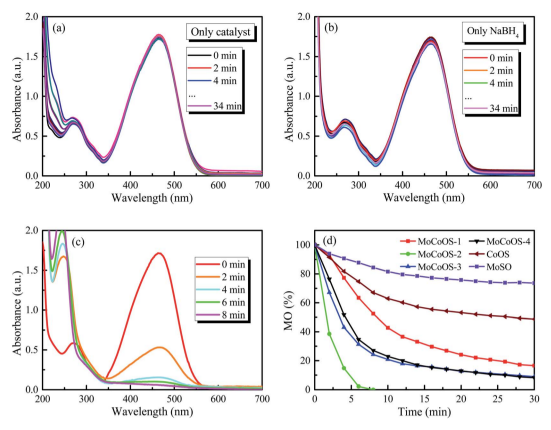


Fig. 9 (a) Reduction of MO with only MoCoOS-2. (b) Reduction of MO with only NaBH₄. (c) Reduction of MO with MoCoOS-2 and NaBH₄. (d) Reduction of MO with Mo(S,O), Co(O,S), and MoCoOS catalysts in the presence of NaBH₄.

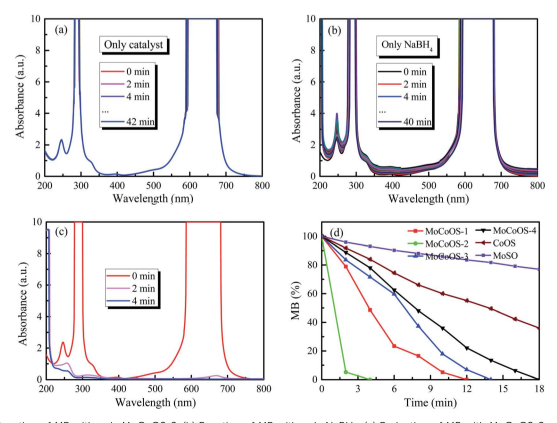


Fig. 10 (a) Reaction of MB with only MoCoOS-2 and NaBH₄. (d) Reduction of MB with MoCoOS-2 and NaBH₄. (d) Reduction of MB with Mo(S,O), Co(O,S), and MoCoOS catalysts in the presence of NaBH₄.

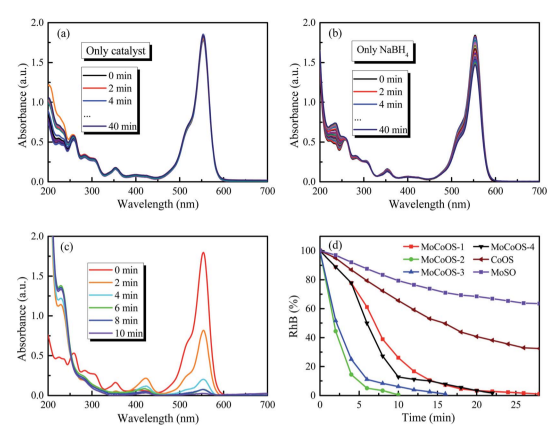


Fig. 11 (a) Reaction of RhB with only MoCoOS-2. (b) Reaction of RhB with only NaBH₄. (c) Reduction of RhB with MoCoOS-2 and NaBH₄. (d) Reduction of RhB with Mo(S,O), Co(O,S), and MoCoOS catalysts in the presence of NaBH₄.

(min⁻¹) is the apparent catalytic rate constant determined from the plot of $ln(C_0/C_t)$ versus illumination time, as shown in Fig. 8e. According to the figure, the kinetic constants of the reduction reaction of 4-NP with MoCoOS-1, MoCoOS-2, MoCoOS-3, and MoCoOS-4 are 0.0615, 0.211, 0.121, and 0.129 min⁻¹, respectively. The reaction constants of MoCoOS-2 are 3.43, 1.71, and 1.63 times higher than those of MoCoOS-1, MoCoOS-3, and MoCoOS-4, respectively. In order to further verify the practicability of the MoCoOS-2 catalyst, the catalyst was subjected to six runs. As shown in Fig. 8f, the catalytic efficiency only drops to 90.1% after the sixth run. Table S9† illustrates the comparison between some previous studies and the catalytic reduction of 4-NP by MoCoOS-2.

