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Construction of a hierarchical CoP@ZnIn₂S₄ heterojunction for photocatalytic hydrogen evolution†

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To relieve the present energy scarcity and environmental challenges, hydrogen formation *via* water decomposition utilizing sunlight is desirable. However, due to poor charge separation and low hydrogen production efficiency, conventional semiconductor photocatalysts are limited in practical applications. Herein, by introducing a spatial decoupling strategy, a core-shell CoP@ZnIn₂S₄ heterojunction photocatalyst is built through *in situ* formation of two-dimensional ZnIn₂S₄ (ZIS) nanosheets on a spiky CoP surface, achieving effective separation of the redox sites, resulting in a substantial enhancement of photocatalytic hydrogen formation efficiency. Notably, the optimized 5-CoP@ZIS photocatalyst exhibits a hydrogen production rate of 14.98 mmol g⁻¹ h⁻¹ with full solar spectrum illumination, which is 6.3-fold higher than the individual ZIS. These results exhibit that the increased efficiency is explained by the creation of the CoP@ZIS core-shell heterostructure, which strongly prevents the self-assembly and aggregation of the ZIS nanosheets, and offers enhanced light absorption and an abundance of reaction sites. The close interfacial contact facilitates the light-induced electron separation and transfer from ZIS to CoP, which synergistically yields an efficient hydrogen production performance. This work offers a highly effective method for the fabrication of powerful photocatalysts to realize light energy conversion.

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1. Introduction

As fossil energy usage rises and environmental issues become progressively more prominent, the pursuit for clean and sustainable alternative energy sources becomes increasingly pressing. Researchers have extensively recognized and predicted hydrogen as a renewable and clean new energy source.^{1–6} Recently, researchers have developed a variety of hydrogen production technologies, including electrolysis of water, hydrogen synthesis from fossil raw materials, biomass hydrogen generation, and photocatalytic decomposition of water. Among them, solar-driven semiconductor-based fabrication of hydrogen through water decomposition is one of the greatest viable hydrogen production strategies due to the universality and inexhaustibility of sunlight. To date, numerous semiconductors with various bandgap energies, including oxides (ZnO, TiO₂ and Fe₂O₃),^{7–9} sulfides (CdS, ZnS

and MoS₂),^{10–12} and carbon-based materials (graphene, g-C₃N₄),^{13,14} *etc.*, have been produced and thoroughly investigated for photocatalytic water decomposition to produce hydrogen. However, on account of their restricted ability to absorb light and slow carrier dynamics, conventional single-component semiconductor photocatalysts typically exhibit inferior photocatalytic activity.¹⁵ For example, ZnO, as a significant n-type semiconductor, has made many advances in photocatalytic hydrogen production due to its non-toxicity, easy-to-control morphology, and high redox potential. However, on account of its wide bandgap (~3.37 eV), the corresponding PHE process can only be carried out in the UV irradiation region, and most of the sunlight cannot be utilized, which severely limit the improvement of its photocatalytic hydrogen production efficiency.¹⁶ As a result, it is critical to design and build efficient and robust photocatalysts.

Metal sulfides, particularly ZnIn₂S₄ (ZIS), are recognized to be potential photocatalysts with their minimal toxicity, appropriate bandgap width, and distinctive electro-optical characteristics, which have lately sparked renewed interest in photocatalytic generation of hydrogen.^{17–19} However, due to the rapid complexation speed of photogenerated carriers and the scarcity of active surfaces, it has a low light utilization.²⁰ Furthermore, due to the accumulation of photogenerated cavities in ZIS-based catalysts, S²⁻ ions may combine with the

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cavities and generate S, leading to serious photocorrosion processes, thereby shortening the durability of the photocatalysts and limiting their practical applications.²¹ Impressively, the unique ultrathin 2D ZIS nanosheets not only have an extensive specific surface area for exposing numerous active regions, but also a significantly reduced photogenerated carrier migration distance.²² However, individual 2D ZIS nanosheets are subject to self-assembly and aggregation, causing the development of 3D ZIS nanoflowers, which greatly restrict the ability of photogenerated carriers to separate, as well as minimize their photocatalytic capacity.²³ Therefore, it is critical to develop a strategy that can successfully prevent the aggregation of 2D ZIS nanosheets while also improving the photocatalytic activity.

