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Synergetic microstructure engineering by induced ZB/WZ twin boundaries and S vacancies in a $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ -based S-scheme photocatalyst for highly efficient photocatalytic hydrogen production†

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Surface-abundant active sites, rapid charge transport and associated prolonged electron lifetime are vital factors that determine efficient photocatalysis. A series of different $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solutions, including single crystalline $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ (ZCS), single crystalline $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ with S vacancies (ZCS-V), twin structured- $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ (T-ZCS) and twin-structure $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ with S vacancies (T-ZCSv) were successfully prepared in the present work by manipulating the conditions of the hydrothermal reaction. Experimental results confirm that the optimized T-ZCSv photocatalyst that possesses a hexagonal wurtzite/zinc blende (WZ/ZB) twin structure and rich-surface S vacancies exhibits an excellent photocatalytic hydrogen production efficiency of approximately $551.74 \mu\text{mol h}^{-1}$. The outstanding performance of the optimized T-ZCSv is attributed to the prolonged electron lifetime and effectively facilitated separation and migration of charge carriers. These are provided by the periodically aligned WZ/ZB interfacial homojunctions that form the S-scheme staggered energy band structure across the junction and abundant S vacancies that serve as electron trapping sites in the T-ZCSv. Furthermore, T-ZCSv are uniformly dispersed on 2-methylimidazole zinc salt [zeolitic imidazolate framework-8 (ZIF-8 polyhedron)], which not only could inhibit the aggregation of T-ZCSv but also expose more active sites for photocatalytic-redox reactions. Finally, a possible charge separation and transfer mechanism explaining the optimum activity of the outperforming sample is proposed on the basis of the results obtained from a range of investigation methods [scanning electron microscopy and energy-dispersive spectroscopy (SEM-EDS), transmission electron microscopy and high-resolution transmission electron microscopy (TEM/HRTEM), X-ray diffraction (XRD) technique, ultraviolet-visible (UV-vis) diffuse reflection spectroscopy, and electron paramagnetic resonance (EPR) spectroscopy]. This study demonstrates the development of a structurally unique $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ (with twin structure and S vacancies) and a $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ -based metal-organic framework (MOF) for photocatalytic applications.

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Introduction

Recently, metal sulfide semiconductors, especially ternary zinc cadmium sulfide ($\text{Zn}_x\text{Cd}_{1-x}\text{S}$) solid solutions, have been widely used in photocatalytic hydrogen production due to their excellent photocatalytic performance with an appropriate energy band structure and efficient photo-charge separation.^{1–4} Compared to pristine zinc sulfide (ZnS) and cadmium sulfide (CdS), ZnS has a broader band gap (~ 3.6 eV), across which the electrons are only excited by ultraviolet (UV) light, thus limiting the utilization of the full sun spectrum. Some fatal drawbacks also inhibit the efficiency of CdS for hydrogen production, including low resistance to photo-corrosion and rapid

recombination of photogenerated charge carriers.^{5–7} Therefore, a series of ZnCdS solid solutions of tunable band gaps and band edges are prepared by changing the ratio of Zn/Cd, which can effectively solve the above problems and improve photocatalytic activity and stability.^{8–10} It is worth noting that some typical research work of single crystal ZnCdS solid solution with hexagonal wurtzite (WZ) and cubic zinc blende (ZB) phases was respectively prepared by Wang and Chan *et al.*^{11,12} As they hypothesized, single-crystalline ZnCdS exhibited better hydrogen production efficiency than individual CdS and ZnS. These earlier works addressed the basic challenges of sulfide photocatalysts to some extent, such as severe photocorrosion, poor stability and visible light absorbance, and a non-adjustable band gap. Nevertheless, the single-crystal phase (defect-free crystal) of $\text{Zn}_x\text{Cd}_{1-x}\text{S}$ still suffers from low separation efficiency of space charge and a short lifetime owing to its untargeted migration and short diffusion lengths.

A twin crystal structure, as one of the crystal phase engineering types, containing two different crystalline phases can be formed in single-twinned nanocrystals between the wurtzite (WZ) and zinc blende (ZB) segments.¹³ Obtaining twin structures in one-dimensional nanocrystals provides an alternative alignment of long-range-order twinned planes in a twin-superlattice arrangement at the nanoscale and introduces ingenious WZ/ZB structural segments at particular dimensions.^{14,15} The parallel WZ/ZB twin boundaries form a one-dimensional ordered homojunction because of the realignment of the band structures across the WZ/ZB coherent twin boundary interfaces. Compared with the traditional charge transfer mechanism, Yu *et al.* first proposed the concept of Step-scheme (S-scheme) photocatalytic principle which solves the contradiction of the photocatalytic charge transfer path reported previously.¹⁶ They first designed WO_3 (2D) and $\text{g-C}_3\text{N}_4$ (2D) S-scheme heterojunction photocatalysts in 2019, which showed high photocatalytic H_2 -production performance, which was approximately 1.8 times that of pure $\text{g-C}_3\text{N}_4$ due to the efficient charge separation and transfer in the photocatalytic reaction process.¹⁷ The suitable staggered band configurations of the S scheme photocatalytic system possess optimized redox capability, which could effectively accelerate the recombination of relatively useless photo-generated charges but inhibit the recombination of useful holes and electrons, and facilitate better charge transport properties owing to its band-to-band potentials and highly ordered structural charge transport pathways, respectively.^{18,19} Consequently, the interfacial resistance of photo-generated charges for coherent twin boundaries is one order of magnitude lower than that of the conductors and semiconductors with crystal structures of large-angle grain boundaries.^{20,21} Therefore, the structurally induced nano twin crystals enhance the photocatalytic activity. An imitative recreation of this structural feature is realized by achieving a periodic alignment of wurtzite/zinc blende (WZ/ZB) structures together with regular spacing in the $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ -pseudobinary structure.²² The WZ/ZB homojunction is a conventional type II staggered band in the twinned superlattice structure of the ZnCdS solid solution. It still preserves the

transfer properties of charge carriers such as perfect nanocrystals.²³ The highly ordered homojunction structure and its interfacial electrostatic field lead to an effective separation of photo-charges and transfer to the surface of the samples. Consequently, they participate in redox reactions. Thus, the S scheme system inhibits charge recombination and optimizes the utilization rate of photogenerated charge carriers.¹⁴ Du *et al.* prepared a series of twinned $\text{Zn}_x\text{Cd}_{1-x}\text{S}$ nanomaterials employing $\text{Cd}^{2+}/\text{Zn}^{2+}$ acetate and thioacetamide as precursors.²⁴ The results indicated that an appropriate amount of tri-ethanolamine quantity resulted in the $\text{Zn}_x\text{Cd}_{1-x}\text{S}$ solid solution with a high-density WZ/ZB twinned superlattice structure, which greatly improved the separation of charge carriers and photocatalytic performance.

Furthermore, the development of novel functional photocatalysts with optimum optoelectronic characteristics has been achieved by exploiting the unique advantages of vacancy engineering for sustainable energy and environmental applications. It is also possible to tune the photocatalytic ability of semiconductors by manipulating the distribution and concentration of defects. Vacancy engineering pertaining to the introduction of typical sulphur (S) vacancies into the crystal structure of semiconductor photocatalysts has been shown to lead to higher absorption of visible light, prolonged electron lifetimes and abundant reactive sites.^{25–27} However, the synergy between the twin crystal structure and vacancies to improve the photocatalysis has not been reported yet. Therefore, the acquisition of WZ/ZB twin structure-based homogeneous junctions and the introduction of S vacancies should be a practical way to boost the activity of hydrogen production of ZnCdS-based photocatalysts. Moreover, zeolitic imidazolate frameworks (ZIFs), a sub-class of MOFs, are promising in the synthesis of novel functional materials by serving as excellent precursors and/or templates in compounds/composites such as metal sulfides, metal oxides, interesting porous carbon materials, and carbon-metal/metal oxide hybrids with systematically controlled compositions.^{28,29} Besides, the composites based on ZIF/ZIF derivatives act as potential light-harvesting sites and electronic mediators. Loading ZnCdS onto the surface of ZIF-8 using direct *in situ* growth may be an effective strategy to achieve the dispersion of the catalyst and improve its catalytic activity.

Herein, we have prepared a series of $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solutions, including single-crystal $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ without S vacancies and twin structure (ZCS), S-vacancy-rich single-crystal $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ (ZCS-V), twinned $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ without S vacancies (T-ZCS) and S-vacancy-rich twin-crystal $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ (T-ZCSv) using a facile and feasible method. The photocatalytic activity of the samples was evaluated by hydrogen production, and T-ZCSv is the best member of this series with the highest photocatalytic hydrogen production activity, reaching approximately 18 times that of the bare ZCS. A comparative study is carried out among various ZIF-8@T-ZCSv composites containing different amounts of uniformly distributed T-ZCSv on the surface of ZIF-8 polyhedra. The uniform distribution of T-ZCSv prevents the agglomeration of T-ZCSv, thereby facilitating the exposure of more active sites, which contribute to enhancing

the photocatalytic hydrogen production rates of ZIF-8@T-ZCSv composites. Optimized ZIF-8@T-ZCSv (1300 mg) exhibited a superior H_2 generation rate of $\sim 583.25 \mu\text{mol h}^{-1}$. Furthermore, the characterization techniques employed could verify the formation mechanism of WZ/ZB twin boundaries and S vacancies in $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ -based S scheme photocatalysts and reveal the reasons for promoting the photocatalytic hydrogen production efficiency. This work mainly emphasizes the critical role of the S vacancy-defects and the charge transfer mechanism across twin-crystals in improving the photocatalytic ability of the $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solution supported by the ZIF-8 substrate, which develops a new idea for designing efficient photocatalysts to achieve sustainable hydrogen production.

Experimental

Synthesis of the $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solution with different structural characteristics

Preparation of pristine $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ (ZCS) and $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ with sulfur vacancies (ZCS-V). The $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solution with sulfur vacancies was synthesized in a hydrothermal reaction. Initially, $\text{Cd}(\text{CH}_3\text{COO})_2 \cdot 2\text{H}_2\text{O}$ (5 mmol, 1332.7 mg) and $\text{Zn}(\text{CH}_3\text{COO})_2 \cdot 2\text{H}_2\text{O}$ (1097.5 mg, 5 mmol) were dissolved in de-ionized water (50 mL) containing appropriate amounts of $\text{N}_2\text{H}_4 \cdot \text{H}_2\text{O}$, and the solution was stirred until it became a clear and transparent solution. Then a required quantity of thiourea (12 mmol, 913.5 mg) was added to the above solution and stirred thoroughly for one hour at ambient temperature, and a 100 mL volume Teflon lined stainless steel autoclave was filled with this solution where the hydrothermal process was carried out at 220°C overnight. Once it was cooled down to ambient temperature naturally, the solution was centrifuged to separate solid particulate products by forming a supernatant. The solid product was cleaned with ethanol and deionized water alternately. The obtained product was finally vacuum dried at 70°C for 5 h. The resulting $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solutions are supposed to possess sulfur vacancies and were designated as ZCS-V. In contrast, pure $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ was produced under the same conditions without adding $\text{N}_2\text{H}_4 \cdot \text{H}_2\text{O}$ in the initial mixture and labeled as ZCS.

