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Zi-Rong Tang, Yi-Jun Xu *et al.* Cocatalyst-modified In ${}_2\text{S}_3$ photocatalysts for C–N coupling of amines integrated with ${\sf H}_{_2}$ evolution

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Photocatalytic hydrogen (H₂) production coupled with selective oxidation of organic compounds into high-value-added organic intermediates has expansive prospects in the utilization and transformation of solar energy, which meets the development requirements of green chemistry. In this work, high-efficiency hole cocatalyst PdS-decorated In₂S₃ flower-like microspheres are fabricated for the effective visible-lightdriven C-N coupling of amines to imines coupled with H₂ evolution. Owing to the establishment of the internal electric field, which further boosts the transfer of photoexcited holes to PdS, PdS–In₂S₃ exhibits distinctly enhanced photocatalytic redox performance, which is 39.8 times higher for H_2 and 14.3 times higher for N-benzylidenebenzylamine than that of the blank In_2S_3 , along with high selectivity and stability. Furthermore, the practicability of dehydrogenation coupling of various aromatic amines to the corresponding C–N coupling products on PdS–In $_2S_3$ has been demonstrated and a plausible reaction mechanism has been proposed. This work is anticipated to stimulate further interest in establishing an innovative photoredox platform for selective organic synthesis coupled with $H₂$ evolution in a green and

1 Introduction

Combining photocatalytic hydrogen $(H₂)$ production with organic transformation to generate appreciating compounds in a sustainable and green way, which eschews the consumption of non-renewable fossil fuels and extreme or high-energy-consuming reaction conditions, has garnered considerable attention in environmentally friendly and economically sustainable industrial production.¹ Imines, as one of the most important intermediates for the synthesis of pesticides, pharmaceuticals, and biologically active molecules, have inspired ongoing efforts from both academic and industrial scientists.^{2,3} However, acidic (or alkaline) and hightemperature reaction conditions are usually needed to drive and catalyze the traditional imine synthesis reaction, with these reactions suffering from unsatisfactory yields and product selectivity.4–⁶ Recently, the photocatalytic oxidative dehydrogenation coupling of amines in a pure oxygen (O_2) atmosphere was reported, which possesses milder and more economical conditions than conventional synthesis methods.

sustainable way.

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However, it first requires the formation of superoxide radicals by electron attack on $O₂$ molecules to carry out amine oxidation, which not only loses the energy of the photogenerated electrons, but also exacerbates the uncertainty of the reaction, such as through excessive oxidation and side reactions, thereby reducing the selectivity of the reaction.⁷⁻¹¹ In contrast, the coproduction of H_2 and value-added imines in one photoredox cycle under anaerobic conditions is a prospective scheme to fully utilize the electrons and holes generated by light for cooperative reactions to simultaneously produce valuable organic chemicals and clean fuel.¹² PAPER

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Coupling of amines integrated with H₂ evolution

Yu Chen, Chang Tang ^{Lo} and Yi-Jun Jing-Yu Li, Ming-Y

> Indium sulfide (In_2S_3) , as a typical III-VI group metal sulfide¹³⁻¹⁶ with an excellent visible light response,¹⁷ low toxicity, 18 stable chemistry, 19 and a suitable band edge position,²⁰ has been used as photocatalyst for contaminant degradation,²¹ CO₂ reduction,²² organic coupling,^{23,24} and H₂ evolution.²⁵ However, the photocatalytic redox efficiency of blank In_2S_3 is severely hindered by the absence of catalytically active sites and the rapid recombination of photogenerated charge carriers.26,27 PdS has been proven as an oxidation cocatalyst with excellent hole-trapping ability, which can effectively capture and retain holes from other photoexcited semiconductors.²⁸⁻³² Therefore, the modification of PdS as a cocatalyst onto In_2S_3 for judiciously constructing PdS-In₂S₃ composites could be an appealing strategy.³³⁻³⁵ Such strategy

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Cocatalyst-modified $In₂S₃$ photocatalysts for C-N coupling of amines integrated with H_2 evolution \dagger

could not only effectively reduce the recombination of photoexcited carriers for maximizing the utilization of photogenerated electron–hole pairs, but also change the activation energy of the interfacial photoredox reaction,³⁶ further enhancing the photocatalytic performance of the catalyst.