The loading of co-catalysts and the building of heterojunctions are serviceable solutions for solving the charge separation challenge. By coupling suitable co-catalysts with ZIS nanosheets, electrons and holes move to distinct surface sites on the catalysts, enabling an excellent separation of both oxidizing and reducing sites. Meanwhile, a high conductivity co-catalyst may efficiently avoid the accumulation of photogenerated carriers, reduces the hydrogen production overpotential, and integrates the advantages of each component into the complex, as well as performs special functions that a single photocatalyst cannot.²⁴ In recent years, hierarchical core-shell heterostructures with ZIS nanosheets as the shell have been widely studied, with tight interfacial contact, shortened charge transfer distances, and huge specific surface areas exposing abundant active sites, all of which are conducive to photocatalytic efficiency enhancement. Moreover, the self-assembly and aggregation phenomena of 2D ZIS nanosheets can be mitigated to a degree. For example, Zhou *et al.* have designed the core-shell type $\text{Cu}_{2-x}\text{S}@Zn\text{In}_2\text{S}_4$ heterojunction, in which the empty core-shell structure provides an increased specific surface area with numerous reaction sites, the photocatalytic response range is wider, and the photothermal impact is prominent, which promotes the photocatalytic reactions.²⁵ Lou *et al.* have built a hierarchical $\text{Co}_9\text{S}_8@Zn\text{In}_2\text{S}_4$ heterogeneous cage as an effective visible light hydrogen evolution photocatalyst. This unique structure effectively promotes photogenic charge separation and migration, offers a huge specific surface area, leads to a wealth of active sites for photocatalytic reactions, and displays superior photocatalytic water decomposition capability and durability.²⁶

In light of their strong electronic conductivity, superior chemical stability, and comparatively low hydrogen production overpotential, transition metal phosphides have aroused the interest of scientists as co-catalysts.²⁴ Cobalt phosphide (CoP), a particular kind of transition metal phosphide, has attracted plenty of interest with respect to its abundant raw materials and straightforward fabrication procedure. However, the majority of these investigations used CoP as an isolated active center anchored onto the outer layer of the catalyst; this approach falls short of the requirement

for high spatial separation in photocatalytic systems and lessens the efficiency of interfacial electron transfer in the reaction.^{27–30} We plausibly believe that the architecture of heterojunctions affects photocatalytic reactions significantly, and co-catalysts with significant specific surface areas are desirable for fast electron transfer in the reaction. Thus, we recommend using CoP as a precursor with lamellar ZIS to boost photocatalytic H_2 generation activity.

Herein, we created a core-shell hierarchical structure of the CoP@ZIS heterojunction photocatalyst through *in situ* growth. As expected, the tight interfacial contact between CoP and ZIS facilitates photogenerated electron migration from ZIS to the CoP surface, effectively shortening the transfer distance and optimizing electron-hole pair separation and migration efficiency. Notably, the optimized 5-CoP@ZIS sample exhibits a remarkable hydrogen yield of $14.98 \text{ mmol g}^{-1} \text{ h}^{-1}$ under full solar spectrum treatment, which is 6.3 times greater than pure ZIS. Moreover, after 4 times of cycling, 5-CoP@ZIS maintained a high hydrogen production rate, demonstrating good stability. This work presents an appealing idea for fabricating extremely efficient ZIS-based photocatalysts for H_2 evolution *via* water decomposition.

2. Results and discussion

The synthetic route of CoP@ZIS spiny nanoflowers is schematically represented in Fig. 1. To begin with, a simple hydrothermal technique was used to create the precursor of $\text{Co}(\text{CO}_3)_{0.5}(\text{OH})_{0.11}\text{H}_2\text{O}$ spiny nanoflowers. The $\text{Co}(\text{CO}_3)_{0.5}(\text{OH})_{0.11}\text{H}_2\text{O}$ precursor was thermally oxidized to porous Co_3O_4 , which was subsequently phosphated to create porous CoP spinous nanoflowers. Finally, the CoP@ZIS composite was constructed through *in situ* formation of ultrathin ZIS nanosheets on the outer layer of CoP employing a low temperature solvent heat treatment, which used ZnCl_2 , $\text{InCl}_3 \cdot 4\text{H}_2\text{O}$ and thioacetamide (TAA) as raw materials.

Fig. S1a† shows the scanning electron microscopy (SEM) picture of the $\text{Co}(\text{CO}_3)_{0.5}(\text{OH})_{0.11}\text{H}_2\text{O}$ precursor. Obviously, the $\text{Co}(\text{CO}_3)_{0.5}(\text{OH})_{0.11}\text{H}_2\text{O}$ precursor is assembled from smooth nanothorns into nanoflowers with a majority size ranging from 8–14 μm , while adjacent nanothorns possess adequate spaces, which is conducive to coating of the target product ZIS.^{21,31} After thermal oxidation, the obtained Co_3O_4 retains the basic shape of the original $\text{Co}(\text{CO}_3)_{0.5}(\text{OH})_{0.11}\text{H}_2\text{O}$ substrate with some rough surface (Fig. 2a–c). The (311) crystal plane of Co_3O_4 has a lattice spacing of 0.243 nm, as seen in Fig. 2d. After further phosphating, the surface of CoP becomes rougher (Fig. 2e, f and S3a†). Obviously, the presence of numerous cavities in CoP nanothorns (Fig. 2g, S3b and c†) is undoubtedly advantageous for the quick separation and transmission of photogenerated electron-hole pairs, and the existence of cavities may be attributed to the water vapor adsorbed by the CoP and the carbon source escaping in the form of gas during the high-temperature calcination process. A high-resolution transmission electron microscopy (HRTEM) picture (Fig. 2h) of