Preparation of twin crystal $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ (T-ZCS). A one-step hydrothermal process was performed to obtain the twin-structure $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$. For the twin crystal $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$, initially, cadmium acetate dihydrate (5 mmol, 1333 mg) and $\text{Zn}(\text{CH}_3\text{COO})_2 \cdot 2\text{H}_2\text{O}$ (5 mmol, 917.4 mg) were dissolved in alcohol (60 mL) by stirring for 30 min. Then, thioacetamide (35 mmol, 2630 mg) was dispersed in the solution by magnetic stirring for another 30 min. The suspension was placed in a Teflon autoclave (100 mL), kept at 160°C for 15 h and cooled to ambient temperature. The product was obtained by centrifugation and washed alternately with alcohol and deionized water four times. Finally, the as-cleaned wet powder was heated to 70°C for drying overnight, resulting in a twin-structure $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ dry powder, denoted as T-ZCS.

Preparation of twinned $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ with sulfur vacancies (T-ZCSv). The twinned $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ containing sulfur vacancies

were also synthesized through the hydrothermal reaction at 180°C for 24 h. Using uniform stirring, a certain amount of $\text{Cd}(\text{CH}_3\text{COO})_2 \cdot 2\text{H}_2\text{O}$ (10 mmol, 2665.3 mg), $\text{Zn}(\text{CH}_3\text{COO})_2 \cdot 2\text{H}_2\text{O}$ (10 mmol, 2195.1 mg) and thioacetamide (25 mmol, 1878.3 mg) were dissolved in de-ionized water (40 mL) stepwise with 15 min intervals to form a clear and transparent solution. Then 4 mol L^{-1} NaOH (10 mL) was quickly dropped into the solution, resulting in a visible and violent chemical reaction. A homogeneous suspension was achieved by stirring for 30 min, which was then poured into a Teflon autoclave (100 mL) and treated hydrothermally at 180°C for a day and night. The autoclaves are then cooled to a normal temperature. Product samples from hydrothermal treated solutions were obtained by centrifugation, washing with DI water and alcohol four times alternately, and vacuum drying at 80°C for 5 h. The prepared twin crystal $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ containing sulfur vacancies was labeled as T-ZCSv. The critical preparation details of four different $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solutions are summarized in Table S1.†

Synthesis of ZIF-8 and ZIF-8@T-ZCSv

A typical ageing effect resulted in the formation of ZIF-8@T-ZCSv. First, the $\text{Zn}(\text{NO}_3)_2 \cdot 6\text{H}_2\text{O}$ (734.4 mg) crystals were completely dissolved in methanol (50 mL) by stirring for 15 min. This methanol solution containing zinc nitrate was quickly mixed with another 50 mL of methanol solution which had 0.81 g of 2-methylimidazole, 0.81 g of 1-methylimidazole and different quantities of T-ZCSv (30, 50, 100, 700, 1300, 1900 and 2500 mg), and kept stirring for a few minutes. Subsequently, the obtained solution was aged under ambient conditions without any disturbance for a day and night. Finally, the resulting sediment was repeatedly cleaned with ethanol and dried overnight. ZIF-8 is prepared in a similar process to that of ZIF-8@T-ZCSv but without the addition of T-ZCSv.

Characterization

The JSM-7401F instrument of SEM-EDS was employed to monitor the morphology, grain size, particle dispersion, and micro-distribution of the elements of the samples. JEM-2100F TEM/HRTEM equipment (180 kV of source voltage) was used to investigate the micromorphology and crystal lattice of the samples. The crystalline phases and crystalline nature of the photocatalysts were monitored using a MiniFlex 600, XRD instrument operating with $\text{Cu-K}\alpha$ radiation, voltage $\sim 40 \text{ kV}$, and current of 30 mA between 5° and 90° . The identification of surface oxidation states and chemical valency of the photocatalysts, measurement of relative elemental content and qualitative analysis of each element was performed using ESCALAB 250xi and Thermo Fisher's X-ray photoelectron spectroscope (XPS). The physical characteristics of the surface (specific surface area, pore volume, pore structure and pore size distribution) of the photocatalysts were measured using the JW-BK 100 B instrument. A HITACHI U-3310 UV-Vis diffuse reflection spectrometer (BaSO_4 served as reference material) was utilized to measure the absorbance of light by photocata-

lysts in the wavelength ranging from 200 to 800 nm, and their energy band gaps were calculated by subsequent conversion (KM plots) of the equations. A Bruker A300 electron paramagnetic resonance (EPR) spectroscope monitored the EPR signals pertaining to lattice defects in the sample at ambient temperature.

Measurement of photocatalytic hydrogen production

Photocatalytic hydrogen production was performed through an automated CEL-SPH2N system (China Education Au-light) equipped with a 250 mL reaction vessel. In the typical procedure, 10 mg (or 50 mg) of samples and sacrificial agents (1.57 g of Na_2SO_3 and 4.2 g of $\text{Na}_2\text{S}\cdot 9\text{H}_2\text{O}$) were dissolved in a 50 mL aqueous solution uniformly and taken in a quartz plate capped reaction vessel. Subsequently, the vessel was carefully mounted on the magnetic stirrer of the CEL-SPH2N system, which was set to rotate at 600 rad min^{-1} and connected to the GC-7920 online-based gas chromatograph (with highly pure N_2 as carrier gas). The system was strongly evacuated prior to irradiation for 30 min. Visible light irradiation obtained by a 300 W xenon lamp with a wavelength filter ($\geq 420 \text{ nm}$) was employed to perform the photocatalytic reaction. The cross-sectional area of the lamp was about $1.857 \times 10^{-3} \text{ m}^2$, *i.e.* the illumination area and the distance from the lamp to the reacting solution were set at 0.125 m. A cold water-flow system was used to stabilize the reaction temperature at 4°C . The online gas chromatograph measured the amount of hydrogen produced automatically every 40 min during the photocatalytic reaction, which was conducted for 200 min. The applied parameters of gas chromatography were 60 mA of bridge current, 100°C of sample inlet temperature, 150°C of TCD temperature, and 60°C of column furnace temperature.

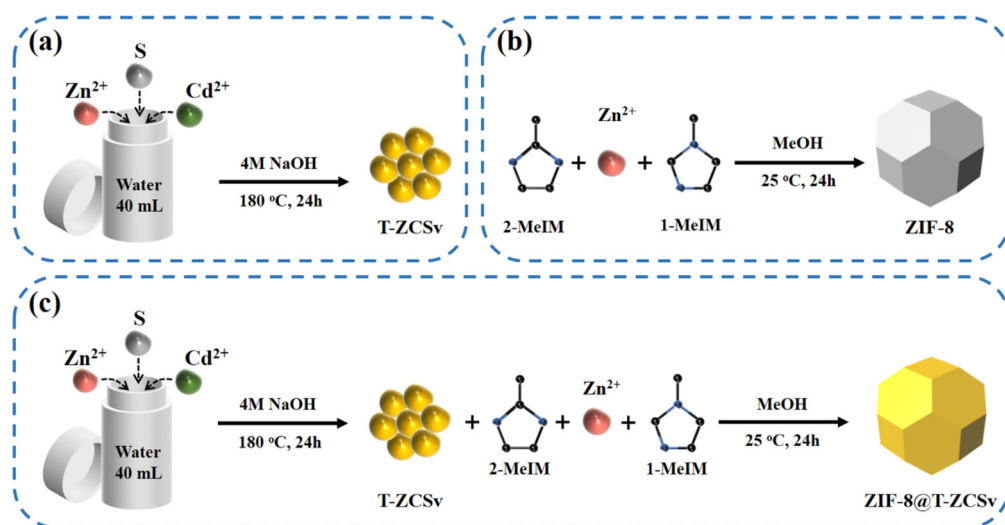
Photoelectrochemical test

Photoelectrochemical tests [transient photocurrent, electrochemical impedance spectra (EIS) and Mott-Schottky] were

conducted using the CHI-660E (Chen Hua Instrument Co., Ltd) electrochemical workstation with a three-electrode system. The working electrode was prepared as follows. The powder sample (5 mg), DI water (500 μL), ethanol (450 μL) and Nafion aqueous solution (50 μL) were taken in a beaker and sonicated for 1 h, and the mixture was dripped uniformly onto the fluorine-doped tin oxide (FTO) glass ($1.0 \times 1.8 \text{ cm}^2$) using a pipette. Subsequently, the FTO glass coated with the sample was dried at room temperature for 12 h. The three-electrode configuration contained a working electrode of a sample-coated fluorine-doped tin oxide glass substrate, a counter electrode that is a Pt wire, and a reference electrode formed with a calomel electrode or saturated Ag/AgCl . The electrolyte is a solution of Na_2SO_4 (0.1 M). The lamp used at 15 cm of the working electrode to irradiate is the same as that of the photocatalytic reaction.

Results and discussion

Typical procedures for the synthesis of T-ZCSv, ZIF-8 and ZIF-8@T-ZCSv are illustrated in Scheme 1(a–c). First, T-ZCSv was synthesized using $\text{Zn}(\text{CH}_3\text{CO}_2)_2\cdot 2\text{H}_2\text{O}$, $\text{Cd}(\text{CH}_3\text{CO}_2)_2\cdot 2\text{H}_2\text{O}$ and CH_3CSNH_2 , which serve as precursors of Zn^{2+} , Cd^{2+} and S respectively by a facile hydrothermal reaction (4 M NaOH) at 180°C for 24 h [Scheme 1(a)]. The synthesis of ZIF-8 is illustrated in Scheme 1(b), according to which a certain proportion of 1-methylimidazole, 2-methylimidazole and $\text{Zn}(\text{NO}_3)_2\cdot 6\text{H}_2\text{O}$ are aged in methanol solution for 24 h to obtain ZIF-8 polyhedra that possess a uniform size and favorable structure. Combining the above two experimental schemes, ZIF-8@T-ZCSv was prepared by employing different quantities of T-ZCSv solid solutions added to the reaction solution before the process of aging ZIF-8. Consequently, ZIF-8@T-ZCSv with different proportions of T-ZCSv were obtained and labeled as ZIF-8@T-ZCSv (30 mg), ZIF-8@T-ZCSv (50 mg), ZIF-8@T-ZCSv



Scheme 1 Synthesis processes of T-ZCSv (a), ZIF-8 (b), and ZIF-8@T-ZCSv (c).

(100 mg), ZIF-8@T-ZCSv (1300 mg), ZIF-8@T-ZCSv (1900 mg), and ZIF-8@T-ZCSv (2500 mg). These ZIF-8@T-ZCSv samples are expected to possess a uniform distribution of T-ZCSv over the surface of the ZIF-8 polyhedra.