Herein, we successfully constructed a PdS–In₂S₃ binary photocatalyst through a combination of solvothermal treatment and in situ chemical deposition for the dualfunctional photocatalytic redox reaction of dehydrocoupling of amines to imines and H_2 . The conversion of benzylamine (BA) reached more than 95.5% over the optimal PdS–In₂S₃ photocatalyst after 4 h of reaction, which is roughly 36.1-fold higher than that of blank In_2S_3 , along with a terrific N-benzylidenebenzylamine (N-BDBA) selectivity of above 99.0%. The formation of the internal electric field provides a powerful impetus for PdS to extract holes from $In₂S₃$, and thus effectively reduces the photogenerated carrier recombination, which leads to more photogenerated holes and electrons for the coproduction of imines and H_2 . In addition, such a photocatalytic system is applicable to various amines, furnishing the corresponding imines with excellent selectivity and long-term stability. Mechanism studies revealed that the carbon radical $(\text{Ph}(\cdot \text{CH})\text{NH}_2)$ plays a key role in the reaction, and a plausible reaction mechanism has been suggested accordingly. It is anticipated that this work will provide insights into the rational design of In_2S_3 based composite photocatalysts to facilitate the visible-lightdriven coproduction of high-value-added chemicals and clean $H₂$ fuel.

2 Results and discussion

As shown in Fig. 1a, a series of PdS-modified In_2S_3 (PdS- In_2S_3 nanospheres with different mass ratios were successfully synthesized by the combination of the

Fig. 1 (a) Schematic presentation of the synthesis of PdS–In₂S₃ samples. SEM images of (b) blank In_2S_3 and (c) 3% -PdS- In_2S_3 . (d) HRTEM image of 3% -PdS-In₂S₃. (e) Element mapping results of 3% -PdS-In₂S₃.

solvothermal method and the in situ chemical deposition method. The specific morphology information of the composites was confirmed by scanning electron microscopy (SEM) and transmission electron microscopy (TEM). Fig. 1b and S1 \dagger present the SEM images of the blank In_2S_3 nanospheres with diameters ranging from 2.3 to 7.7 μm, where several thin nanosheets can be observed at the edges. The morphology of In_2S_3 is not influenced by the deposition of PdS, which can be seen in the SEM image of PdS–In₂S₃ (Fig. 1c). From the high-resolution TEM (HRTEM) image in Fig. 1d, the obvious lattice fringes of 0.277 nm and 0.618 nm are scanned in PdS–In₂S₃, corresponding to the (400) and (111) planes of In_2S_3 , respectively.^{15,23,25} Meanwhile, the lattice stripes of 0.324 nm and 0.185 nm belonging to the (200) and (222) planes of PdS, respectively, are also observed in the binary samples.30,37 Additionally, the results of aberration-corrected high-angle annular dark-field scanning TEM (HAADF-STEM), elemental mapping (Fig. 1e), and energy-dispersive X-ray (EDX) spectroscopy (Fig. S2†) analyses suggest a homogeneous distribution of Pd, S, and In in the samples, demonstrating that PdS is evenly decorated onto the In_2S_3 nanospheres. In addition, the results of inductively coupled plasma-mass emission spectroscopy (ICP-MS) analysis demonstrate that the actual content of PdS in the different samples is close to the theoretical value (Table S1†). **Paper**

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Fig. 2 (a) XPS survey spectra of blank In_2S_3 and 3%-PdS-In₂S₃. The high-resolution XPS spectra of (b) S 2p, (c) In 3d, and (d) Pd 3d of 3%- PdS-In₂S₃ and In₂S₃ composites. (e) XRD patterns and (f) DRS results for In_2S_3 and the PdS-In₂S₃ samples with varying amounts of PdS.

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In order to study the valence states and element compositions of the composites, X-ray photoelectron spectroscopy (XPS) analysis was carried out. All elements (Pd, In, and S) related to In_2S_3 and PdS can be detected in the survey spectra (Fig. 2a), which are well-matched the with EDX and element mapping results. The peaks at 161.60 eV and 162.77 eV in the high-resolution S 2p XPS spectrum in Fig. 2b are categorized as S $2p_{3/2}$ and S $2p_{1/2}$, respectively, which are attributed to S^{2-25} The binding energies of In 3d in Fig. 2c demonstrate that the spin–orbit doublet peaks at 444.95 eV and 452.49 eV are attributed to In $3d_{5/2}$ and In $3d_{3/2}$, respectively, which match well with In^{3+} in $In_2S_3.^{26}$ As for Pd 3d in 3%-PdS-In₂S₃, the peaks at 342.16 eV and 336.93 eV belong to the binding energies of Pd $3d_{3/2}$ and Pd $3d_{5/2}$ (Fig. 2d), which are associated with the Pd^{2+} of PdS.^{37,38} Notably, with the modification of PdS, the characteristic elemental peaks associated with $In₂S₃$ shift toward lower binding energies, suggesting that electron migration between In_2S_3 and PdS affects the surface electron density of the photocatalysts.39–⁴¹ X-ray diffraction (XRD) analysis was