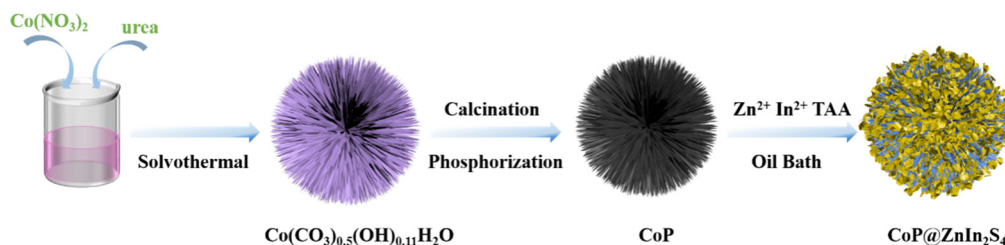


Fig. 1 Schematic diagram for the synthesis of core-shell CoP@ZIS heterostructure.

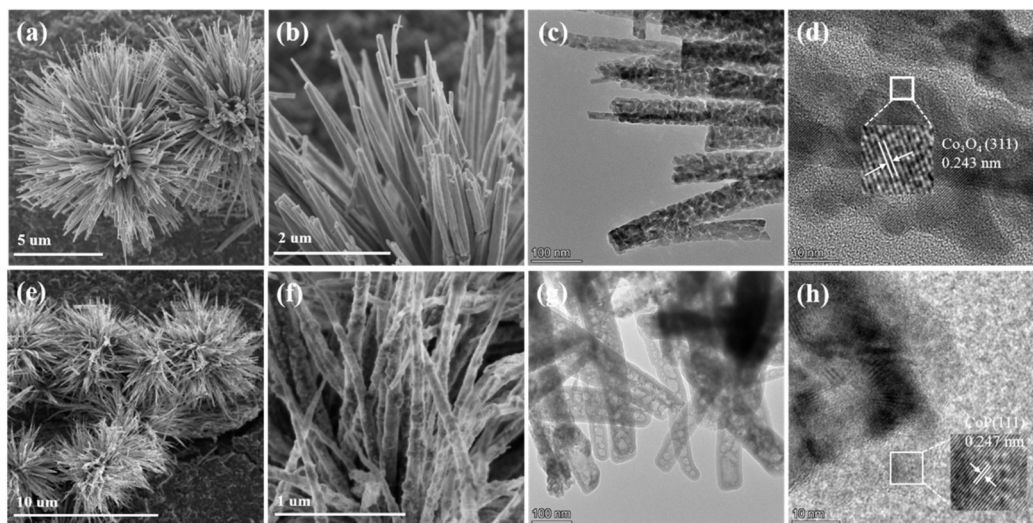


Fig. 2 SEM images of (a and b) Co_3O_4 and (e and f) CoP. TEM images of (c) Co_3O_4 and (g) CoP. HRTEM images of (d) Co_3O_4 and (h) CoP.

CoP reveals a distinct lattice spacing of 0.247 nm, belonging to the CoP (111) plane. According to the EDX elemental analysis (Fig. S3d-f†), Co and P are evenly distributed in CoP spiny flower spheres. When ZIS is coupled with CoP, it is obvious that each CoP nanothorn is evenly wrapped in ZIS nanosheets and exhibits a close contact interface (Fig. 3a and b), which is further revealed by TEM results (Fig. 3c and d). The synthesized ZIS possesses intersecting ultrathin nanosheets which are distributed uniformly on the CoP surface to construct a core-shell framework, effectively preventing the aggregation of individual ZIS, which offers improved multiple light reflections and thus optimizes the utilization of sunlight.^{21,32} Importantly, the HRTEM image (Fig. 3e) collected at the contact of the CoP nanothorn and the ZIS nanosheet reveals the distinct crystal fringes of 5-CoP@ZIS with d values of 0.317 nm and 0.247 nm that may be nicely attributed to the (102) and (111) lattice planes of ZIS and CoP, respectively. These results imply a close contact between CoP and ZIS. Additionally, the corresponding elemental mapping images (Fig. 3f-k) confirm the coexistence and even distribution of Co, P, S, In and Zn elements, with Co and P elements being mainly distributed in the middle core region and S, In and Zn elements being distributed throughout the entire CoP, further confirming the successful synthesis of core-shell structure CoP@ZIS. The photocatalyst's precise structural design prohibits self-assembly and aggregation of 2D ultra-thin ZIS nanosheets, ensuring structural stability as well

as rapid separation and migration of photogenerated electrons, hence boosting photocatalytic activity.²³ Instead, in the absence of CoP, ZIS formed nanosheet aggregates (Fig. S4a-d†).

