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Title	Photocatalytic hydrogen generation with simultaneous organic degradation by composite CdS-ZnS nanoparticles under visible light
Author(s)	Wang, X; Peng, WC; Li, XY
Citation	International Journal of Hydrogen Energy, 2014, v. 39 n. 25, p. 13454-13461
Issued Date	2014
URL	http://hdl.handle.net/10722/202671
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Submitted to:International Journal of Hydrogen Energy (HE-D-13-03747)Date:March 6, 2014

## Photocatalytic hydrogen generation with simultaneous organic degradation by composite CdS-ZnS nanoparticles under visible light

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#### 1

#### Abstract

2 A visible light-driven CdS-ZnS photocatalyst in the form of nanoparticles with a 3 heterogeneous structure was synthesized using the stepped microemulsion method. The 4 composite CdS-ZnS was capable of simultaneous photocatalytic hydrogen production and 5 organic degradation under visible light. The ZnS deposition on CdS helped suppress the 6 recombination of electron/hole pairs generated on the more reactive CdS, leading to faster 7 hydrogen production and improved stability of the CdS-ZnS in comparison to the bare CdS 8 catalyst. Deposition of Ru on the catalyst surface further increased its photo-reactivity by 9 about 4 times for hydrogen production. The heterostructured nanoparticles were effective in 10 photocatalytic hydrogen production together with the degradation of model organic 11 substances, including formic acid, methanol, and ethanol. The highest hydrogen production 12 rate was achieved by the (CdS-ZnS)/Ru catalyst at 266 mmol/m<sup>2</sup>-h in the formic acid 13 solution with an energy conversion efficiency of 3.05% in visible light, and the 14 corresponding organic degradation rate in terms of the removal of chemical oxygen demand 15 (COD) was estimated at 4272 mg COD/m<sup>2</sup>-h.

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Keywords: Hydrogen production, organic photolysis, CdS-ZnS, photocatalyst, visible light,
solar energy.

19

#### 20 Introduction

21 Hydrogen is one of the most promising clean and renewable energy carriers. It has a 22 high combustion value and a near-zero level of pollutant and greenhouse gas emissions. 23 Photocatalytic hydrogen generation from water is an attractive and environmentally-friendly 24 method to harvest the solar energy [1]. However, while visible light ( $\lambda$ >420 nm) covers a 25 large portion of the solar spectrum, most photocatalysts, such as TiO<sub>2</sub>, function only under the 26 energy-intensive UV light. Efforts have been made to develop novel photocatalysts, such as 27 metal oxides (e.g. ZnO) and metal sulfides (e.g. CdS), that response to both UV and visible 28 lights for water photolysis and hydrogen production [2, 3]. However, the solar energy 29 conversion efficiency of these photocatalysts for hydrogen generation is still rather low due to 30 mainly the recombination of photo-generated electron/hole pairs [4]. Moreover, an increase in 31 reactivity of the photocatalyst would often result in a decrease in stability of the catalyst, 32 leading to a rapid loss of its catalytic capability to photo-corrosion [5].

A well-structured co-catalyst that integrates the functions from two or more catalyst materials may offer solutions to the above problems. A highly photo-sensitive material with a narrow band gap, such as CdS, will provide a great photo-reactivity for  $H_2$  generation, while the use of a relatively less active material with a wider band gap, such as ZnS, can effectively 37 reduce the electron/hole recombination and thereby protect the more active catalyst during the photocatalytic process [6]. In addition to hydrogen evolution ( $H^+$  reduction), the 38 39 photocatalytic reactions in water also render a strong oxidation power that may be utilized for 40 wastewater treatment. In fact, photocatalytic oxidation has been developed as an advanced 41 oxidation technology for treatment and pre-treatment of various pollutants in wastewater [7]. 42 During the photocatalysis under solar light, model organic pollutants such as alcohols can 43 function as electron donors for hydrogen evolution, whilst the organics are degraded [8, 9]. In 44 such a photocatalytic application, both the purposes of hydrogen production and wastewater 45 treatment can be achieved using the solar energy [5, 7].

46 CdS and ZnS are known as photocatalysts owing to their high photo-sensitivities [10, 47 11]. In addition to a sole catalyst material, research has been carried out to integrate CdS and 48 ZnS or with other co-catalysts to increase the photo-reactivity of the catalysts [12-16]. 49 However, most of the material integration was achieved in the form of a homogeneous 50 system, e.g., a solid solution, which would only tailor the band gaps of the two catalyst 51 materials [12-16]. In the present study, a composite CdS-ZnS catalyst with a heterogeneous 52 structure was synthesized. The heterostructured catalyst was shown as a visible light-driven 53 photocatalyst with a much improved photo-reactivity and photo-stability. A number of model 54 organic substances were tested as electron donors for hydrogen production. The aim of the study was to achieve both photocatalytic hydrogen generation and organic wastewater 55 56 treatment under visible light.