3.7. Reduction activity for MO, MB, and RhB

The UV-vis absorption patterns of the reduction of MO, MB and RhB by MoCoOS catalysts are shown in Fig. 9, 10 and 11, respectively. Throughout the reaction, 5 mg of MoCoOS catalyst was used to reduce 100 mL of 50 ppm organic dyes in the presence of 50 mg NaBH₄ as the reducing agent. Fig. 9a, 10a and 11a show the reduction of dye solutions in the presence of only the MoCoOS-2 catalyst for the reduction of dyes MO, MB, and RhB, respectively. The reduction reaction without the aid of NaBH₄ did not occur. Fig. 9b, 10b and 11b show the catalyst-free reduction reaction in the presence of NaBH4. There are no noticeable changes for reactions without heterogeneous

catalysts. According to Fig. 11b, the reaction rate is calculated to be 0.0086 ppm min⁻¹ per mg of NaBH₄. Fig. 9c, 10c, and 11c shows the reduction reaction of the three dyes with the MoCoOS-2 catalyst in the presence of NaBH₄. Fig. 9d, 10d and 11d show the reduction effects of MoCoOS catalysts for MO, MB and RhB, respectively. The MoCoOS-2 catalyst shows better reducibility than other samples under dark conditions. In the presence of both MoCoOS-2 catalyst and NaBH₄, almost complete reduction of MO, MB, and RhB was achieved within 8, 4, and 10 min, respectively. The first-order reaction rate constants of reduction reactions of MO with MoCoOS-1, MoCoOS-2, MoCoOS-3, and MoCoOS-4 catalysts are 0.066, 0.608, 0.102, and 0.095 min⁻¹; reduction rate constants of MB with MoCoOS catalysts are 0.250, 1.478, 0.176, and 0.118 min⁻¹; and reduction rate constants of RhB with MoCoOS catalysts are $0.159, 0.482, 0.280, \text{ and } 0.180 \text{ min}^{-1}, \text{ respectively. The reduc-}$ tion performance of the MoCoOS-2 catalyst for the dyes is obviously better than those of other catalysts.

3.8. Reduction of Cr(vi)

For the Cr(v1) reduction test, 5 mg of MoCoOS catalyst was used to reduce 100 mL of 20 ppm Cr(vi) in the presence of 50 mg NaBH₄ as the reducing agent. Fig. 12a and b show the reduction of Cr(vi) under the NaBH₄-free and the catalyst-free conditions, respectively. The results show that the use of either NaBH4 or MoCoOS catalyst alone does not cause the reduction of Cr(v1)

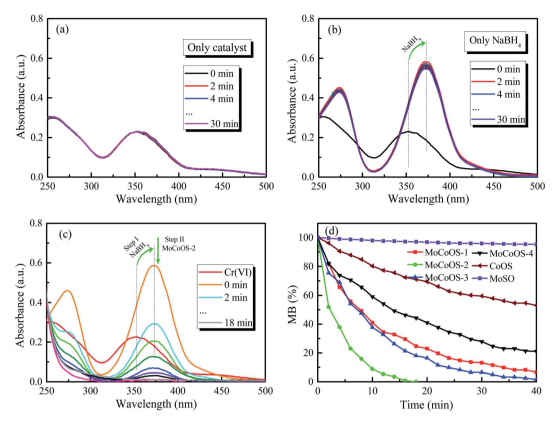


Fig. 12 (a) Reaction of Cr(vi) with only MoCoOS-2, (b) reaction of Cr(vi) with only NaBH₄, (c) reduction of Cr(vi) with MoCoOS-2 and NaBH₄, (d) reduction of Cr(vI) with Mo(S,O), Co(O,S), and MoCoOS catalysts in the presence of NaBH₄.