We employed X-ray diffraction (XRD) to examine the structure of the crystals and composition of the phases of the as-obtained products. Fig. S1b† shows the prepared $\text{Co(CO}_3)_{0.5}(\text{OH})_{0.11}\text{H}_2\text{O}$ (JCPDF# 48-0083). Fig. S2† reveals that Co_3O_4 (JCPDS# 42-1467) was successfully produced. The peaks located at 31.597° , 36.321° , 46.238° , 48.118° and 56.796° in Fig. 3l can be ascribed to the (011), (111), (112), (211) and (301) planes of orthorhombic CoP (JCPDS# 65-1474), correspondingly,³³ demonstrating the successful production of CoP after phosphating the Co_3O_4 . Furthermore, all diffraction peaks observed for pure ZIS match well with the characteristic peaks of ZIS (JCPDS# 65-2023). Notably, all the CoP@ZIS composites mainly display ZIS characteristic diffraction peaks, whereas the peak intensity of CoP is modest, which may be owing to the low concentration of CoP and the fact that it is wrapped in ZIS nanosheets.³⁴ The degree of intensity for the (011) diffraction peak gradually increases as CoP concentration increases, while the strength of the ZIS diffraction peaks declines. It is also notable that the diffraction peak of (011) belonging to CoP in CoP@ZIS migrates to a higher angle. The modest change in peak locations and decline in peak intensity in the heterojunctions reveal that there is a certain interaction between CoP and



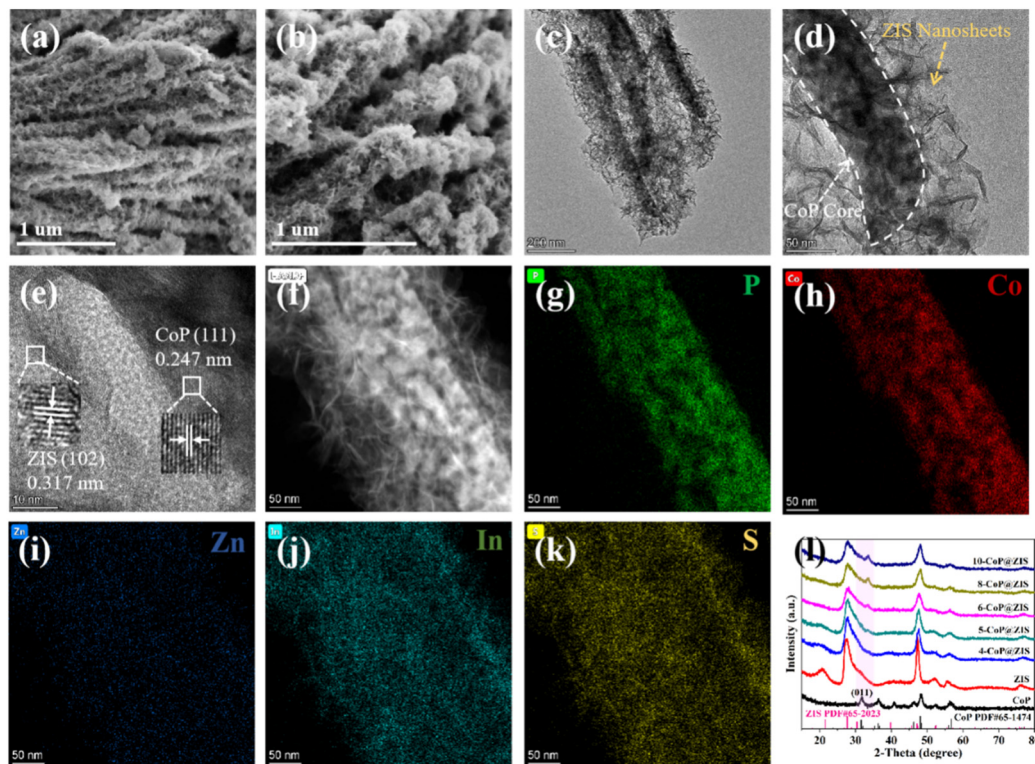


Fig. 3 (a and b) SEM, (c and d) TEM, (e) HRTEM, (f) HAADF-STEM, and (g–k) corresponding elemental mapping images of 5-CoP@ZIS. (l) XRD patterns of the different samples.

ZIS.³⁵ This conclusion is compatible with SEM and TEM findings, suggesting that CoP and ZIS are successfully recombined.