57

#### 58 Materials and Methods

#### 59 Synthesis of the CdS-ZnS catalysts

60 The stepped microemulsion technique was used to synthesize CdS-ZnS photocatalyst

3

61 nanoparticles with a heterogeneous structure. The synthesis was conducted in a water/Triton 62 X-100/1-butanol/n-hexane system, with cadmium nitrate (Cd(NO<sub>3</sub>)<sub>2</sub>, 99%, Sigma-Aldrich) and zinc nitrate (Zn(NO<sub>3</sub>)<sub>2</sub>, 99%, Sigma-Aldrich) used as the Cd and Zn precursors, 63 respectively, and sodium sulfide (Na<sub>2</sub>S·xH<sub>2</sub>O, Sigma-Aldrich) as the S<sup>2-</sup> source. To form the 64 CdS nanoparticles, 40 ml of 0.1 M Cd(NO<sub>3</sub>)<sub>2</sub> was placed in the microemulsion (W/O) 65 66 solution and 50 ml of 0.1 M Na<sub>2</sub>S was added dropwise under a vigorous stirring condition. The mixture was stirred for 15 min and 10 ml 0.1 M Zn(NO<sub>3</sub>)<sub>2</sub> was then added to react with 67 excess S<sup>2-</sup> to form ZnS that deposited on CdS particles. The mixture was stirred for 6 h at 68 69 room temperature. A Cd to Zn precursor ratio of 0.8 to 0.2 was used for the microemulsion 70 process to form  $(CdS)_{0.8}/(ZnS)_{0.2}$ . The CdS-ZnS solid precipitates formed in the solution were 71 centrifuged and washed with DI water and alcohol. For comparison, a homogeneous solid 72 solution of Cd<sub>x</sub>Zn<sub>1-x</sub>S with a Cd to Zn ratio of 0.8:0.2 was also prepared using the one-step microemulsion method. A pre-determined amount of 0.1 M Cd(NO<sub>3</sub>)<sub>2</sub> and a pre-determined 73 amount of 0.1 M Zn(NO<sub>3</sub>)<sub>2</sub> were mixed together and placed in the microemulsion (W/O) 74 75 solution. An excessive amount of 0.1 M Na<sub>2</sub>S was added drop-wise to the W/O liquor to form 76 the precipitates of Cd<sub>0.8</sub>Zn<sub>0.2</sub>S under a vigorous stirring condition. Similarly, the solid 77 particles were recovered and washed. The dry catalyst powder obtained from each batch was annealed at 723 K in a furnace (LHT 02/16 LBR, Nabertherm) supplied with pure nitrogen 78 79 for 2 hr. Afterward, the powder was grounded by ball mill for 5 min and then stored in dark 80 before use.

81 Ruthenium (Ru) was deposited on the surface of the catalyst using *in-situ* photo-82 deposition in an acetic acid solution of RuCl<sub>3</sub> (Aldrich). Photo-deposition was carried out by 83 illuminating ( $\lambda$ >420 nm, 300 W Xe lamp) the CdS-ZnS catalyst particles suspended in the 84 RuCl<sub>3</sub> solution for 20 min. The load of Ru coating was around 5% of the CdS-ZnS content, 85 and the resulting composite catalyst was denoted as (CdS-ZnS)/Ru. 86

#### 87 Characterization of the photocatalysts

88 The size distribution of the catalyst particles was measured by a laser diffraction particle size analyzer (Delsa<sup>™</sup> Nano, Beckman Coulter). The BET surface area of the photocatalyst 89 90 was determined by a surface area analyzer (SA3100, Beckman Coulter). The diffuse 91 reflection spectrum (DRS) of the photocatalyst was obtained using a UV-vis 92 spectrophotometer (Lambda 25, Perkin Elmer) that was converted from the reflection 93 function to the absorbance function following the Kubelka-Munk method [17]. The 94 crystalline phases and structural features of the catalysts were analyzed by an X-ray 95 diffraction (XRD) system (D8 Advance, Burker AXS) with the Cu Ka irradiation from 10 to 96 90 degrees. The morphology of the catalyst particles was examined and their selected area 97 electron diffraction (SAED) pattern was obtained using a transmission electron microscope 98 (TEM) (Tecnai G2 20 S-TWIN, Philips FEI). In addition, the TEM equipped with an energy-99 dispersion spectroscopy (EDS) was employed to obtain the element mapping distribution for 100 the photocatalyst.