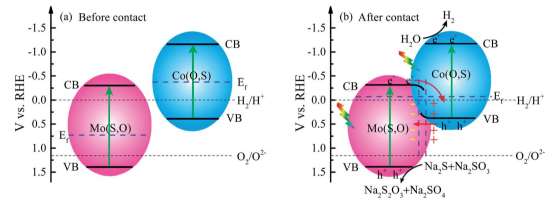


Fig. 13 (a and b) The energy band arrangement in the MoCoOS Z-scheme heterojunction

within 30 min. Fig. 12c shows that the peak intensity at 250 nm and 300 nm decreases for the catalytic reduction of Cr(v1) with the co-existence of the 5 mg MoCoOS-2 catalyst and 50 mg NaBH₄. Fig. 12d shows the catalytic reduction of Cr(v_I) by four MoCoOS catalysts, which have the first-order reaction kinetic rate constants of 0.069, 0.267, 0.098, and 0.047 min⁻¹. The firstorder kinetic constant of the MoCoOS-2 catalyst is 3.83, 2.73 and 5.72 times higher than those of MoCoOS-1, MoCoOS-3 and MoCoOS-4 catalysts, respectively. Table S10† illustrates the comparison between some previous studies and MoCoOS-2 catalytic reduction performance for Cr(vI).

For all the reduction reactions, the MoCoOS catalyst has shown an excellent catalytic effect for several typical water pollutants such as 4-NP, MO, MB, RhB, and Cr(vI). Compared with Ni@carbon,87 the reduction efficiency of hexavalent chromium is enhanced 4 times. In addition to the excellent catalytic activity, the prepared MoCoOS catalyst also shows high stability for repeated runs. To further study the influence of pH change in the reaction environment, we tested the reaction of 4-NP, MO, MB, RhB, and Cr(v1) with MoCoOS-2 under different pH conditions with the results shown in Fig. S11.† The reduction of pollutants can be enhanced at pH < 7. The photocatalytic

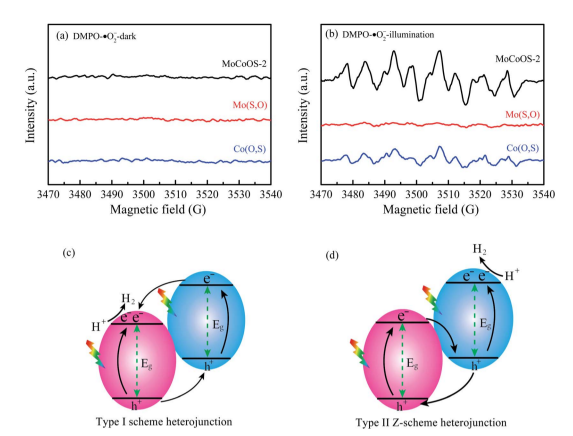


Fig. 14 EPR spectra of (a) DMPO-'O and (b) DMPO-'O₂- for Mo(S,O), Co(O,S), and MoCoOS catalysts under darkness and illumination for 10 min. (c and d) The two types of heterojunction.

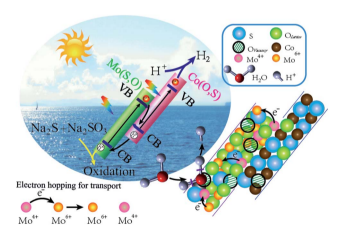


Fig. 15 Mechanism diagram of photocatalytic hydrogen production in the presence of the MoCoOS catalyst.

reduction can be completed within 20 min even under the most difficult conditions at pH = 9.