X-ray photoelectron spectroscopy (XPS) was employed to further explore the detailed chemical make-up of the composite surface as well as the relevant valence-state of elements. The survey XPS result of 5-CoP@ZIS clearly demonstrates the peaks of Co 2p, P 2p, S 2p, In 3d and Zn 2p (Fig. S5†). This is parallel to the finding of the elemental mapping discussed before. In the high-resolution XPS result of Zn 2p (Fig. 4a), the peaks for Zn 2p_{3/2} and Zn 2p_{1/2} in ZIS are located at 1022.17 eV and 1045.17 eV, respectively,³⁶ whereas those for the 5-CoP@ZIS sample are shifted to 1022.44 eV and 1045.45 eV, correspondingly. In the In 3d XPS spectra of Fig. 4b, the In 3d_{5/2} and In 3d_{3/2} peaks for ZIS are centered at 445.29 eV and 452.82 eV, respectively.³⁷ Similarly, the peaks centered at 445.42 eV and 452.95 eV in the In 3d XPS spectra of 5-CoP@ZIS are attributed to In 3d_{5/2} and In 3d_{3/2}, correspondingly. The S 2P XPS result for pure ZIS in Fig. 4c includes two peaks at 161.88 eV and 163.08 eV, which can be ascribed to S 2p_{3/2} and S 2p_{1/2}, correspondingly,³⁸ whereas those for the 5-CoP@ZIS sample are shifted to 161.97 eV and 163.15 eV, correspondingly. It is noteworthy that, when CoP is coupled with ZIS, the peak strength of Co2p and P 2p in 5-CoP@ZIS is low due to the instrument's detection accuracy, as well as the fact that CoP is located in the core of 5-CoP@ZIS and the depth of ZIS is massive, but the main Co2p_{3/2} and Co2p_{1/2} species remain readily

identified. The CoP sample's Co 2p XPS spectra can be analyzed into six peaks (Fig. S6†). The values observed at 778.83 eV and 793.74 eV commonly belong to Co 2p_{3/2} and Co 2p_{1/2} for CoP, correspondingly, and those at 782.19 eV and 797.98 eV corresponded to Co 2p_{3/2} and Co 2p_{1/2} for oxidized Co species induced by the inevitable surface oxidation of CoP. Besides, two satellite peaks at 785.31 eV and 802.65 eV are triggered by high-spin Co²⁺ ion oscillations.³⁹ For the 5-CoP@ZIS composite, the Co 2p peaks are shifted toward the low binding energy. In Fig. 4d, the P 2p XPS result of bare CoP shows three separated peaks, with the peak at 134.34 eV being linked with oxidized P species, and the two peaks at 129.65 eV and 130.75 eV belonging to P 2p_{3/2} and P 2p_{1/2}, respectively, relating to the phosphorus anion in CoP.⁴⁰ The P 2p peaks for the 5-CoP@ZIS hybrid, like the Co element, exhibit a negative change as compared to CoP. As previously described, the Zn 2p, In 3d and S 2p peaks in 5-CoP@ZIS all move slightly towards higher binding energy compared with the corresponding peaks of the original CoP and ZIS, while the peaks of Co 2p and P 2p move in the opposite way, indicating that when CoP and ZIS are coupled, the inner chemical environment of these parts changes and charges migrate in their contact section.⁴¹ In general, increased binding energy matches decreased electron density.⁴² As a result, when CoP and ZIS are coupled, ZIS serves as an electron donor and CoP serves as an electron acceptor, and electrons move from ZIS to CoP, accelerating charge carrier separation and migration, thus increasing the