The geometry, size, microscopic morphology, and distribution of the ZIF-8 polyhedra, pure T-ZCSv nanoparticles and ZIF-8@T-ZCSv (1300 mg) composite are analyzed by SEM images. As shown in Fig. 1a and b, the prepared ZIF-8 exhibits a typical polyhedron morphology with a stereoscopic framework, smooth surfaces, and a uniform particulate size of approximately 1.5 μm . SEM images of two different magnifi-

cations for twinned $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ nanoparticles containing sulfur vacancies are presented in Fig. 1c and d. The particle size distribution curve with the corresponding histogram is shown as an inset. These figures show that T-ZCSv nanoparticles possess inhomogeneous sizes with diameters of 20–130 nm and an average dimension of ~ 60 nm (inset). The particles do not possess a specific geometric shape. Fig. 1c and d show irregularly aggregated pure T-ZCSv nanoparticles with poor uniformity in their dispersion, which is not beneficial for the exposure of photocatalytic active sites. The SEM images of ZIF-8@T-ZCSv (1300 mg), prepared by an ageing

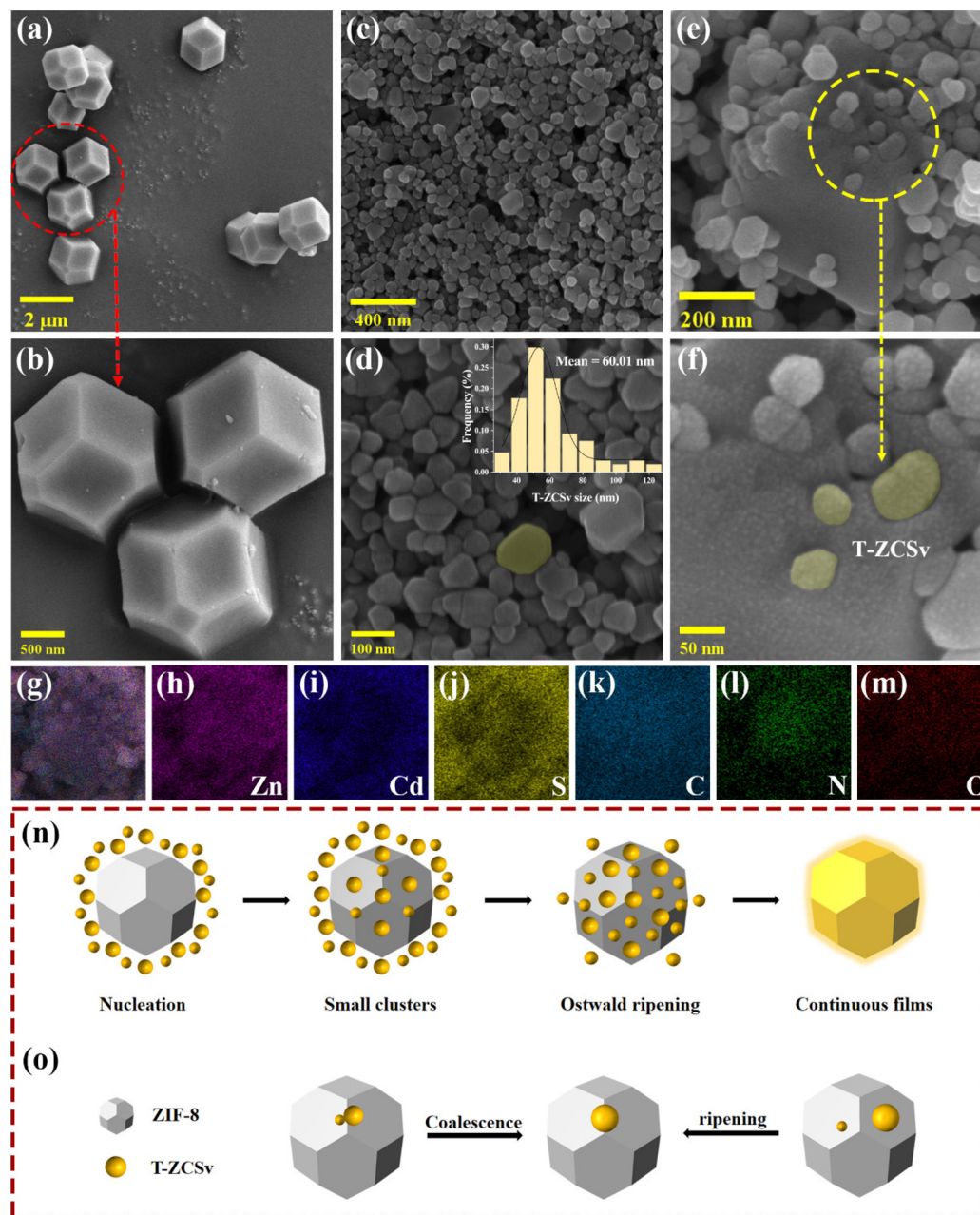


Fig. 1 SEM images of ZIF-8 (a and b), T-ZCSv and the particle size distribution plot (c and d), ZIF-8@T-ZCSv (1300 mg) (e and f); the EDS element mapping of ZIF-8@T-ZCSv (1300 mg) (g–m); the model of the Ostwald ripening process (n) and comparison of ripening and coalescence (o).

effect, are presented in Fig. 1e and f with different magnifications. The well-formed ZIF-8 polyhedron in the 3D space (Fig. 1e) supports the dispersion of pure T-ZCSv nanoparticles [highlighted with a yellow color in Fig. 1f] and their distribution over the surfaces of the ZIF-8 polyhedron, indicating a significantly reduced agglomeration of pure T-ZCSv nanoparticles. The successful formation of the ZIF-8 polyhedra and T-ZCSv nanoparticle-based composite is confirmed by the dents made by the T-ZCSv nanoparticles over the ZIF-8 polyhedra's surface, demonstrating that the surfaces of the ZIF-8 polyhedron serve as a substrate for T-ZCSv nanoparticles and prevent the nanoparticles from aggregation. This strategic composite remarkably enhanced the dispersion of T-ZCSv and facilitated more reactive sites thereby excellent exposure to the light and the aqueous environment for quick photocatalytic reaction with high light absorbance over the surface of the T-ZCSv nanoparticles. The corresponding EDS elemental distribution in ZIF-8@T-ZCSv (1300 mg) is shown in Fig. 1(g–m), depicting the existence of zinc, cadmium, sulphur, carbon, nitrogen and oxygen throughout ZIF-8@T-ZCSv (1300 mg). Fig. 1n and o depict the mechanism of formation of the ZIF-8@T-ZCSv composite with highly dispersed nanoparticles on the surfaces of ZIF-8 based on the Ostwald ripening process. According to this process, the ZIF-8 polyhedron acts as the substrate of T-ZCSv nanoparticles, and some of the small T-ZCSv nanoparticles gradually dissolve into each other and form small clusters over the surface of ZIF-8. The smaller T-ZCSv nanoclusters can be converted into larger agglomerates at the expense of the mass of tiny T-ZCSv nanoparticles. This process makes the average size of the T-ZCSv nanoparticles gradually increase and uniformly deposit on the surfaces of ZIF-8, resulting in the formation of ZIF-8@T-ZCSv composite, which is in good agreement with the SEM studies.

The microscopic morphology and crystalline lattice structure of T-ZCSv and ZIF-8@T-ZCSv (1300 mg) were further measured by TEM images. Fig. (2a and b) show the T-ZCSv nanoparticles agglomerated together and composed of many irregular granules at different magnifications. The bright parallel straight-line patterns are seen on the surface of the nanoparticles rounded by the red dotted circles in the TEM images with 100- and 25 nm scale bars. These straight lines can be attributed to the wurtzite twin boundaries (WZ-TB) of the sphalerite zinc blende (ZB) lattices. Fig. 2b represents the magnified image of a rounded nanoparticle (highlighted with a translucent yellow area) with clearly visible twin boundaries. To confirm the twinning of sphalerite–Zinc Blende and Wurtzite-TBs, we analyzed the HRTEM images of the samples. Fig. 2c is the HRTEM image of the T-ZCSv's lattice structure and its generated Fast Fourier Transform (FFT). Different regions of the nanoparticles are indexed using the FFTs generated from the portions. The nanocrystal regions I, II, III and IV are confirmed as sphalerite–Zinc Blende lattice structures of the ZnS crystalline phase segments by indexing the corresponding FFT patterns. Regions I and II are indexed based on the zone axes (viewed along) $[110]$ and $[1\bar{1}0]$ respectively, depicting the mirrored symmetrical alignment of atomic

planes of the ZB structure in these two regions. Similarly, regions III and IV are indexed with the same zone axes, respectively. These four regions are separated by atomic stacking fault (SF), which form WZ-TBs or dislocation boundaries (indicated by white dashed arrows) with parallel distribution,^{13,24} confirming that the occurrence of $(111)/[11\bar{2}]$ -slips interrupts the continuity in the ZB structure, resulting in twin structures. These twin crystals can be associated with homojunctions originating from the WZ twin boundary slips in the T-ZCSv. Fig. 2g schematically represents the ZB phase with a face-centred cubic (FCC) structure of “...ABCABC...” order in the $\langle 111 \rangle$ direction of its tightly packed structure. The WZ phase with a hexagonal structure possesses “...ABABAB...” order of atomic planes in the closely-packed $[0001]$ direction.³⁰ The appearance of periodic stacking faults in the “...ABCABC...” ordered arrangement of the FCC structure along the $\langle 111 \rangle$ crystal plane introduces discontinuities in the sphalerite ZB's structure. Consequently, dislocations are created at discontinuities with “...CAC...” or “...BAB...” stacking of atomic planes (Fig. 2h). These dislocations distinguish the neighboring lattice segments with “A” as a mirror symmetry which are thus known as coherent twin boundaries or coherent twin planes. These “...CAC...” or “...BAB...” twin planes belong to the wurtzite structure, and therefore the twin boundaries are called wurtzite twin boundaries (WZ-TBs), forming twinned superlattices. Fig. 2(c and c-1) depicts mirror-symmetric lattice (zigzag) stripes distributed on both sides of the twin boundaries, and the order of twin lattice arrangement changed to “...CAC...” or “...BAB...” or *vice versa*. Several wurtzite structural phases are identified on many T-ZCSv nanocrystals in the present samples. The wurtzite structure can also occur with two or three diatomic layers, as shown in Fig. S1† for the T-ZCSv nanocrystals. These ZB/WZ twin boundaries produce interfacial-homogeneous junctions, which remarkably influence the electronic properties of the T-ZCSv nanocrystals and thus effectively boost the vectorial separation and transport of charge carriers.^{15,31} Fig. 2d and e are the TEM images of ZIF-8@T-ZCSv (1300 mg) monitored at different resolutions that describe the T-ZCSv nanocrystals held over the structure of ZIF-8. The granular T-ZCSv (translucent yellow area in Fig. 2e) is distributed around the ZIF-8 polyhedron and is held with ZIF-8 by close contact (marked with a red dashed line). Fig. 2d confirms that the T-ZCSv nanocrystals are well dispersed over the surfaces of ZIF-8 compared to the independent clusters of the T-ZCSv nanocrystals (Fig. 2a). Fig. 2d and e show no clusters of T-ZCSv nanoparticles, suggesting that the dispersion of T-ZCSv was significantly improved due to the introduction of ZIF-8, which is consistent with SEM results. Furthermore, Fig. 2f shows twinned ZB crystal structures associated with WZ-twin boundaries in the T-ZCSv nanocrystals for ZIF-8@T-ZCSv (1300 mg). The longitudinal columns of the ZB structures distinguished by the WZ structure are confirmed by the FFT pattern indexed based on the $[011]$ zone axis shown as an inset. The patterns indicate the appearance of $(111)/[11\bar{2}]$ -slips in the twinned nanocrystals, confirming that the twinning characteristics of T-ZCSv's structure is