performed to characterize the properties of the crystalline phase with the different samples. The XRD patterns of the In_2S_3 and PdS– In_2S_3 containing different amounts of PdS are shown in Fig. 2e, and the main characteristic diffraction peaks of pure In_2S_3 (JCPDS no. 65-0459) at 27.4°, 33.2°, and 47.7° are detected in all materials, demonstrating that the characteristic structure of $In₂S₃$ is not damaged by the deposition of $PdS¹⁸$ However, the corresponding characteristic diffraction peaks of PdS are not identified in the pattern, which is due to the low load weight of PdS ³⁶ Furthermore, UV/vis diffuse reflectance spectroscopy (DRS) was used to measure the optical absorption properties of the different composites. As portrayed in Fig. 2f, the pristine In_2S_3 exhibits favorable visible light absorption, which is further enhanced with the increased addition of PdS. Moreover, on the basis of Tauc plots of $(ahv)^2$ versus photo energy (hv) ,^{42,43} the band gap energy (E_{g}) of In₂S₃ was derived to be 2.18 eV (Fig. S3†).

The photocatalytic activity of the pristine In_2S_3 and the PdS–In₂S₃ photocatalysts was measured by a dual-function

catalyzed performance of bare In_2S_3 and x%-PdS- In_2S_3 (x = 0.2, 0.5, 1, 3, 5, and 7). (c) DRS spectrum of 3%-PdS- In_2S_3 and yields of the product over 3%-PdS-In₂S₃ under different monochromatic lights. (d) Time courses of N-BDBA and H₂ evolution over 3%-PdS-In₂S₃. (e) Photocatalytic recycling tests over 3%-PdS–In2S3. Reaction conditions: 5 mg of 3%-PdS–In2S3 photocatalyst, 0.1 mmol of BA, and 10 ml of acetonitrile (MeCN) were added to a quartz glass reactor, then the suspension was degassed under Ar for 0.5 h, and finally illuminated by visible light for 2 h.

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photoredox system involving $H₂$ evolution and selective oxidation of BA to N-benzylidenebenzylamine (N-BDBA) (Fig. 3a). Qualitative and quantitative analyses of the liquid products were performed with the gas chromatography-mass spectrometry (GC-MS) method (Fig. S4†). As plotted in Fig. 3b, after exposure to visible light ($\lambda > 420$ nm) for 2 h, the blank $In₂S₃$ showed poor catalytic activity for N-BDBA generation and H_2 production (2.68 µmol and 1.05 µmol, respectively). After being modified by the cocatalyst PdS, the performance of the $PdS-In_2S_3$ binary composites for producing H_2 and N-BDBA is significantly improved. The amount of H_2 and N-BDBA produced over blank In_2S_3 and $x\%$ -PdS-In₂S₃ exhibits a volcano-type curve with the increase of the PdS amount. Particularly, 3% -PdS-In₂S₃ shows the

highest H_2 and N-BDBA yields of 41.62 µmol and 38.46 µmol, which are ca. 39.8- and 14.3-fold higher than that of the blank In_2S_3 . Excessive introduction of PdS leads to reduced photoactivity, which is mainly due to the fact that excessive PdS loading may hinder the active site and the light absorption of In_2S_3 ^{29,44} Furthermore, the molar ratios of the N -BDBA and H_2 during the process were measured to be *ca*. 1.0, revealing that the dehydrogenation reaction proceeds in chemically stoichiometric conditions.¹ For investigating the connection of photo-absorption with catalytic performance, wavelength correlation experiments were conducted in which the light intensity is the same. As can be seen in Fig. 3c, the photocatalytic activity of the 3% -PdS-In₂S₃ composite decreases with the increase in wavelength, which explicitly

 a Reaction conditions: 5 mg of 3%-PdS-In₂S₃ photocatalysts and 0.1 mmol of reactants were added in 10 mL of acetonitrile (MeCN). The suspension was degassed under Ar for 0.5 h and then illuminated by visible light for 2 h at normal temperature and atmospheric pressure.
^b Conversion (%) = $(n_0 - n)/n_0 \times 100\%$. ^c Selectivity (%) is based on the GC-MS n_2)/(n_0 − n) × 100%. \bar{f} The electrons-to-holes ratio involved in the redox process was derived by the following equation of e[−]/h⁺ = 2n(H₂)/(n_0 − n), where n_0 and n are the amounts of the initial and the residual amines, respectively, n_1 and n_2 represent the amount of imine and by-products, respectively, and t represents the reaction time $(2 h)$.

demonstrates the dependence on visible light to drive the process.