101

#### 102 Photocatalytic H<sub>2</sub> production in different model organic solutions under visible light

103 The photocatalytic hydrogen production experiments were conducted in a circular photo 104 cell made of Pyrex class with a quartz window on the top. A 300 W Xe lamp (wavelength 105 250-750 nm) was used in a light source setup (PLS-SXE, Trustech) to simulate the solar light. 106 A cutoff ( $\lambda < 420$  nm) filter was installed to yield only visible light (light intensity ~ 70 107 mW/cm<sup>2</sup> measured by a light power meter, I400, Trustech) over a lighting area of 33 cm<sup>2</sup> 108 from the top of the photo cell. A certain amount of the photocatalyst, i.e. 0.15 g of the CdS-109 ZnS or (CdS-ZnS)/Ru powder, was suspended in 150 mL of water or an organic solution. 110 Different model organic compounds, including formic acid, methanol and ethanol, were 111 tested as electron donors for photocatalytic hydrogen production. The solution had an organic 112 content of 10% and was kept at pH~7. The gas produced during the photo-tests was collected, 113 and the H<sub>2</sub> and CO<sub>2</sub> contents were analyzed by a gas chromatograph (GC HP5890 Series II, 114 Hewlett Packard). Each run of the photo-test lasted for around 4 hrs. The H<sub>2</sub> production rate 115 for a test was calculated from the slope of the linear regression of the accumulated hydrogen 116 production vs. time. Each test was repeated at least once, and the average results are reported 117 for the duplicate. The reactivity of the photocatalyst was evaluated in terms of the specific 118 hydrogen production rate (R) and the energy conversion efficiency ( $\eta$ ), as described by the 119 following equations:

120 
$$R_A = \frac{\Delta m_{H_2}}{A\Delta t}, \text{ or}$$
(1)

121 
$$R_A = \frac{\Delta m_{H_2}}{W\Delta t}, \text{ and}$$
(2)

122 
$$\eta = \frac{R \,\Delta H_c}{I} \tag{3}$$

where  $R_A$  and  $R_w$  are the area-based and weight-based specific hydrogen production rates, respectively,  $\Delta m_{H2}$  is the moles of H<sub>2</sub> production measured,  $\Delta t$  the duration of the photoreaction, *A* the irradiation area (33 cm<sup>2</sup>), *W* is the amount (weight) of the catalyst in the photo cell,  $\Delta H_c$  is the combustion value of H<sub>2</sub> (286 kJ/mol) and *I* the light density. The hydrogen production rates reported below were normally obtained from the first 2 or 3 test runs for the newly prepared photocatalysts.

129 The quantum yield (QY) of the photocatalytic reaction was also calculated as follows, 130 which has been used to evaluate the reactivity of a photocatalyst for  $H_2$  production under light 131 irradiation over a broad band of wavelength, 132

$$QY = \frac{m_{re}}{m_{ip}} = \frac{2m_{H_2}}{m_{ip}}$$
(4)

where  $m_{re}$  is the amount of reacted electrons which is two times of the moles of H<sub>2</sub> produced ( $m_{H2}$ ), and  $m_{ip}$  is the amount of total incident photons for the catalyst in the photo cell. The amount of incident photons for the setup of the photocatalytic tests was determined by the method of ferrioxalate actinometer employing potassium ferrioxalate (K<sub>3</sub>[Fe(C<sub>2</sub>O<sub>4</sub>)<sub>3</sub>]) [18].

To evaluate the reproducibility of the experiment and the stability of the catalyst, the photocatalytic hydrogen generation test was repeated for 10 times for each type of the catalysts. In addition, the amount of  $Cd^{2+}$  leaching into the solution during the photocatalytic experiments was measured by an atomic absorption spectrometer (AAnalyst 300, Perkin Elmer).