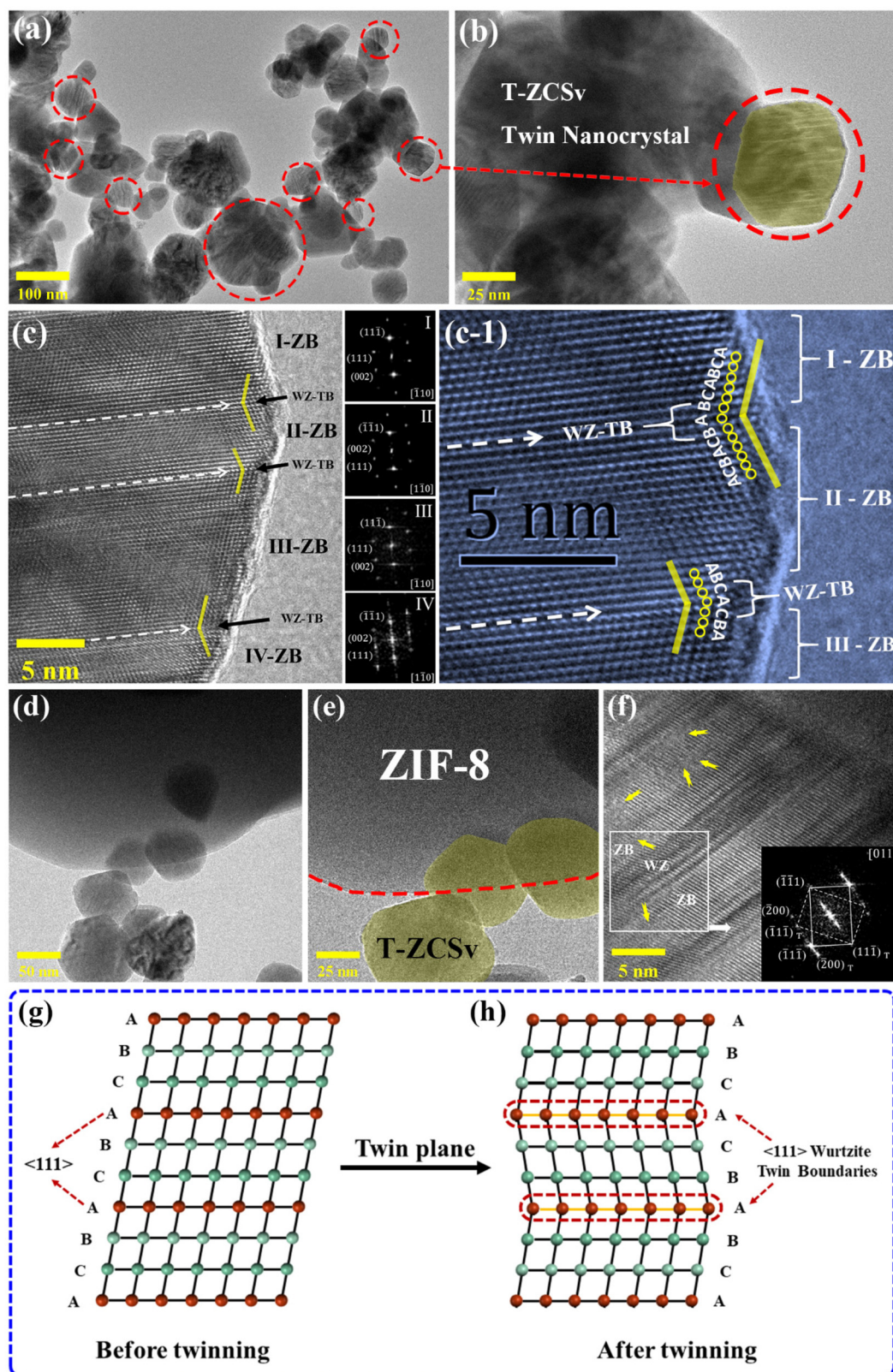


Fig. 2 TEM images of T-ZCSv (a–c) and ZIF-8@T-ZCSv (1300 mg) (d–f) at different magnifications. The corresponding structural model of the twinning nanocrystal (g and h).

unaffected by the presence of ZIF-8. Furthermore, the lattice of the ZIF-8@T-ZCSv (1300 mg) composite exhibits smudged regions marked with yellow arrows representing the lattice defects which can be attributed to the S vacancies. These distinctive structural features are expected to display unusually high activity in photocatalytic reactions.

XRD technique was employed to study the characteristic crystallinity and crystal phases for the series of photocatalysts. Four different kinds of $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solution, including ZCS, ZCS-V, T-ZCS and T-ZCSv samples exhibit different crystal properties in Fig. 3a. The XRD pattern of the S vacancy containing-ZCS-V and ZCS exhibits the diffraction peaks corresponding to wurtzite CdS (JCPDF#41-1049) and wurtzite ZnS (JCPDF#36-1450), respectively.³² For the ZCS, the characteristic diffraction peaks located around $2\theta \sim 24.8^\circ$ (100), 26.4° (002), 28.2° (101), 43.8° (110), and 51.9° (112) are attributed to the planes given within the parentheses, which belong to the hexagonal wurtzite $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ phase. It is observed that the peak intensity of ZCS-V decreases, and the position of the peaks shifts to a higher angle in comparison with that of ZCS, which is ascribed to the structural modification due to the presence of S vacancies and corresponding surface disorder. However, the positions of the diffraction peaks pertaining to ZCS and ZCS-V are located at the angular positions that lie between CdS

and ZnS, which means that the prepared $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ is formed into a solid solution of ZnS and CdS rather than an ordinary physical combination of ZnS and CdS.^{14,25} Besides, the XRD patterns of T-ZCS and T-ZCSv with twin crystal exhibit two sets of characteristic peaks corresponding to Wurtzite CdS (JCPDF#41-1049) and Zinc-Blende ZnS (JCPDF#05-0566), respectively.³³ The position of characteristic peaks for T-ZCS are located between that of CdS and ZnS *i.e.*, at 25.4° , 26.9° , 28.4° , 44.6° and 52.9° which is ascribed to the theoretical $0.5:0.5 = \text{Zn}:\text{Cd}$ (molar ratio). The peaks belong to the (100), (002), (101), (110), (112) planes of wurtzite CdS and (111), (220), (311) planes of zinc blende ZnS. Some additional peaks of ZnS observed for T-ZCS are assigned to different solubility limits and nucleation rates, and this phenomenon has been confirmed in our previous reports.³⁴ Moreover, a set of sharp peaks of T-ZCSv are found at 27.6° , 45.6° , and 54.1° , which are attributed to (111), (220), and (311) planes of zinc blende ZnS and (002), (110), and (112) planes of wurtzite CdS, indicating that the T-ZCSv solid solution was successfully prepared. Two sets of diffraction peaks attributed to the ZB and WZ phases are observed for T-ZCS and T-ZCSv, suggesting the presence of the W/Z phase junctions and twin crystalline structure in the solid solution.

The XRD patterns of ZIF-8 and twinned T-ZCSv nanocrystals with S vacancy, including their composites with different

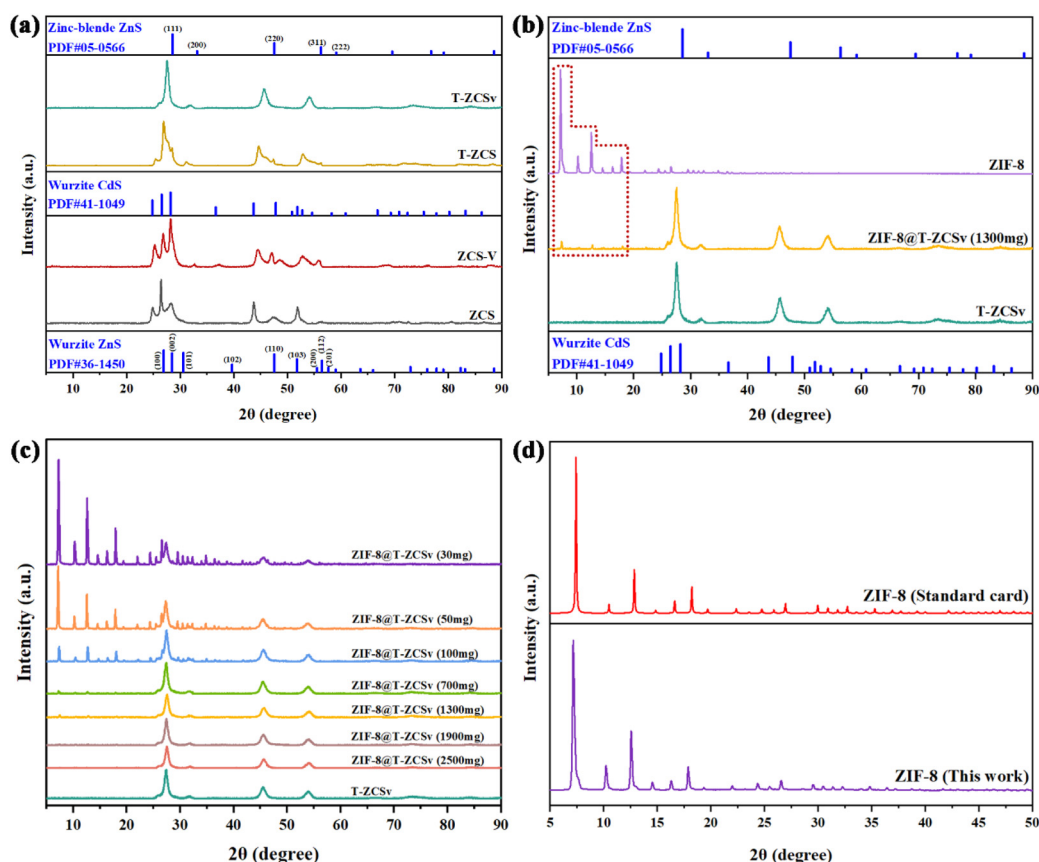


Fig. 3 XRD patterns of ZCS, ZCS-V, T-ZCS and T-ZCSv with different characteristics (a), ZIF-8, T-ZCSv and ZIF-8@T-ZCSv (b), ZIF-8@T-ZCSv with different quality of T-ZCSv (c) and ZIF-8 with its standard cards (d).

amounts of T-ZCSv, are presented in Fig. 3(b–d). The diffraction pattern of the ZIF-8 polyhedron exhibits three sharp and intense diffraction peaks at 7.2° , 12.6° and 17.9° which are well matched with the CIF file of ZIF-8 standard card (CCDC: 864309†) of Mercury software, confirming the high crystallinity of the synthesized ZIF-8.²⁹ Additionally, the typical peaks corresponding to T-ZCSv and ZIF-8 were observed in the diffraction pattern of the ZIF-8@T-ZCSv composites. The relative intensity of the ZIF-8's distinct peaks seem to decrease gradually with the content of T-ZCSv and almost disappeared when the mass of T-ZCSv exceeded 1900 mg due to the relatively low concentration of ZIF-8. In other words, the intensity of T-ZCSv's diffraction peaks increases significantly with the ratio of T-ZCSv. However, the position of diffraction peaks pertaining to either ZIF-8 and T-ZCSv show no observable shift for ZIF-8@T-ZCSv, indicating that the formation of the ZIF-8@T-ZCSv composite with different masses of T-ZCSv did not affect the crystal structure and atomic alignment of each of the components and confirmed the formation of the composite using an ageing method. No characteristic diffraction peaks of the compounds other than T-ZCSv and ZIF-8 were found in the composites, suggesting that no new crystalline phases or compounds were produced during the synthesis of ZIF-8@T-ZCSv and confirming that the introduction of ZIF-8 did not affect the crystallinity and purity of T-ZCSv. XRD studies indicate that the ZIF-8 acts as a substrate in composites, providing space to improve redistribution and thus facilitates a wider exceptional dispersion of T-ZCSv. The results are consistent with those of the SEM and TEM studies.