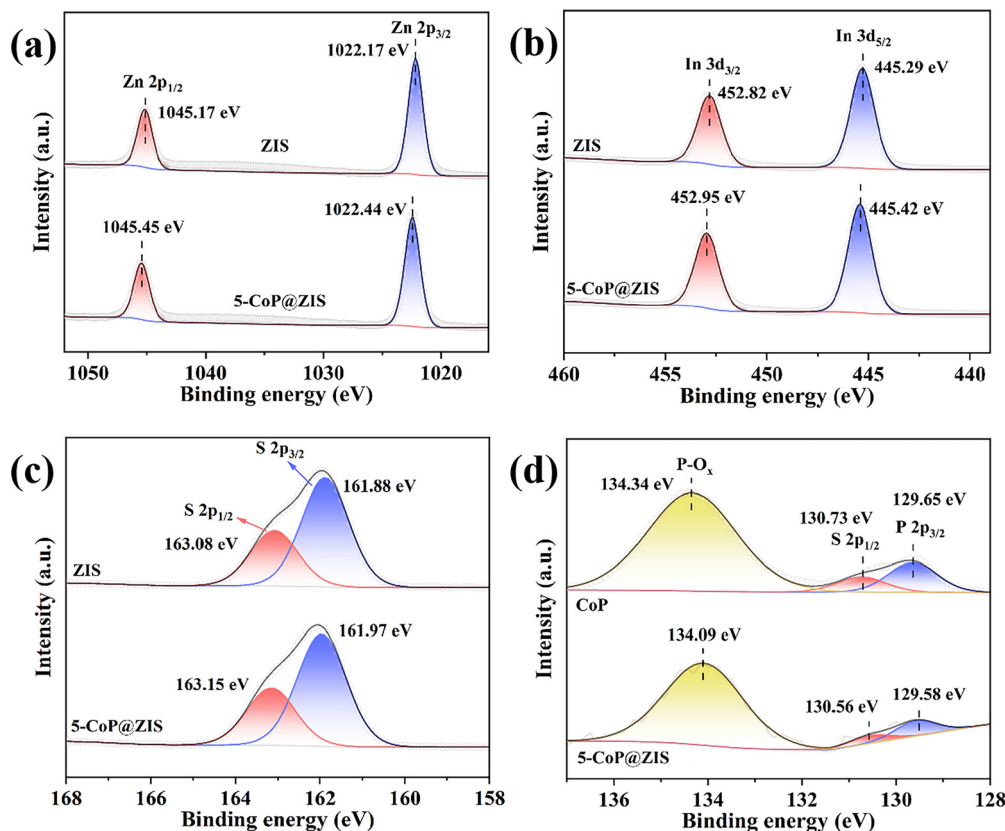


Fig. 4 High-resolution XPS spectra of (a) Zn 2p, (b) In 3d, (c) S 2p and (d) P 2p of samples.

efficiency of 5-CoP@ZIS photocatalytic decomposition of water. All of the aforementioned peak changes can be explained by the creation of the p-n heterojunction between CoP and ZIS components.

To measure the photocatalytic H₂ production capacity of the synthesized products, photocatalytic water decomposition for H₂ evolution was performed using triethanolamine (TEOA, 10 vol%) as a sacrificial agent. The time-dependent graph of the hydrogen production rate for various photocatalysts is depicted in Fig. 5a. The rate of H₂ generation increases progressively as the time of light irradiation increases. The average production rate of H₂ is shown in Fig. 5b. Due to the self-assembled 2D ZIS nanosheet structure and slow carrier migration, the average H₂ formation rate of bulk ZIS is quite low (only 2.36 mmol g⁻¹ h⁻¹). And CoP produces minimal H₂ gas, indicating that it merely functions as a cocatalyst. On the contrary, CoP@ZIS possesses a substantially higher photocatalytic H₂ creation capability than either CoP or ZIS alone. Besides, the H₂ production rate increases with the content of CoP in CoP@ZIS, then decreases. This may be due to excess CoP serving as a site for charge carriers to recombine,²¹ reducing the photocatalytic capacity. Therefore, the amount of CoP in the heterostructure has a significant impact on photocatalytic performance, and only a moderate amount CoP can demonstrate optimum photocatalytic hydrogen evolution performance. When the CoP content

increases to 5 mg, the H₂ generation rate at 5-CoP@ZIS reaches the maximum (14.98 mmol g⁻¹ h⁻¹), which is nearly 6.3-fold higher compared to the original ZIS. The synthesized CoP@ZIS heterojunctions combine numerous features that are favorable for photocatalytic hydrogen evolution reactions. First, the ZIS nanosheet shell is tightly bound to CoP and is evenly distributed on the CoP surface by ultra-thin nanosheets, which can prevent 2D ZIS nanosheet self-assembly and aggregation, enlarge the sample's active surface area, and supply greater active sites for the creation of hydrogen.⁴³ Second, the CoP@ZIS complex's light absorption capacity will be greatly boosted. Third, the CoP@ZIS heterojunctions will considerably increase charge transfer and migration, so that more electrons take part in the hydrogen evolution reaction, hence increasing photocatalytic activity.

Meanwhile, several kinds of controlled investigations were performed to investigate the impact of various sacrificial reagents on the performance of the 5-CoP@ZIS catalyst. The rates of generation of hydrogen are 14.98 mmol g⁻¹ h⁻¹, 5.47 mmol g⁻¹ h⁻¹ and 1.59 mmol g⁻¹ h⁻¹ when the sacrificial agents are TEOA (10 vol%), lactic acid (10 vol%) and 0.25 M Na₂SO₃-0.35 M Na₂S, respectively, as displayed in Fig. 5d. Without the introduction of sacrificial agents, the yield of photocatalytic hydrogen generation is minimal. The findings indicate that the sacrificial agent is critical to the photocatalytic reaction system, and TEOA can be used as an