142

#### 143 **Results and Discussion**

#### 144 Characterization of the photocatalysts

The CdS-ZnS catalysts are nanoparticles that ranged from 40 to 340 nm with a number-145 146 based mean size of around 100 nm according to the particle size analysis. The TEM images 147 show that the catalysts had primary particles of about 60 nm (Figure 1). Moreover, rather than 148 a homogeneous solid solution, the CdS-ZnS catalysts synthesized by the two-step 149 microemulsion process appeared to be of a heterogeneous structure with ZnS deposited on the 150 surface of the CdS crystals (Figure 1A-C). According to the TEM-EDS mapping result, the 151 composite catalysts were Cd-based nanoparticles with Zn distributing mostly on the particle 152 surface (Figure 1D-F). Based on the SAED patterns given in Figure 2, the comparison 153 between pure CdS and CdS-ZnS confirmed the presence of cubic phase ZnS (d-Spacing = 154 0.320 nm (111), 0.195 nm (220), 0.165 nm (222)) in the composite catalyst [19].

155 A good crystal phase of CdS can be found in the CdS-ZnS catalyst according to its XRD 156 pattern (Figure 3). More specifically, the crystal structure of CdS-ZnS is dominated by the 157 hexagonal CdS phase. Due to likely the low ZnS content in the CdS-ZnS particles, the peak 158 of ZnS could not be clearly identified in the XRD spectra. A similar result has been previously reported for the CdS-ZnS catalyst by Soltani et al. [20]. For the solid solution of 159 160  $Cd_{0.8}Zn_{0.2}S$ , the diffraction peaks were found to shift to higher angles in comparison to CdS, 161 which is consistent with previous findings [21]. In contrast, the heterostructured CdS-ZnS did 162 not show such a shift of the diffraction peaks from those of CdS.

The diffuse reflection spectra of CdS-ZnS and its base materials (CdS and ZnS) are presented in Figure 4 to show the sensitivity of the photocatalysts to visible light. As expected, ZnS did not response to visible light, while CdS had an intensive absorption band in the visible region with a band gap of 2.23 eV. CdS-ZnS also responded well to visible light with a band gap of 2.31 eV, although it blue shifted slightly, which is consistent with the results of Soltani et al. [20]. The similar refection spectra between CdS and CdS-ZnS indicate that the photo-reactivity of CdS was well remained in the composite CdS-ZnS catalyst.

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#### 171 Photocatalytic hydrogen production by the CdS-ZnS catalyst under visible light

The CdS-ZnS nanoparticles were capable of photocatalytic hydrogen generation together with organic degradation under visible light (Figure 5A). Bare CdS had a hydrogen production rate of only  $13.7\pm1.2 \mu$ mol/h in the formic acid solution, while ZnS was not able to produce hydrogen under visible light. In comparison, CdS-ZnS was shown as a much effective visible light-driven photocatalyst that could produce hydrogen at a rate of  $189.5\pm3.5$ µmol/h. Coating of ZnS on the CdS surface formed a heterogeneous nanoparticle structure, resulting in a significant increase of the hydrogen production activity of the photocatalyst. In comparison, simply incorporating Zn into the CdS structure was not effective to enhance the photo-activity of the catalyst. The H<sub>2</sub> production rate by the structured CdS-ZnS ((CdS)<sub>0.8</sub>/(ZnS)<sub>0.2</sub>) was found to be about 10 times of that by the Cd<sub>0.8</sub>Zn<sub>0.2</sub>S solid solution (180 vs. 18 mmol/m<sup>2</sup>-h) in the S<sup>2-</sup>/SO<sub>3</sub><sup>2-</sup> solution under the same photocatalytic conditions.

183 The composite CdS-ZnS heterostructure apparently brought about a synergetic function 184 of the catalyst materials. The use of the more photo-sensitive CdS ensured the reactivity of 185 the photocatalyst [3], while the ZnS functioned to suppress the recombination of electron/hole 186 pairs formed on CdS, making the electros more available for H<sup>-</sup> reduction and H<sub>2</sub> evolution [8, 9]. Moreover, the use of the outer ZnS could help increase the stability of the CdS-based 187 188 photocatalyst. After 10 runs of the photocatalytic tests, the CdS-ZnS catalyst still maintained 189 86% of its hydrogen productivity at a level of 166.0 µmol/h, whilst the bare CdS had only 46% 190 of its reactivity remaining for hydrogen production at a low rate of 6.3 µmol/h (Figure 5B). The ZnS coating could also minimize the leaching of  $Cd^{2+}$  from the catalyst nanoparticles 191 192 during the photocatalytic process. Between test runs 2 and 5, the CdS-ZnS catalyst had an average  $Cd^{2+}$  leaching rate of 68.5  $\mu$ g/h, while the pure CdS particles had a high  $Cd^{2+}$ 193 leaching rate of 334.6  $\mu$ g/h. Between runs 6 and 10, the leaching of Cd<sup>2+</sup> from the CdS-ZnS 194 195 catalyst decreased to a much lower level at 9.5 µg/h, in comparison to bare CdS that had a Cd<sup>2+</sup> leaching rate of 103.2 µg/h. Coating of ZnS could effectively protect the more reactive 196 197 CdS catalyst and hence greatly improved its stability against photo-corrosion.