The surface physical adsorption properties of ZIF-8 polyhedra, ZIF-8@T-ZCSv composites, and pure T-ZCSv nanoparticles were explored *via* the BET test at 77 K. The N_2 adsorption–desorption plot of ZIF-8 (Fig. 4a) showed a type I isotherm conforming to the BDDT (Brunauer–Deming–Demin–Teller) classification. N_2 starts to fill the micropores of ZIF-8 in the region of low pressure ($0 < P/P_0 < 0.05$), resulting in a rapid increase in gas adsorption and a plateau region without an

observable increase ($0.05 < P/P_0 < 0.97$). Besides, the adsorption isotherm showed a slow increasing trend when the relative pressure (P/P_0) approaches 1.0, and no hysteresis loop was observed for ZIF-8, suggesting that ZIF-8 possesses micropores. On the other hand, the N_2 adsorption–desorption isotherms of T-ZCSv showed a rise in the high-pressure region after 0.9, which is attributed to capillary coagulation and defined as a typical type IV with a narrow H3-type hysteresis loop, confirming the presence of a mesopore structure. The N_2 adsorption–desorption isotherms of different ZIF-8@T-ZCSv composites showed a similar shape to that of ZIF-8 and T-ZCSv based on the mass of T-ZCSv. The isotherm of the composite with a lower T-ZCSv mass depicts close features of ZIF-8, and the composite with a higher T-ZCSv mass exhibits a similar shape to T-ZCSv's isotherm. The measured physical properties, such as the total surface area of the samples, the average volume and size of the pores for ZIF-8, T-ZCSv and various ZIF-8@T-ZCSv composites, are given in Table 1. The specific surface area of ZIF-8, ZIF-8@T-ZCSv (100 mg), ZIF-8@T-ZCSv (700 mg), ZIF-8@T-ZCSv (1300 mg), ZIF-8@T-ZCSv (1900 mg), ZIF-8@T-ZCSv (2500 mg) and T-ZCSv were measured and found to be $1302.011 \text{ m}^2 \text{ g}^{-1}$, $448.441 \text{ m}^2 \text{ g}^{-1}$, $121.227 \text{ m}^2 \text{ g}^{-1}$, $77.586 \text{ m}^2 \text{ g}^{-1}$, $45.239 \text{ m}^2 \text{ g}^{-1}$, $34.187 \text{ m}^2 \text{ g}^{-1}$ and $17.328 \text{ m}^2 \text{ g}^{-1}$, respectively. The specific surface area for the ZIF-8@T-ZCSv composites is gradually reduced with the

Table 1 Physical surface properties of various samples

| Samples | Surface area ($\text{m}^2 \text{ g}^{-1}$) | Pore volume ($\text{cm}^3 \text{ g}^{-1}$) | Pore size (nm) |
|------------------------|--|--|----------------|
| ZIF-8 | 1302.011 | 0.753 | 2.313 |
| ZIF-8@T-ZCSv (100 mg) | 448.441 | 0.424 | 3.784 |
| ZIF-8@T-ZCSv (700 mg) | 121.227 | 0.208 | 6.872 |
| ZIF-8@T-ZCSv (1300 mg) | 77.586 | 0.160 | 8.919 |
| ZIF-8@T-ZCSv (1900 mg) | 45.239 | 0.164 | 14.531 |
| ZIF-8@T-ZCSv (2500 mg) | 34.187 | 0.140 | 16.356 |
| T-ZCSv | 17.328 | 0.155 | 34.387 |

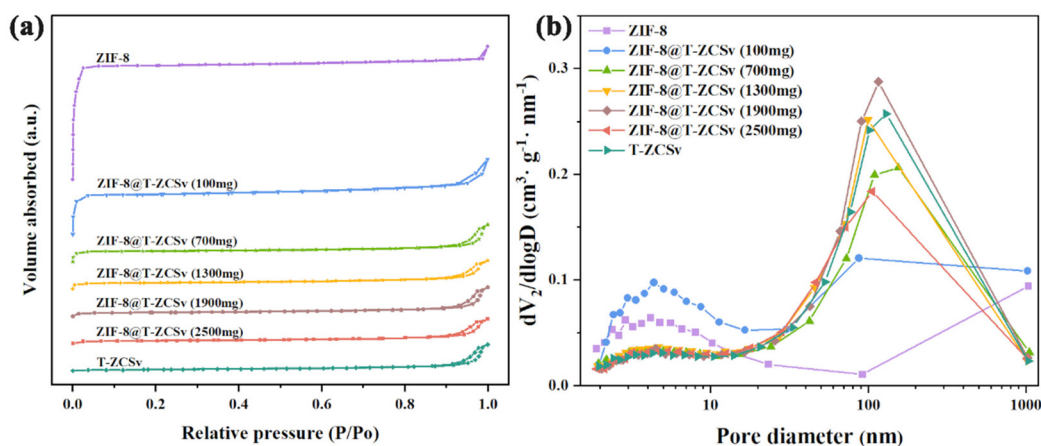


Fig. 4 N_2 adsorption/desorption analysis spectra (a) and the pore size distribution (b) of the pristine T-ZCSv, different ZIF-8@T-ZCSv, and ZIF-8 samples.

amount of T-ZCSv. Additionally, the pore size increased with the amount of T-ZCSv, indicating a good dispersion of T-ZCSv particles over the surface of ZIF-8 and preventing the aggregation of T-ZCSv particles. The variation in surface physical characteristics for the ZIF-8@T-ZCSv series confirms the successful synthesis of the composite with the expected uniform distribution of T-ZCSv particles on the surface of ZIF-8. This enabled one of the members, ZIF-8@T-ZCSv (1300 mg), to possess more reactive center sites that adsorb reactants for high photocatalytic activity.

XPS was employed to determine the chemical compositions and chemical states of ZIF-8, T-ZCSv, and ZIF-8@T-ZCSv composites with different contents of T-ZCSv in Fig. 5. The typical XPS survey spectra (Fig. 5a) of various samples show the existence of zinc, cadmium, sulphur, and carbon in T-ZCSv and Zn, C, N elements in ZIF-8, and the coexistence of C, N, Zn, Cd, and S in ZIF-8@T-ZCSv, indicating the successful formation of ZIF-8@T-ZCSv composites. The XPS spectra were calibrated to the C 1s peak of 284.8 eV (Fig. 5b) of ZIF-8. The high-resolution comparative spectra of ZIF-8 and three composites in the

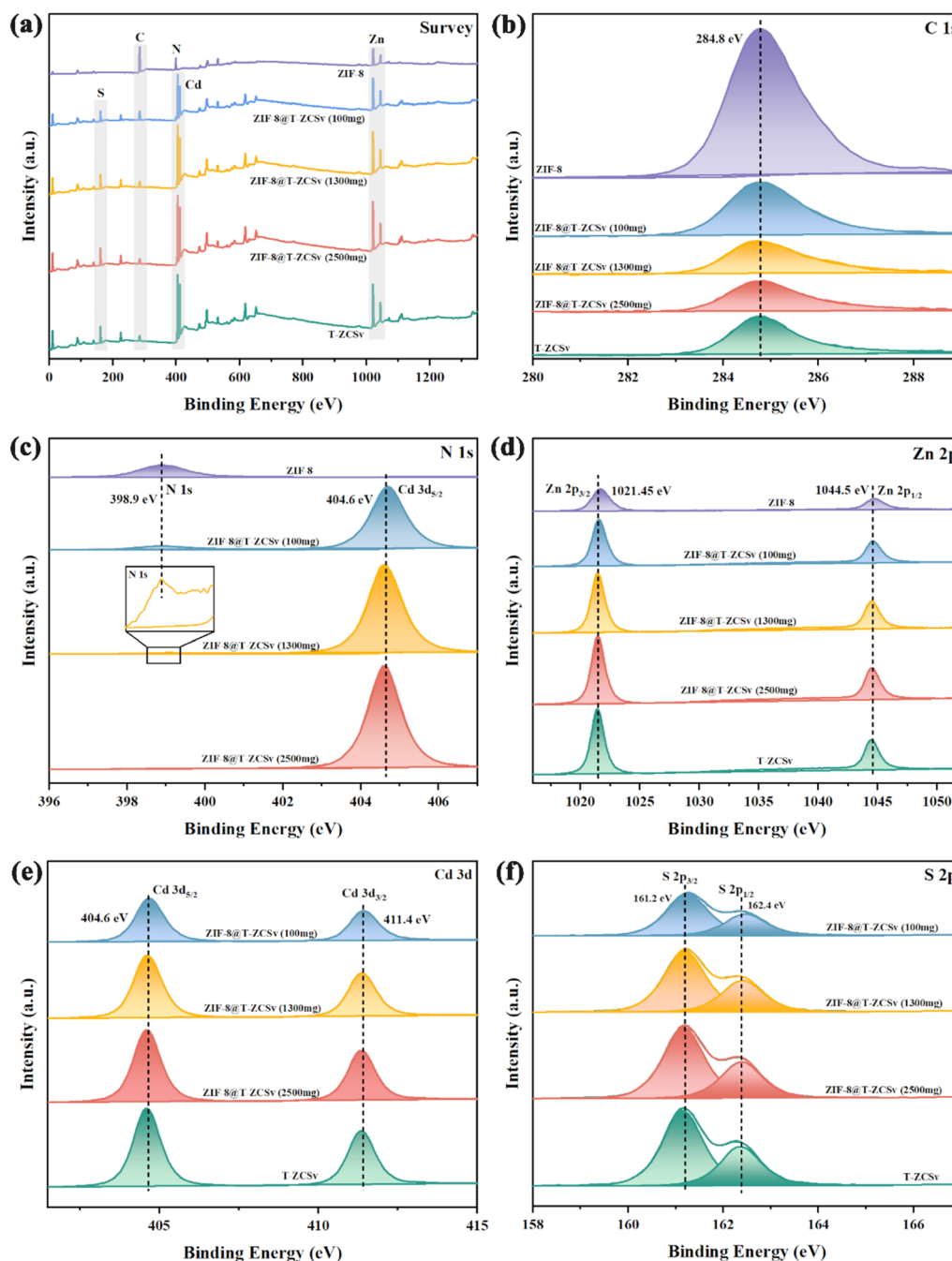


Fig. 5 Survey spectra (a) and high-resolution XPS spectra of pristine T-ZCSv and ZIF-8@T-ZCSv for C 1s (b), N 1s (c), Zn 2p (d), Cd 3d (e), and S 2p (f).

region from 396 eV to 407 eV (Fig. 5c) showed N 1s for ZIF-8 at 398.9 eV and N 1s and Cd 3d_{5/2} for the composites. The relative intensity of N 1s of ZIF-8 decreased with the T-ZCSv content as the relative content of ZIF-8 decreased in the ZIF-8@T-ZCSv composites, but the N 1s peak exists in all compounds (Inset).