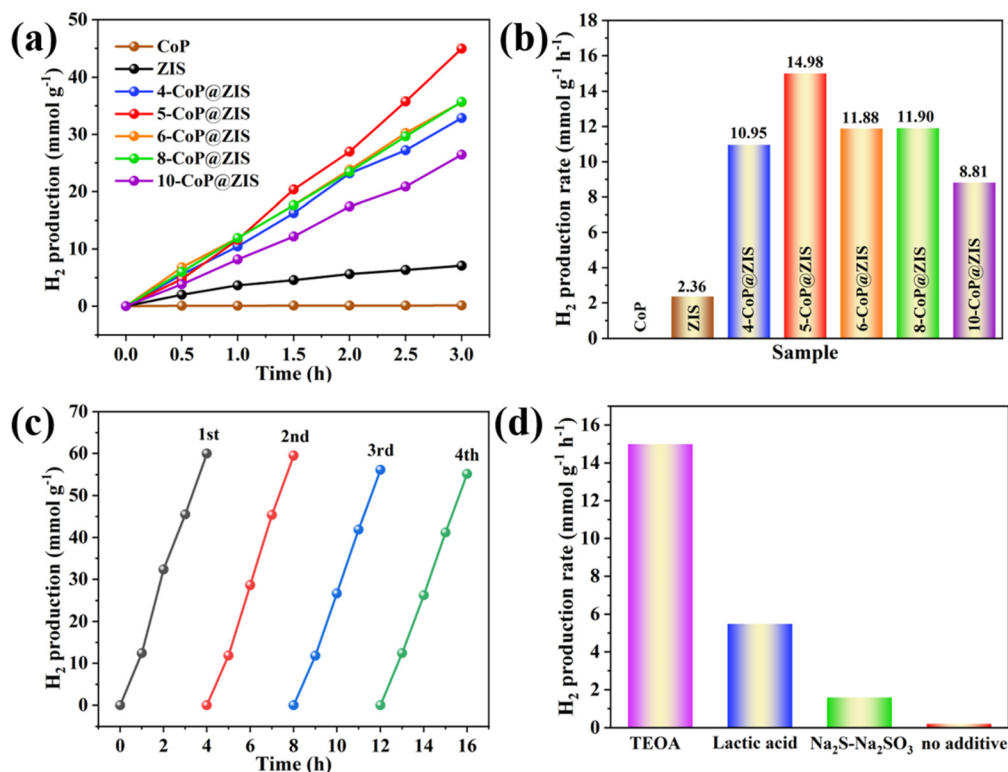


Fig. 5 Photocatalytic H₂ evolution performance: (a) time-yield plots of H₂ evolution. (b) H₂ evolution rate for different samples. (c) Cyclic tests of 5-CoP@ZIS. (d) Photocatalytic hydrogen evolution performance with different sacrificial reagents on 5-CoP@ZIS.

appropriate sacrificial agent in the present reaction system for effective hydrogen generation.

Besides the powerful photocatalytic H₂ production activity, recycling studies were done to test the stability of this 5-CoP@ZIS catalyst. As shown in Fig. 5c, the results reveal no substantial decline in hydrogen formation after four continuous rounds. Furthermore, the XRD patterns of 5-CoP@ZIS after the reaction show no discernible alteration from the original data, and the SEM images reveal that the morphology of 5-CoP@ZIS remains unchanged considerably after the activity test (Fig. S7[†]). The findings further confirm the satisfactory stability of the prepared 5-CoP@ZIS catalyst.

To deeply investigate the causes of the increase in photocatalytic activities, photophysical methods were utilized to evaluate the migration behaviors of charge carriers. Steady-state photoluminescence (PL) studies reveal the

recombination efficiency of free excitons, and the stronger PL spectrum strength represents the faster recombination process of photogenerated electrons and holes.¹⁸ As represented in Fig. 6a, individual ZIS shows a significant PL peak, indicating an extensive degree of charge carrier combination. The PL intensity of 5-CoP@ZIS drops dramatically compared with ZIS, manifesting a rapid separation of photogenerated electrons and holes, that's in good agreement with the photocatalytic activity data. In addition, the transient photocurrent response (Fig. 6b) and electrochemical impedance spectra (Fig. 6c) were recorded to explore the photo-induced electron/hole separation and transfer efficiency. Fig. 6b displays that 5-CoP@ZIS possesses an increased photocurrent density compared with pure ZIS, demonstrating that 5-CoP@ZIS has stronger charge carrier separation and migration dynamics.⁴⁴ The Nyquist plots

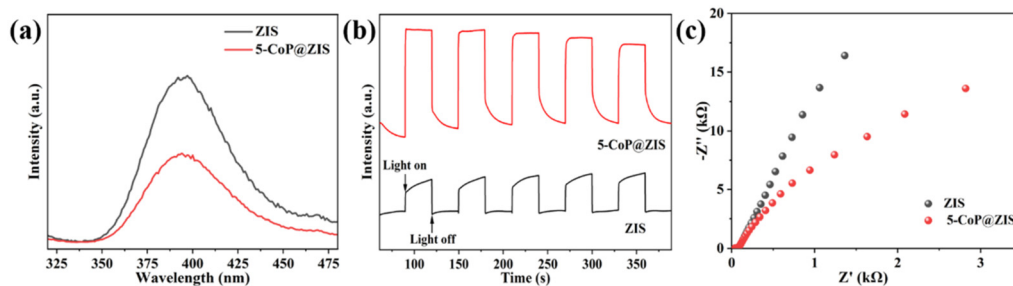


Fig. 6 (a) Steady-state PL spectra. (b) Transient photocurrent response curves. (c) EIS plots of ZIS and 5-CoP@ZIS.