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# Photocatalytic hydrogen generation together with organic degradation by the CdS-ZnS catalysts

201 Deposition of ruthenium on the CdS-ZnS surface further enhanced the hydrogen 202 productivity of the photocatalyst under visible light (Figure 6). With the Ru deposition on the 203 catalyst surface, the photocatalytic hydrogen production rate increased about 4 times. For 204 both CdS-ZnS and (CdS-ZnS)/Ru in pure water, no hydrogen was produced in the absence of 205 the model organics. The presence of organic enabled the catalyst to effect photocatalytic  $H_2$ 206 generation, as the model organics functioned as electron donors for the reduction of  $H^+$  ions 207 to realize hydrogen evolution.

208 The highest H<sub>2</sub> production rate was achieved with the (CdS-ZnS)/Ru catalyst at 266±4 mmol/m<sup>2</sup>-h in formic acid with an overall energy conversion efficiency of  $3.05\pm0.05$  % and a 209 210 quantum yield of around 20%. In water pollution control, the chemical oxygen demand 211 (COD) is commonly used to measure the concentration of organic matter in water. In theory, 212 COD is the amount of oxygen that would be consumed for organic oxidation. During the 213 organic oxidation, electrons donate electrons that are accepted by oxygen, and every four 214 moles of electrons donated by the organic matter correspond to one mole of oxygen 215 consumed. During the photocatalytic process, oxidation (organic oxidation) and reduction (H<sup>+</sup> 216 reduction) also occur simultaneously. The amount of electrons provided by the organic would 217 be equal to the amount of electrons received by  $H^+$  for  $H_2$  evolution. Thus, theoretically, the 218 amount of organic degradation in terms of COD removal can be estimated as follows in 219 relation to the amount of H<sub>2</sub> production,

$$220 \qquad COD \quad removal = \frac{m_{H_2} M_{O_2}}{2} \tag{5}$$

where  $M_{O_2}$  is the molar mass of  $O_2$  (32 g/mol). The theoretical photocatalytic COD reduction rate for formic acid was 14.1 mg/h by the (CdS-ZnS)/Ru catalyst (Figure 7A), corresponding to an area-based specific rate of 4272 mg COD/m<sup>2</sup>-h under visible light. Meanwhile, CO<sub>2</sub> was produced at a rate of 226 mmol/m<sup>2</sup>-h in the gaseous phase during the photocatalytic test, which agrees well with the theoretical COD reduction rate. The molar ratio of hydrogen to carbon dioxide productions was approximately 1:1, suggesting a complete decomposition of 227 formic acid [22].

However,  $CO_2$  production was not detected during the photocatalytic H<sub>2</sub> generation in both methanol and ethanol solutions. It is likely that the photocatalytic reactions resulted in organic degradation and intermediate formation other than complete organic mineralization [8]. When the CdS-ZnS catalyst was excited by visible light, the photo-generated holes would attack methanol or ethanol, leading to the following organic degradations [8, 23],