Furthermore, the high-resolution Zn 2p comparative spectra of all the samples depict two sharp peaks located at ~1021.4 and 1044.5 eV (Fig. 5d) corresponding to the doublet of Zn 2p_{3/2} and Zn 2p_{1/2}. The Cd 3d binding energy region compared to the spectra of three composites and T-ZCSv (Fig. 5e) contains a doublet of Cd 3d_{5/2} (404.6 eV) and Cd 3d_{3/2} (411.4 eV). The S 2p region of the spectra (Fig. 5f) for three composites and T-ZCSv are resolved into two typical binding energy peaks at ~161.2 and 162.4 eV, which correspond to S 2p_{3/2} and S 2p_{1/2}, respectively. The separation of the Zn 2p, Cd 3d and S 2p doublet-peaks is approximately 23.1 eV, 6.8 eV and 1.2 eV, respectively, confirming the existence of Zn²⁺, Cd²⁺ and S²⁻ states in the twinned T-ZCSv with WZ and ZB. Moreover, these peaks become more intense with the amount of T-ZCSv, and no observable shift in the peak positions for the composites, evidencing the increased mass of T-ZCSv negligibly affected structure of the ZIF-8 framework. However, for T-ZCSv nanoparticles, the S 2p peaks shifted to a lower binding energy position. It can be explained based on the existing S vacancies, which act as strong electron-deficient groups, and cause a distortion in the equilibrium of the electron cloud of the T-ZCSv's surface due to the low coordination resulting from the escape of S atoms.^{35,36} Consequently, electron transport is boosted and the lifetime of electrons is prolonged during the photocatalytic reaction. The investigation confirmed that ZIF-8 serves merely as the substrate that provides more space for T-ZCSv to avoid the aggregation and does not affect the twinned structure and vacancies. This structural feature dramatically enhances the dispersion of T-ZCSv to facilitate more active sites for exposure, which agrees with the SEM and BET analysis.

EPR spectra (Fig. 6) were recorded to further investigate the S vacancies over the surface of ZCS, ZCS-V, T-ZCS, and T-ZCSv. Vacuum deoxidation treatment was performed to avoid the interference of adsorbed oxygen on the surface of the samples for all products prior to the EPR test. There are no obvious paramagnetic signals of ZCS and T-ZCS in the EPR spectra. However, the spectra of ZCS-V and T-ZCSv exhibit two typical symmetrical EPR signals peaked at $g = 2.003$, indicating the appearance of S vacancies on the surface of these two samples,³⁷ converging with XPS studies. EPR studies demonstrate that an appropriate amount of NaOH and hydrazine hydrate with a strong alkali property, used as a reducing agent in the hydrothermal reaction process, can control the formation of sulfur vacancies by providing a powerful reducing environment with relatively low formation energy favorable for the formation of S vacancies.^{32,38} Besides, excess S sources (thiourea and thioacetamide) will accumulate on the surface of the nanocrystal and considerably hinder the formation of the crystals, which is also one of the crucial reasons for the formation of S vacancies.³⁹ The introduction of S vacancies plays

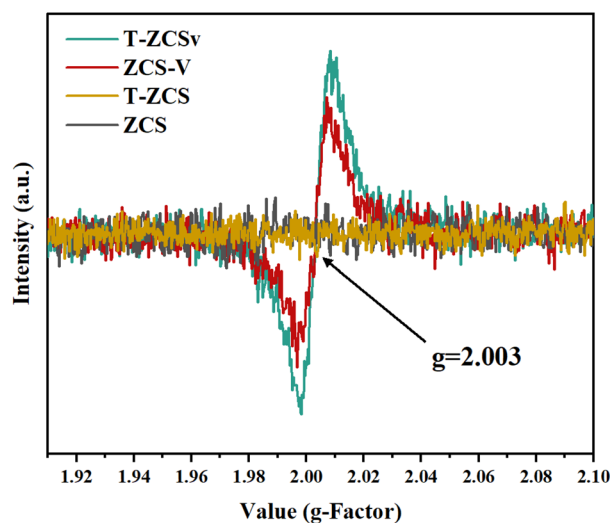


Fig. 6 EPR spectra of the ZCS, ZCS-V, T-ZCS and T-ZCSv.

a critical role in promoting photocatalytic activity. Firstly, S vacancies may change the band structure of electron energy, induce the reorganization of the crystal lattice, and create unsaturated active sites that act as electron traps. These active sites could also contribute to enhancing the space-charge separation and effectively improve the kinetic properties of photo-generated charges. Besides, S vacancies accelerate the electron migration from T-ZCSv to the vacancy sites due to their electron-deficient solid characteristics. Consequently, an effective separation and migration of photogenerated electrons and holes can be achieved over the surface of T-ZCSv, which effectively supports the photocatalytic production of hydrogen.

Fig. 7 shows the UV-vis spectra and measured Tauc plots to investigate the light absorption properties and band gap of four types of Zn_{0.5}Cd_{0.5}S solid solutions with distinctive characteristics (Fig. 7a and c), ZIF-8 and ZIF-8@T-ZCSv (1300 mg) (Fig. 7b and d). The absorption edges correspond to the fundamental absorption of ZCS, T-ZCS, ZCS-V and T-ZCSv were located at about 549, 517, 510 and 500 nm, respectively in Fig. 7a, indicating that bound electrons of all samples can be excited by visible light and thus the samples showed excellent visible absorption capacity. The absorption edges of ZCS-V, T-ZCS and T-ZCSv showed a blue shift toward a shorter wavelength compared to those of ZCS. T-ZCSv exhibits the largest blueshift over 49 nm, resulting from the coexistence of twinned structure and S vacancies associated with the defective state energy level.⁴⁰ The optical absorption characteristics of ZIF-8 (Fig. 7b), show an absorption edge at 248 nm (inset), but the ZIF-8@T-ZCSv (1300 mg) composite exhibits the same absorption edge as T-ZCSv (500 nm). This demonstrates that the ZIF-8 polyhedron, as a metal-organic framework, can be excited by ultraviolet light and thus has poor visible-light photocatalytic performance. Consequently, the presence of ZIF-8 polyhedron in the ZIF-8@T-ZCSv (1300 mg) composite has negligible influence on the visible light sensitivity of T-ZCSv, demonstrating that the ZIF-8 acts as a substrate frame-

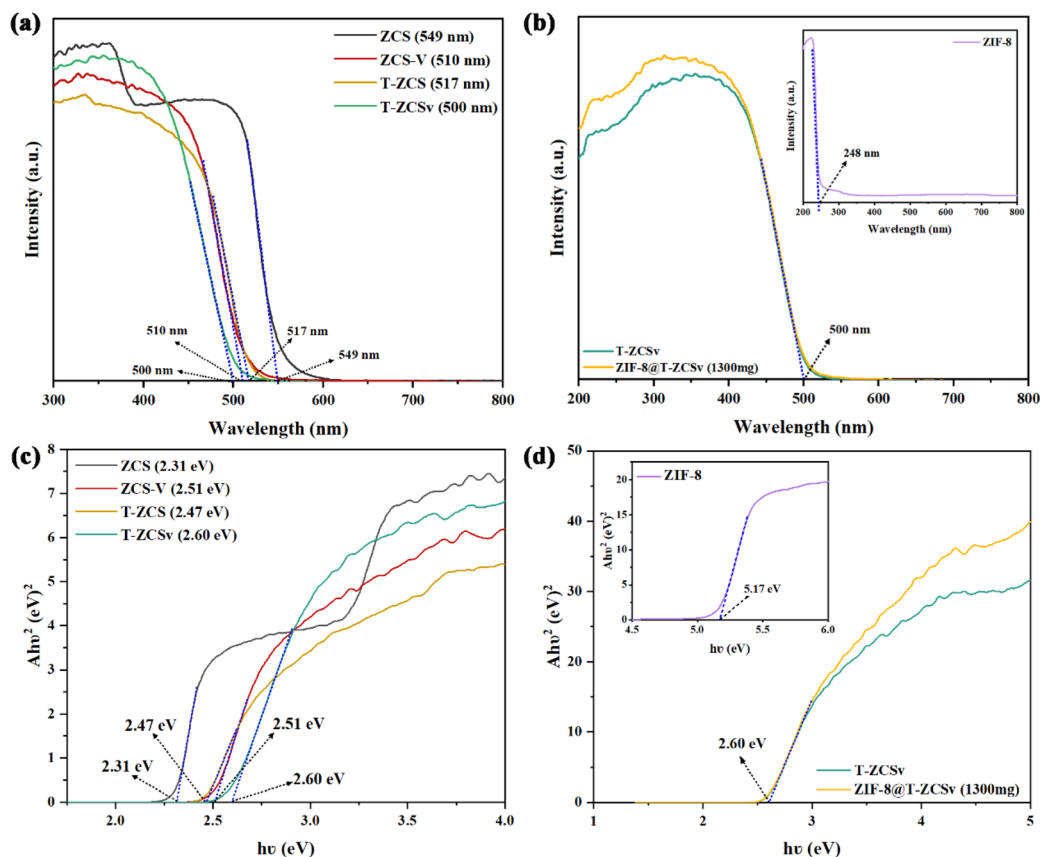


Fig. 7 UV-vis diffuse reflectance spectra and band gap evaluation from the corresponding Tauc plots: (a and c) ZCS, ZCS-V, T-ZCS and T-ZCSv samples; (b and d) ZIF-8 and ZIF-8@T-ZCSv.

work to improve the dispersion ability of T-ZCSv, which is supposed to lead to efficient photocatalysis. In addition, the corresponding band gap energies of various samples were calculated using the Tauc plots with the Kubelka–Munk function, according to the following formula:

$$\alpha h\nu = A(h\nu - E_g)^n$$

where α is the absorption coefficient; h , the Planck's constant; ν , the light frequency; E_g , the band gap energy; $n = 2$ for the indirect-band gap semiconductors and $n = 1/2$ for direct-band gap semiconductors. Based on the Tauc plot given in Fig. 7c, the direct band gap values of the samples were calculated. The direct-band gaps of ZCS, T-ZCS, ZCS-V and T-ZCSv are 2.31 eV, 2.47 eV, 2.51 eV and 2.60 eV, respectively. Additionally, the conduction band (CB) of various samples were obtained by Mott–Schottky curves in Fig. S2(a–d).† The CB potentials (E_{CB}) of ZCS, T-ZCS, ZCS-V and T-ZCSv are measured as -0.39 V, -0.42 V, -0.43 V and -0.60 V (vs. NHE), respectively. According to the equation: $E_{VB} = E_g + E_{CB}$, the corresponding VB potentials (E_{VB}) of ZCS, T-ZCS, ZCS-V and T-ZCSv were 1.92 V, 2.09 V, 2.04 V and 2.00 V (vs. NHE) and the corresponding energy band structure information are shown in Table S2.† Furthermore, the band gap energy of ZIF-8@T-ZCSv (1300 mg) was measured

and found to be 2.60 eV, which is equal to that of T-ZCSv (Fig. 7d), indicating that the wider band gap-ZIF-8 (5.17 eV) shows little impact on the bandgap structure of T-ZCSv upon the formation of the ZIF-8@T-ZCSv (1300 mg) composite. The different optical absorption of the samples was attributed to the presence of S vacancies, twin crystal structure, and the improved dispersion of the T-ZCSv nanoparticles provided by ZIF-8.