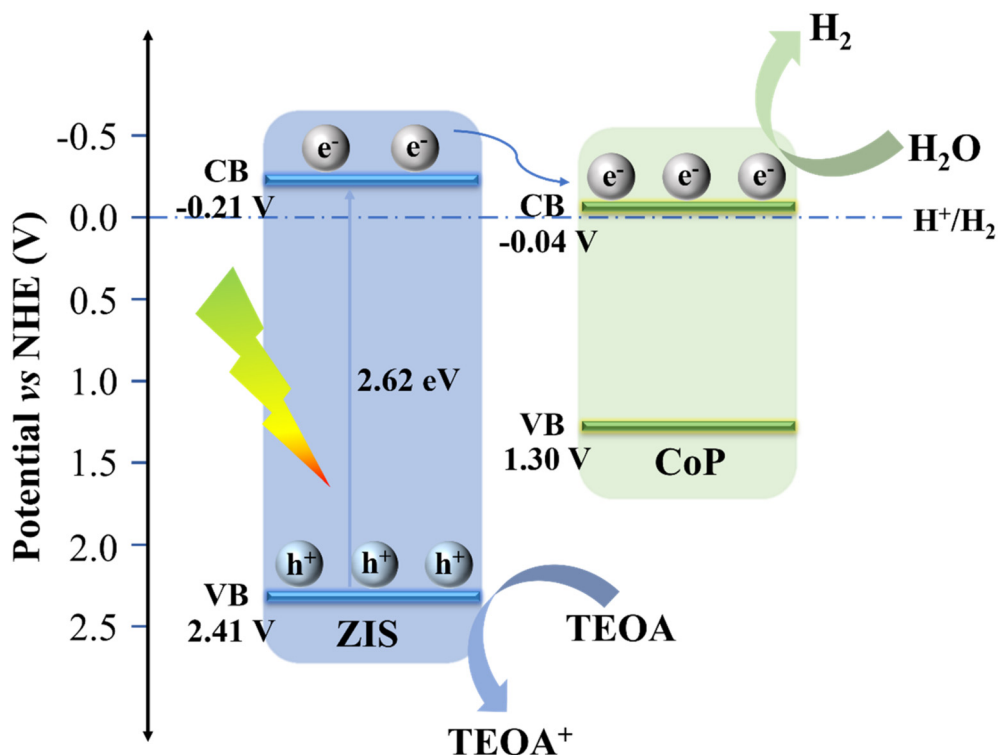


Fig. 8 Schematic diagram of the possible charge flow paths.

quickly move to the CB in CoP, the directional and efficient electron migration from the shell of ZIS to the nucleus of CoP takes place in order to reduce the protons and release the hydrogen, and the holes left in the VB in ZIS are trapped by the sacrificial agent, leading to a long-distance separation of oxidative and reductive sites and increase of photocatalytic performance.⁵⁵ Because of its metal-like nature and narrow band gap, CoP contributes little to the formation of reducing electrons under photoexcitation, and its beneficial function is to increase electron conduction on the surface so as to finish the protonation reaction.⁴² Furthermore, the CoP@ZIS core-shell structure possesses a higher diffusion surface area, which ensures that the photogenerated electrons may transport efficiently at the interface and considerably improves the separation efficiency of electron-hole pairs, thus enhancing PHE capability.

3. Conclusions

In light of the limited charge separation of photocatalysts, efficient H₂ production remains a long-term difficulty. This difficulty is well addressed in this work by the *in situ* fabrication of ZIS nanosheets on spiny CoP nanoflowers using a spatial decoupling technique. The created core-shell CoP@ZIS heterostructure effectively limits ZIS nanosheet aggregation, exposing more active sites spatially separating oxidation and reduction sites. Under complete solar spectrum irradiation, the optimized 5-CoP@ZIS sample acquires an excellent hydrogen yield of 14.98 mmol g⁻¹ h⁻¹,

which is nearly 6.3-fold greater compared to the single ZIS. This study illustrates the feasibility of building CoP@ZIS heterojunction photocatalysts to facilitate charge separation for fast H₂ production so as to solve the increasing energy scarcity and environmental concerns.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

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