Stability and recyclability are the essential criteria for evaluating the performance of photocatalysts. The reaction time curves (Fig. 3d and S5†) show that the conversion of benzylamine gradually increases with the extension of the reaction time, and about 95% of benzylamine can be converted within 4 h, which further illustrates the excellent photo-oxidative reduction efficiency of the composites. As demonstrated in Fig. 3e, after 6 cycles in 12 h, the amount of H_2 and N-BDBA generated by the 3%-PdS–In₂S₃ remained basically unchanged, and no obvious inactivation was observed. Based on these positive results, in order to validate the tolerability of our cooperative photoredox-catalyzed system, the substrate range for the amine dehydrocoupling reaction was extended. As can be seen from Table 1, the PdS– In_2S_3 photocatalysts exhibit high selectivity for different substrates, while the variation of substrates affects the yields of the corresponding imines. Specifically, the transformation of primary amines with electron-absorbing groups is

marginally more moderate than that of imines with electrondonating groups (entries 1–8), and the rate of reaction of the regional isomers increases in the order of ortho < meta < para, revealing a steric effect (entries 4–10). The abovementioned results indicate that both electronic and siteblocking effects significantly affect the reaction.⁴⁵ Noticeably, the catalysts also exhibit favorable activity for heteroatomcontaining amines (entries 11 and 12). In addition, no other by-products were observed by GC-MS analysis (Fig. S6–S15†). The ratio of electrons to holes $(R(e^{-}/h^{+}))$, REH) participating in the photoredox reactions is one of the key indexes defining the catalytic performance, and was calculated to be about 100% in the photo-oxidation reaction described above,¹ implying that the dehydrogenation reaction of the dualfunctional photocatalytic system is chemically stoichiometric, thus unveiling the effectiveness of cooperative photoredoxcatalyzed amine conversion and H_2 production on PdS–In₂S₃.

In an attempt to examine the causes of the enhanced photocatalytic performance of the 3% -PdS–In₂S₃ binary complex compared to the blank $In₂S₃$, a series of

Fig. 4 (a) Polarization curves. (b) Transient photocurrent spectra. (c) CV curves. (d) Decay curves of photovoltage (left) and electron lifetime (right). (e) EIS Nyquist plots. (f) TRPL spectra of In_2S_3 and 3%-PdS-In₂S₃, respectively. (g) CPD of PdS and In₂S₃. (h) The schematic diagrams of the proposed IEF formation and carrier transfer processes (1), (2), and (3).

photoluminescence (PL) measurements were taken, which are closely correlated with the reaction kinetics of photogenerated charge carriers.⁴⁶ Fig. 4a presents the linear sweep voltammetry (LSV) curves of 3% -PdS-In₂S₃ and bare In_2S_3 . With the introduction of PdS, the initial overpotential of the binary sample is increased, thus accelerating the reduction of protons to H_2 .⁴⁷ As sketched in Fig. 4b, an enhanced photocurrent response is observed for 3%-PdS– In₂S₃ compared to that of bare In₂S₃, implying that the photogenerated charge pair recombination in 3% -PdS–In₂S₃ samples is effectively improved. The cyclic voltammetry (CV) curves in Fig. 4c also support the above results.⁴⁸ The opencircuit photovoltage (OCP) and consequent attenuation analyses demonstrate that bare $In₂S₃$ exhibits a shorter electron lifetime than the 3% -PdS–In₂S₃ (Fig. 4d), which suggests that the separation of photogenerated carriers is faster on 3% -PdS-In₂S₃.⁴⁹ Furthermore, electrochemical impedance spectroscopy (EIS) was employed to probe the charge transfer resistance on the contact interface of the sample and working electrolyte.⁵⁰ Fig. 4e exhibits that the curvature radius of the EIS plot of 3% -PdS-In₂S₃ is smaller than that of the blank In_2S_3 , revealing a lower resistance and faster charge transfer between the 3% -PdS–In₂S₃ composite and the electrolyte. The mentioned inferences are further supported by PL and time-resolved PL (TRPL) spectra, which are broadly applied to determine the state of photogenerated charge carriers.⁵¹ As shown in Fig. S16, \dagger the PL spectra of both composites show similar emission peak characteristics at approximately 650 nm (λ_{ex} = 360 nm), and a significant diminution of the PL intensity of 3% -PdS-In₂S₃ is observed by the modification of PdS. Furthermore, by calculating the average charge lifetime (τ_a) and fitting the exponential decay dynamic function (Fig. 4f and Table S2†), the average emission lifetime of In_2S_3 (7.71 ns) was found to be longer than that of 3% -PdS–In₂S₃ (5.88 ns), demonstrating that a faster interface charges transfer is achieved in 3%-PdS– In_2S_3 .³⁴ In a nutshell, the above comparative analysis measurements elucidate that the load of PdS can significantly improve the separation and migration of $In₂S₃$ electron-hole pairs, resulting in a noteworthy enhancement of the photocatalytic performance. **Paper**