The transient photocurrent (TPC) measurements were carried out to investigate the photogenerated charge carrier dynamics, including charge carrier separation and migration in four different $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ photocatalysts. A series of on–off irradiation cycles (each for every 30 s) was employed using a 300 W Xe Lamp ($\lambda \geq 420$ nm). The transient photocurrent density (Fig. 8a) of ZCS-V, T-ZCS and T-ZCSv are significantly higher than that of ZCS, indicating better separation of free electrons and holes provided by the twinned crystal structure and appropriate S vacancies of $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$. Specifically, T-ZCSv exhibits the highest photocurrent density in the series with the order of T-ZCSv > T-ZCS > ZCS-V > ZCS. This trend indicates that the charge transfer in the ZCS with a twinned structure is faster than that of the ZCS with S vacancies. The movement of charges across the interfaces of the ZB and WZ phases contributes to their faster transport.²⁴ The highest TPC density of

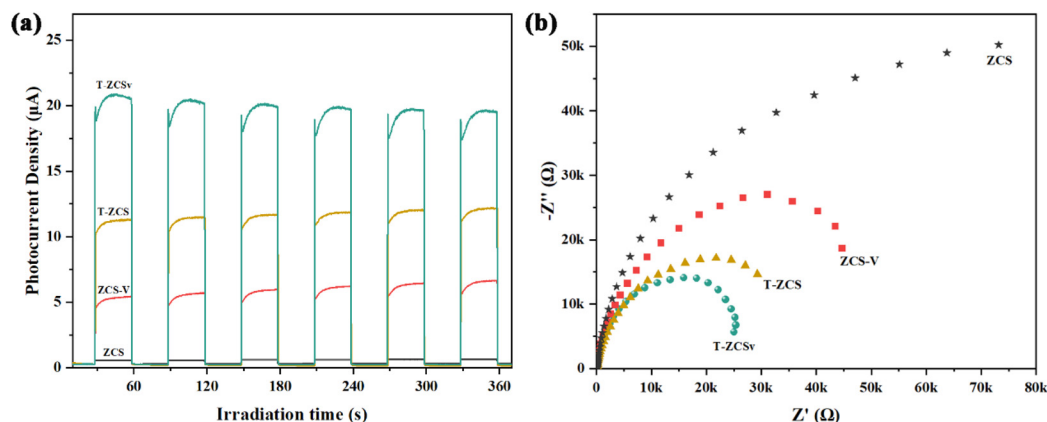


Fig. 8 Photocurrent response (a) and electrochemical impedance spectroscopy (b) of the ZCS, ZCS-V, T-ZCS, and T-ZCSv.

T-ZCSv among the four samples confirms that the twin structure and S vacancies are crucial for further improvement of photocatalytic performance. Electrochemical impedance spectra (EIS) of Nyquist plots (Fig. 8b) display varying curvature radii for different samples of the series, which are in the order $T-ZCSv < T-ZCS < ZCS-V < ZCS$. The shorter the curvature radius, the lower the charge transfer resistance, implying quicker electron and hole transfer and better electrical conductivity. Therefore, T-ZCSv possesses the lowest charge transfer resistance compared to other $Zn_{0.5}Cd_{0.5}S$ photocatalysts. Combining the TPC and EIS measurements, it can be confirmed that the T-ZCSv exhibits faster charge transfer pathways and less probable charge recombination. The introduced twinned crystal structure and surface S vacancies alter the structural properties of $Zn_{0.5}Cd_{0.5}S$, which probably leads to a remarkable improvement in photocatalytic hydrogen production.

To evaluate the rate of photocatalytic production of hydrogen (RPH) of the prepared four kinds of $Zn_{0.5}Cd_{0.5}S$ solid solutions (including ZCS, ZCS-V, T-ZCS and T-ZCSv) and different ZIF-8@T-ZCSv composites in a solution of Na_2SO_3 (0.25 M) and $Na_2S \cdot 9H_2O$ (0.35 M) as sacrificial agents, a Xe lamp (300 W, $\lambda > 420$ nm) was used to irradiate the samples for 200 min. A 50 mg of sample was added to the reacting solution to carry out the photocatalytic H_2 production experiment for different photocatalysts. Fig. 9a represents the RPH against time for ZCS, ZCS-V, T-ZCS and T-ZCSv, whereas Fig. 9b depicts a histogram for the RPH of the different samples. The measured RPH values are compared in Table 2 when 50 mg of the samples are used. Pure ZCS, without S vacancies and twinned crystal structure, exhibited the lowest RPH, i.e., $32.0 \mu mol h^{-1}$ or $0.64 mmol g^{-1} h^{-1}$. However, the ZCS-V with S vacancies and the T-ZCS with twinned-crystal structure showed improved RPH $329.44 \mu mol h^{-1}$ or $6.58 mmol g^{-1} h^{-1}$ and $412.92 \mu mol h^{-1}$ or $8.25 mmol g^{-1} h^{-1}$ respectively, which are 10.3 and 12.9 times that of the pristine ZCS, respectively under the similar experimental conditions. Interestingly, T-ZCSv with S vacancies and twin-crystalline structure displays an impressive hydrogen evolution activity of nearly $551.74 \mu mol h^{-1}$ or $11.03 mmol g^{-1} h^{-1}$.

h^{-1} , which are 1.3, 1.7 and 17.3 times that of T-ZCS, ZCS-V and ZCS, respectively. The enhanced RPH performance is attributed to S vacancies and the twin-crystalline structure containing a WZ/ZB interfacial homogeneous junction, which facilitates better electron trapping and prolonged electron lifetime and promotes charge separation. In addition, sulfur vacancies serve as active sites that adsorb oxygen in H_2O molecules during the photolytic splitting of the water reaction, thus improving the efficiency of photocatalytic hydrogen production.

Selecting T-ZCSv that exhibited the highest hydrogen production activity, a series of ZIF-8@T-ZCSv composites were synthesized by varying the mass of T-ZCSv. T-ZCSv serves as the core catalyst, and the ZIF-8 polyhedron with a large specific surface area acts as a substrate that supports a uniform dispersion of T-ZCSv nanoparticles over its surface. A comparative study of hydrogen production activity carried out against time for T-ZCSv, and ZIF-8@T-ZCSv with varied masses of T-ZCSv is given in Fig. 9c. Fig. 9d is the histogram of hydrogen production activity for the T-ZCSv and ZIF-8@T-ZCSv composites with 10 mg of the samples used in the production of hydrogen. The RPH of the ZIF-8@T-ZCSv composites were found to increase with the amount of T-ZCSv to 1300 mg. The maximum RPH exhibited by ZIF-8@T-ZCSv (1300 mg) is $583.25 \mu mol h^{-1}$ or $58.32 mmol g^{-1} h^{-1}$, which is almost 1.4 times that of T-ZCSv ($439.92 \mu mol h^{-1}$ or $43.99 mmol g^{-1} h^{-1}$). Meanwhile, the optimal ZIF-8@T-ZCSv (1300 mg) was performed for 10 h to verify long-term activity and the results are shown in Fig. S3.† The hydrogen production efficiency of ZIF-8@T-ZCSv (1300 mg) is not only undiminished with the prolongation of the reaction time, but was also still relatively stable and the high activity after 10 h was about $655.31 \mu mol h^{-1}$ or $65.53 mmol g^{-1} h^{-1}$. Additionally, a comparative study of the hydrogen production rates of $Zn_{0.5}Cd_{0.5}S$ -based composites reported in the recent literature was also performed in Table S2.† In our present work, ZIF-8@T-ZCSv (1300 mg) exhibits excellent photocatalytic hydrogen production activity for the composites, which is better than that of other $Zn_xCd_{1-x}S$ -based photocatalysts. The main reason for the improvement of

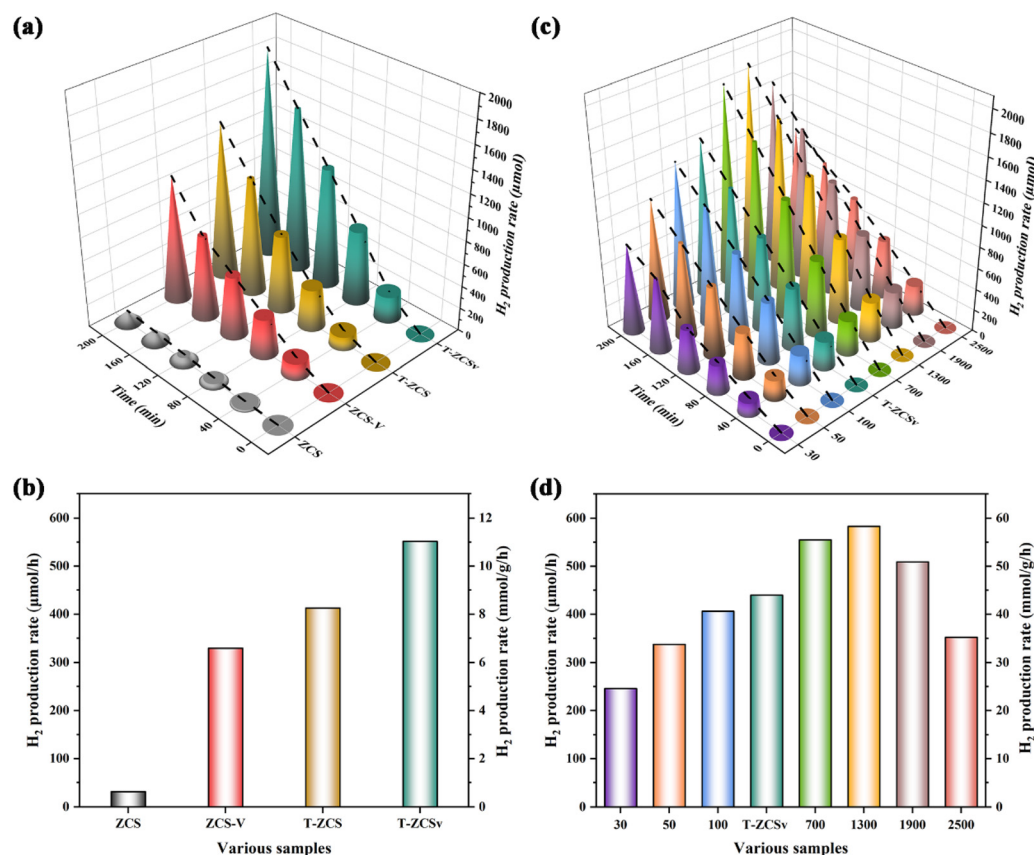


Fig. 9 Photocatalytic hydrogen production performance for various samples: (a and b) ZCS, ZCS-V, T-ZCS and T-ZCSv; (c and d) a series of ZIF-8@T-ZCSv composites with different contents.

Table 2 The H₂ production rates of ZCS, ZCS-V, T-ZCS and T-ZCSv

| Samples | H ₂ production rate ^a (μmol h ⁻¹) | H ₂ production rate (mmol g ⁻¹ h ⁻¹) |
|---------|---|--|
| ZCS | 32.00 | 0.64 |
| ZCS-V | 329.44 | 6.58 |
| T-ZCS | 412.92 | 8.25 |
| T-ZCSv | 551.74 | 11.03 |

^a 50 mg ZCS, ZCS-V, T-ZCS and T-ZCSv were used in the photocatalytic production of hydrogen.