3.9. Mechanism of hydrogen evolution

Under the illumination of light at the appropriate wavelength, the photocatalyst could induce the formation of photogenerated electrons and holes in the conduction band and valence band, and then hydrogen can be produced. In this

study, a filtered xenon lamp with ultraviolet light below 400 nm was used as a light source. The band widths of MoCoOS-1, MoCoOS-3 and MoCoOS-4 catalysts are about 1.7 eV. The reasons for the different PHER rates of MoCoOS are related to the change in the Co precursor content to lead to the different amounts of Mo(S,O) and Co(O,S) in MoCoOS and different S contents in each Mo(S,O) and Co(O,S). With the optimal Co precursor content, the formed MoCoOS-2 catalyst displayed the best charge separation ability, longer charge lifetime, lower transport resistance and the best PHER rate. For Z-scheme photocatalytic water splitting, the reaction kinetics with the reaction steps listed below are explained. At the initial stage, both Mo(S,O) and Co(O,S) in MoCoOS with a bandgap less than 2.0 eV are excited by visible light to generate electron-hole pairs (eqn (1)). The electrons excited in MoCoOS lead to H₂ evolution as shown in eqn (2).

$$Mo(S,O)/Co(O,S) + h\nu \rightarrow Mo(S,O)/C(O,S)(e^- + h^+)$$
 (1)

$$2H_2O + 2e^- \rightarrow H_2 + OH^-$$
 (2)

$$SO_3^{2-} + 2OH^- + 2h^+ \rightarrow SO_4^{2-} + H_2O$$
 (3)

$$2S^{2-} + 2h^+ \to S_2^{2-} \tag{4}$$

$$S_2^{2-} + SO_3^{2-} \rightarrow S_2O_3^{2-} + S^{2-}$$
 (5)

$$SO_3^{2-} + S^{2-} + 2h^+ \rightarrow S_2O_3^{2-}$$
 (6)

$$S^{2-} + 2SO_3^{2-} + 2OH^- + 4h^+ \rightarrow S_2O_3^{2-} + SO_4^{2-} + H_2O$$
 (7)

During the photocatalytic water splitting, Mo(v) can be viewed as Mo(v) with two attached electrons to form a local plasmon. With the plasmonic interaction, the slightly bound electrons can hop through the catalyst with lower resistance for easy charge separation. The surface oxygen vacancy defects facilitate the effective separation of photogenerated carriers, ss, so and as they trap oxygen from water and make the O–H bond of the H_2O molecule weaker. 53,62,90,91 As for the added Na_2S and Na_2SO_3 , they not only avoid the photocorrosion of sulfide but also go through eqn (3)–(7) for the prevention of the excitation of electrons and holes.

As shown in UPS and XPS-VB spectra, the energy band structure is rearranged to form a new Fermi level after hybridization due to the different Fermi levels. The rearrangement leads to downward band bending of Mo(S,O) and upward band bending of Co(O,S) at the heterojunction. The electrons transfer from Mo(S,O) to Co(O,S) with accumulation of electrons on Co(O,S) and negative charge on Mo(S,O), while the excited holes transfer upward at the junction. A depletion region and an internal electrical field are created at the interface of the heterojunction in this process. The directed migration of photogenerated charges is increased. Conclusively, the electrons in the conduction band of Mo(S,O) transfer to the VB of Co(O,S) and quickly annihilate the holes. The mechanism of transfer leads to a higher separation efficiency. Energy band arrangements in the Z-scheme MoCoOS heterojunction are shown in Fig. 13a and b.