233 
$$CH_3OH + h^+ \longrightarrow HCHO + H^+;$$

$$CH_3CH_2OH + h^+ \longrightarrow CH_3CHO + H^+.$$

235 The photocatalytic hydrogen production rates increased as the model organic 236 concentration in solution increased (Figure 5B). For formic acid in the concentration range of 0-10%, the H<sub>2</sub> production rate by CdS-ZnS increased linearly (r = 0.98) with the formic acid 237 238 concentration. Beyond the range (>10%), the  $H_2$  production rate did not show a significant 239 increase with the formic acid concentration. For both methanol and ethanol, the 240 photocatalytic hydrogen production rate also increased nearly linearly with the organic 241 concentration. As the initial concentration decreased to a low level of 500 mg/L, hydrogen still could be produced from methanol and ethanol at  $12.3\pm2.5$  and  $9.6\pm2.0$  mmol/m<sup>2</sup>-h, 242 243 respectively. The specific hydrogen production rates based on the irradiation area or the 244 amount of catalyst were summarized in Figure 6. The hydrogen production rates in the methanol and ethanol solutions were  $220.9\pm5.2$  mmol/m<sup>2</sup>-h and  $122.1\pm3.0$  mmol/m<sup>2</sup>-h, 245 246 respectively, which are more than 2.5 times higher than that reported by Zhang and Zhang [7] 247 for the Ru/CdS/Al-HMS catalyst under similar conditions (350 W Xe lamp). The quantum efficiency of (CdS-ZnS)/Ru (20%) is more than an order of magnitude higher than that of 248 249 Ru/CdS/Al-HMS (1.2%) [7]. Hydrogen production in 10% methanol achieved a production 250 rate of 4.8±0.3 mmol/g-h for visible light irradiation, which is ten times of the specific 251 hydrogen production rate reported for the CuO/Al<sub>2</sub>O<sub>3</sub>/TiO<sub>2</sub> catalyst in 10% methanol under 252 solar irradiation [24]. The higher hydrogen generation and organic degradation efficiency 253 suggests the advantage of the heterostructure of the photocatalyst. Although ZnS cannot be 254 excited directly by visible irradiation, the photo-sensitive CdS can be readily excited by visible light. With its excitation, CdS would function as a photo-sensitizer to induce the 255 256 excitation of ZnS. The more negative conduction band (-1.4 V) of ZnS allows photo-excited 257 electrons to drop to the conduction band of CdS (-0.3 V) [24] and the electron flow from ZnS 258 to CdS supplies more electrons to the conduction band of CdS for transfer to the aqueous phase for H<sup>+</sup> reduction. Meanwhile, the holes left at the valance band of ZnS would attract 259 260 electrons from the chemical solution, resulting in organic oxidation and degradation.

261

#### 262 Conclusions

263 The composite CdS-ZnS nanoparticles with a heterogeneous structure were synthesized 264 as a visible light-driven catalyst capable of both photocatalytic hydrogen production and 265 organic degradation. The ZnS deposited on CdS would suppress the recombination of 266 electron/hole pairs formed on CdS, leading to a faster hydrogen generation rate in comparison 267 to bare CdS. The ZnS coating also helped protect the more sensitive CdS and hence greatly improve its stability against photo-corrosion. The presence of model organic substances, 268 269 including formic acid, methanol and ethanol, enabled photocatalytic hydrogen production 270 under visible light. The highest specific hydrogen production rate was achieved by the (CdS-ZnS)/Ru catalyst at  $266\pm4$  mmol/m<sup>2</sup>-h in the formic acid solution with a photo energy 271 conversion efficiency of 3.05±0.05%. In relation to the hydrogen production, the 272 corresponding photocatalytic organic degradation rate was  $4272\pm67 \text{ mg COD/m}^2$ -h. 273

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#### 275 Acknowledgements

This research was supported by grant HKU714112E from the Research Grants Council (RGC) of the Government of Hong Kong SAR and project 51308230 from the National Natural Science Foundation of China (NSFC). The technical assistance of Mr. Keith C.H. Wong is highly appreciated.

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#### 344 **Figure captions:**

345 Figure 1. TEM examination and EDS element mapping result of the CdS-ZnS catalyst: (A)-

- 346 (C) TEM images showing the heterostructure of the catalyst particles, (D) TEM
  347 image for the EDS mapping of (E) Cd distribution and (F) Zn distribution.
- Figure 2. The SAED pattern of (A) pure CdS and (B) the structured CdS-ZnS.
- Figure 3. XRD spectra of the catalyst materials: the structured CdS-ZnS composite and pureCdS and ZnS.
- Figure 4. Diffuse reflection spectra of the photocatalysts: the structured CdS-ZnS composite
   in comparison with pure CdS and ZnS.
- Figure 5. (A) Hydrogen production in the formic acid solution (10%) by the composite CdS-ZnS, bare CdS and bare ZnS, under visible light; and (B) stability of CdS-ZnS and bare CdS in terms of the hydrogen production rate and leaching of Cd<sup>2+</sup> from the catalysts during the repeated photocatalytic tests (at least 4 hr for each test cycle).
- Figure 6. (A) The area-based and (B) weight-based specific hydrogen production rates of the photocatalysts of CdS-ZnS and (CdS-ZnS)/Ru in different model organic solutions under visible light.
- Figure 7. (A) The theoretical COD removal rate by the CdS-ZnS and (CdS-ZnS)/Ru photocatalysts for different organic solutions under visible light; and (B) the effect of the initial organic concentration on the rate of photocatalytic hydrogen production by CdS-ZnS under visible light.
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