RPH performance is attributed to the effect of substrate ZIF-8 polyhedron contributing to better dispersion ability of T-ZCSv by avoiding agglomeration and therefore exposing more active sites.⁴¹ Nevertheless, due to the relatively low content of T-ZCSv, ZIF-8@T-ZCSv (30 mg), ZIF-8@T-ZCSv (50 mg), and ZIF-8@T-ZCSv (100 mg) display lower RPH performance, *i.e.*, 246.1 μmol h⁻¹ or 24.61 mmol g⁻¹ h⁻¹, 337.27 μmol h⁻¹ or 33.72 mmol g⁻¹ h⁻¹ and 406.48 μmol h⁻¹ or 40.64 mmol g⁻¹ h⁻¹ respectively, which are less than the RPH of T-ZCSv. The lower performance of these samples can be explained based on the shielding effect provided by the dominant ZIF-8 framework to the active sites of T-ZCSv, which causes poor light absorption by the T-ZCSv. Furthermore, the hydrogen pro-

duction efficiency of ZIF-8@T-ZCSv (1900 mg and 2500 mg) composites decreased with the mass of T-ZCSv, indicating the re-emergence of the agglomeration phenomenon of T-ZCSv nanoparticles due to their dominant content. T-ZCSv aggregates loose light-exposable active sites and therefore exhibits poorer absorption of visible light. The specific RPH of the samples are shown in Table 3. The measured values suggest that an appropriate amount of T-ZCSv solid solution is homo-

Table 3 The H₂ production rates for ZIF-8@T-ZCSv composites with different contents

| Samples | H ₂ production rate ^a (μmol h ⁻¹) | H ₂ production rate (mmol g ⁻¹ h ⁻¹) |
|------------------------|---|--|
| ZIF-8@T-ZCSv (30 mg) | 246.10 | 24.61 |
| ZIF-8@T-ZCSv (50 mg) | 337.27 | 33.72 |
| ZIF-8@T-ZCSv (100 mg) | 406.48 | 40.64 |
| T-ZCSv | 439.92 | 43.99 |
| ZIF-8@T-ZCSv (700 mg) | 554.98 | 55.49 |
| ZIF-8@T-ZCSv (1300 mg) | 583.25 | 58.32 |
| ZIF-8@T-ZCSv (1900 mg) | 508.87 | 50.88 |
| ZIF-8@T-ZCSv (2500 mg) | 352.20 | 35.22 |

^a 10 mg T-ZCSv and ZIF-8@T-ZCSv composites were used in the photocatalytic production of hydrogen.

geneously dispersed on the surface of ZIF-8 polyhedra, which is vital to improve photocatalytic efficiency.

To elucidate the electron transport mechanism across the interfacial homojunction for the enhanced RPH of ZIF-8@T-ZCSv composites, the Mott-Schottky curves of T-ZCSv were investigated to determine the conduction band (CB) and valence band (VB) alignments, including the band gap energy (E_g), respectively. The E_g of T-ZCSv containing sulfur vacancies and twin crystal structure with WZ/ZB homogeneous junction is measured as 2.60 eV using the Tauc plot with Kubelka-Munk function as shown in Fig. 10a. Additionally, the Mott-Schottky plots were used to determine the nature of the semiconducting samples and the flat band potential (E_{fb}) of T-ZCSv in 0.1 M Na_2SO_4 electrolyte (frequency used is 1 K Hz). The positive slope of the linear portion in Fig. 10b indicates that T-ZCSv is an n-type semiconductor. The Fermi energy level (E_{fb}) of n-type semiconductors is close to the CB potentials (E_{CB}),^{42–44} so the calculated E_{CB} of T-ZCSv is -0.6 V (vs. NHE). Combining the E_g and E_{CB} of T-ZCSv, the corresponding VB potentials (E_{VB}) of T-ZCSv are estimated as 2.0 V (vs. NHE) with reference to the equation; $E_{VB} = E_g + E_{CB}$, which agrees with the previous report.¹³ Based on the energy band structure measurements, an appropriate mechanism for the improved photocatalytic evolution of hydrogen was proposed in Fig. 10c. The E_{CB} and E_{VB} values of the ZB region are slightly lower (more negative) than those of the WZ region near the interface

of the ZB and WZ portions in the twin-crystalline T-ZCSv solid-solution. Consequently, a new distinctive homogeneous junction like the S scheme heterogeneous junction is formed at the interface. Through *in situ* XPS spectroscopy measurements of the T-ZCSv sample before and after illumination (Fig. S4†), it was observed that the binding energy of Cd 3d shifted towards higher values, while the binding energy of S 2p shifted towards lower values. This observation suggests that Cd acts as an electron donor in the structure, while S acts as an electron acceptor. Upon illumination, S receives electrons from Cd, resulting in the creation of an electric field in the interfacial region. This electric field causes significant bending of the band edges, particularly in the interior or the region near the heterojunction, where the gradient of band bending is more pronounced. Hence, the ISIXPS results successfully elucidate the electron transfer pathway in the T-ZCSv S-scheme heterojunction. The valence band electrons (bound electrons) are transferred to the CB in T-ZCSv upon the visible light illumination, and the corresponding holes have remained in its VB. The photoinduced holes in the VB of the WZ segment and photo-induced electrons in the CB of the ZB segment are inclined to recombine at the interface due to the coulombic attraction between the band structures of WZ and ZB at the WZ/ZB homogeneous junction. Thus, the isolation and transportation of photo charges is achieved by the electric field developed across the homojunction, while the useless electrons and

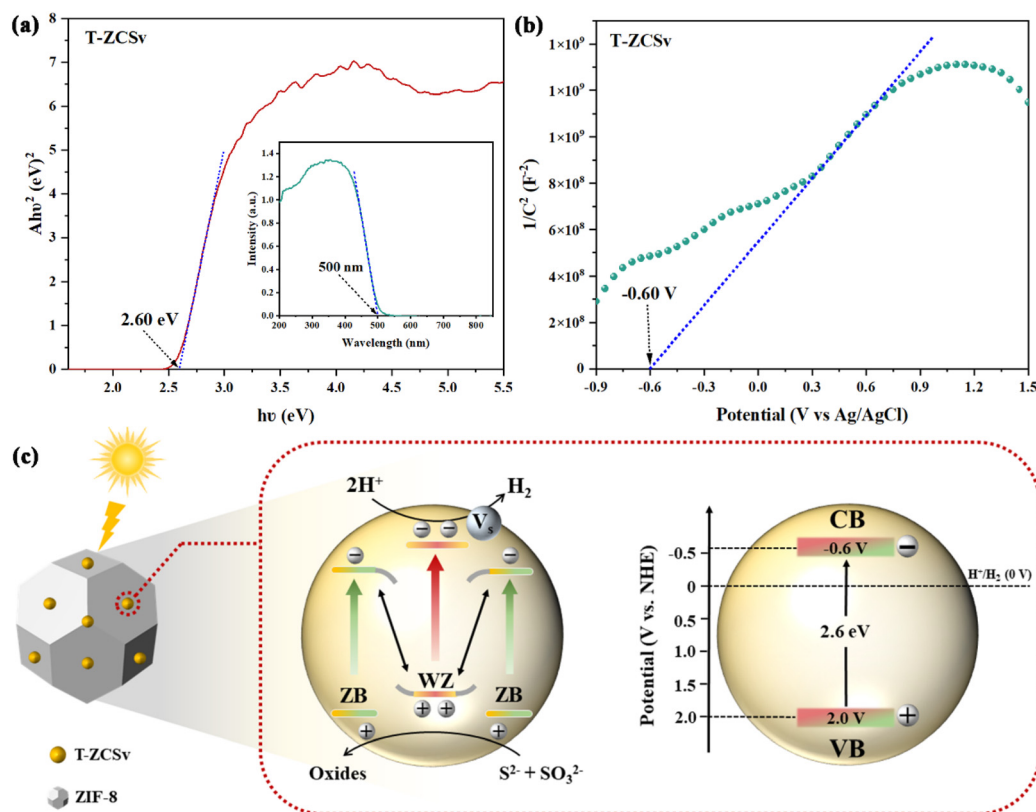
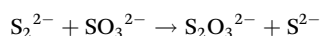
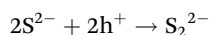
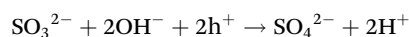
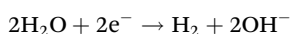
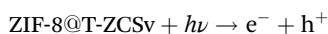


Fig. 10 (a) Band gap energy and UV-vis diffuse reflection spectra (inset) of T-ZCSv; (b) Mott-Schottky curves of T-ZCSv; (c) mechanism for enhanced photocatalytic performance for ZIF-8@T-ZCSv.

holes are removed by recombination and the powerful photo-generated electrons in the CB of WZ were retained to participate in the photocatalytic hydrogen production.^{16,45}

On the other hand, S vacancies trap free electrons and capture the abundant electrons on the CB of T-ZCSv, which not only is beneficial for prolonging the electron lifetime, but also promotes the transportation of photoinduced electrons from bulk to the surface of T-ZCSv for a reduction reaction with H^+ ions of water to generate H_2 (0 V vs. NHE). Meanwhile, the corresponding excess holes in the VB of T-ZCSv are oxidized by sulfide sacrificial agents (S^{2-} and SO_3^{2-}), thus improving the HPR activities. In other words, the synergistic effect of the WZ/ZB twin boundaries and S vacancies contributed to the enhanced photocatalytic activity of T-ZCSv and the ZIF-8@T-ZCSv composites. The reaction process can be described by the following equations:



Furthermore, the unique 3D/0D ZIF-8@T-ZCSv spatial structure possesses an extraordinary feature. The structurally larger ZIF-8 with a higher specific surface area can be exploited as a substrate that provides space and facilitates the proper dispersion of T-ZCSv nanoparticles. This feature significantly reduces the aggregation of T-ZCSv and leads to more surface-active sites exposed for the photocatalytic reaction. In summary, the twinned WZ/ZB interfacial homogeneous junction and the rich surface S vacancies exist in T-ZCSv, and an appropriate amount of ZIF-8 polyhedron substrate [ZIF-8@T-ZCSv (1300 mg)] provides a synergetic effect leading to a remarkable improvement in the hydrogen production efficiency.

Conclusions

A series of different characteristics of $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ solid solutions with WZ/ZB homojunctions and rich-surface S vacancies has been respectively designed and synthesized by simple and practical methods. Synergy of the twinned superlattices and S vacancies of the samples could boost the isolation and transportation efficiency of photoinduced electron-hole pairs and extend the electron lifetime in the absence of any cocatalyst under visible-light irradiation. Furthermore, the introduction of appropriate ZIF-8 polyhedra with a large specific surface area can provide a suitable 3D space dispersion to avoid aggregation of T-ZCSv nanoparticles for exposing more reactive sites in the photocatalytic reaction. In synergy with the above advantages, ZIF-8@T-ZCSv (1300 mg) demonstrated that the optimized photocatalytic H_2 production activity reached

$583.25 \mu\text{mol h}^{-1}$. The sulfur vacancies can effectively trap electrons, the twinned structure homojunction can support the isolation of photogenerated carriers, and ZIF-8 can hinder the agglomeration phenomenon of T-ZCSv. The synergistic effect of these three factors is supposed to result in a dramatic increase in the hydrogen production performance for the ZIF-8@T-ZCSv composite. The research work not only provides a theoretical basis and experimental support to design high-performance $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ -based photocatalysts, but also provides a comparable reference scheme for $\text{Zn}_{0.5}\text{Cd}_{0.5}\text{S}$ -based photocatalysts with unique characteristics.

Author contributions

Yuhao Zhang: conceptualization, methodology, formal analysis, investigation, and writing (original draft). Dingze Lu: conceptualization, funding acquisition, resources, and writing (review and editing). Min Zhou and Kiran Kumar Kondamareddy: supervision, writing (review and editing), and resources. Zhennan Wang and Jing Li: validation, supervision, and writing (review and editing). Huiqing Fan: conceptualization, methodology, supervision, and writing (review and editing). Dezhong Cao: writing (review and editing). Wingkei Ho: conceptualization, visualization, supervision, data curation, and writing (review and editing).

Conflicts of interest

The authors confirm no conflicts of interest.

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