To further ascertain the mechanism of hydrogen production, production of 'OH and ' O_2 ⁻ radicals was investigated by the EPR signal spin-trap with DMPO under visible light irradiation. Because the E_{VB} values of Mo(S,O) and Co(O,S) are higher than the reduction potential of DMPO–'OH, no peak appears under both darkness and illumination. As shown in

Fig. 14a, no signals of DMPO-'O2 can be observed under darkness. After illumination for 10 min, the obvious characteristic DMPO-'O₂ radical peaks of MoCoOS-2 and Co(O₂S) arose, and the DMPO-'O2- characteristic peaks of MoCoOS-2 are higher than those of Co(O,S), which indicates that MoCoOS-2 produces more 'O2 radicals under illumination (Fig. 14b). 92-95 In addition, in Mo(S,O) no DMPO-'O2 signals appeared under illumination. Because the E_{CR} of Mo(S,O) is positive than -0.33 eV, 'O2 can't be produced. Thus, if MoCoOS-2 has the type I Z-scheme heterojunction (Fig. 14c), the DMPO-'O₂ signal intensity of MoCoOS-2 should be weaker than that of Co(O,S) due to the electrons transferred to the CB of Mo(S,O). Generation of fewer 'O₂ radicals from Co(O,S) can be related to the pathway for electron transfer, which involves electron transfer from the CB of Mo(S,O) to electronic hole vacancies in the VB of Co(O,S) and then excitation by illumination. Therefore, the path of e transfer in the MoCoOS-2 photocatalytic system supports the type II Zscheme heterojunction (Fig. 14d), and the kinetic mechanism diagram of photocatalytic hydrogen evolution of the MoCoOS catalyst is shown in Fig. 15.

3.10. The mechanism of pollutant reduction

Fig. 16 shows the possible reaction mechanism of the MoCoOS catalyst for catalytic reduction of pollutants in the water environment with sodium borohydride as the sacrificial agent. Firstly, NaBH₄ dissolves to generate borohydride ions, which adhere to the catalyst surface and covalently bond with it.⁹⁶ Simultaneously, pollutants are adsorbed on the surface of the MoCoOS catalyst and strongly interact with covalently bonded hydrogen atoms, and then electrons are transferred from donor BH₄⁻ to acceptor pollutants. The electron transfer occurs between Mo(IV) and Mo(IVI) in the catalyst to enhance the lifetime of electrons, which further promotes the reduction reaction.^{97,98} Finally, the pollutants are reduced with their products leaving the reaction site. In this reaction process, the catalyst

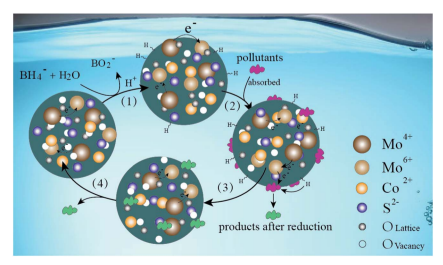


Fig. 16 The schematic reaction of organic dyes in the presence of the MoCoOS catalyst and NaBH₄.

provides a reaction site, and the valence charge-changing Mo ions provide a springboard for electron transfer.

4. Conclusions

MoCoOS catalysts with different Co(O,S) doping amounts were successfully synthesized and characterized as well as utilized for photocatalytic hydrogen production and chemical pollutant reduction. It was found that the synthesized MoCoOS catalyst was a composite consisting of two phases of Mo₄O₁₁ and Co₉S₈ with the incorporation of heteroatoms to form the Mo(S,O)/ Co(O,S) catalyst. The MoCoOS-2 catalyst was stable and had the highest hydrogen evolution with a rate of 153.4 μ mol h⁻¹. The successfully adjusted energy band structure of the two-phase heterojunction Mo(S,O)/Co(O,S) made the hydrogen production more efficient. In addition, MoCoOS-2 with the best PHER performance also displayed the fastest rates in the NaBH₄-aided reduction of MO, MB, RhB, 4-NP, and Cr(vi). Therefore, it is expected that this research of MoCoOS catalysts provides important insights on material design for hydrogen production and pollutant reduction by forming a heterojunction and improving the electron transport efficiency of the bimetallic catalyst.

Conflicts of interest

The authors declare that they have no known competing financial interests or personal relationships that could have appeared to influence the work reported in this paper.

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