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For a deep investigation of the interfacial charge transfer route from PdS to $In₂S₃$, a Kelvin probe device was utilized to determine the contact potential difference (CPD, mV) of the composites in dark conditions, through which the work function (WF) of the In_2S_3 and PdS can be obtained. As shown in Fig. 4g, the CPD of $In₂S₃$ to the gold probe is 470 mV, and the CPD of PdS to the gold probe is 600 mV. Thus, the WFs of PdS and In_2S_3 are determined as 5.70 eV and 5.57 eV, respectively, by the relationship of WF (eV) = $5.1 + e \times$ CPD/1000 (where the e represents an electron and the 5.1 eV is the WF of the gold probe). 52 Compared with PdS, the WF of In_2S_3 is lower, which signifies that electronic escaping in $In₂S₃$ is kinetically easier.⁵² Therefore, upon intimate contact between In_2S_3 and PdS, a spontaneous transfer takes place

via the intimate interface of electrons from $In₂S₃$ to PdS until the equilibrium is reached at their Fermi level (FL). As the electrons are withdrawn, the energy band of In_2S_3 bends upwards, which leads to an accumulation of positive charges on In_2S_3 with the negative charges on PdS, resulting in the formation of the interfacial electric field (IEF) directing from In_2S_3 to PdS.⁵³ When illuminated by visible light, the relocation of photoexcited holes remaining on the valence band (VB) of In_2S_3 to PdS is accelerated by the forceful driving force served by the IEF, while the photogenerated electrons on the conduction band (CB) of $In₂S₃$ are blocked back to In_2S_3 by IEF and the upward energy band bending (Fig. 4h).

In order to better reveal the potential dual photoredox reaction mechanism of the photocatalytic process, a range of control and quenching tests were carried out. As shown in Fig. 5a, no product was detected in the absence of catalyst or light, indicating that the reaction is a light-driven catalytic process. Upon incorporation of the electron scavenger carbon tetrachloride (CCl₄), the evolution of H_2 stops, which implies that photoexcited electrons play a vital role in H_2 evolution. Significantly, the N-BDBA product is significantly hindered by the involvement of the hole scavenger triethanolamine (TEOA), whereas the REH value of the reaction is calculated to be ca. 1.0, which suggests that the oxidation and reduction reactions of the system are intimately interrelated. Thermodynamic constraints require that the CB and VB potentials of In_2S_3 should be lower than the reduction potential of H^{\dagger}/H_2 (-0.41 V vs. NHE) and higher than the oxidation potential of BA/BA⁺⁺, respectively, in order to make these two half-reactions proceed successfully. By analyzing the Mott–Schottky curve (Fig. S17†), the band gap structure of In_2S_3 is obtained. The positive slope of the linear plot indicates that $In₂S₃$ is an n-type semiconductor. The VB potential of In₂S₃ of −0.81 V vs. Ag/AgCl can be transformed to −0.61 V vs. the normal hydrogen electrode (NHE) according to the equation of $E_{\text{NHE}} = E_{\text{Ag/AgCl}} + 0.20 \text{ V}^{20}$ Thereby, the CB value of In₂S₃ can be evaluated as −0.61 V, while the VB value of In₂S₃ is 1.57 V depending on the equation $E_{VB} = E_{CB} + E_{g}^{2.5}$ Moreover, depending on the CV curve, the oxidation potential of BA is measured to be 0.84 V vs. Ag/AgCl (Fig. S18†), which is equal to 1.04 V vs. NHE. Therefore, the dual-function photocatalytic system is thermodynamically feasible.

Additionally, in situ Fourier transform-infrared spectroscopy (FT-IR) was employed to obtain insights into the dynamic evolutionary characterization of the reaction (Fig. 5b and c). With the progress of the reaction, characteristic peaks appear at 3278 cm^{-1} and 3366 cm^{-1} , belonging to the symmetric and asymmetric stretching vibration of the N–H bond, respectively, and at 2858 cm^{-1} and 2920 cm−¹ , belonging to the symmetric stretching vibration bands of the methylene bond. The decrease in the intensity of the above characteristic peaks indicates that benzylamine is being consumed.⁵⁴ Meanwhile, the intensity of the newly formed peak at 1645 cm−¹ corresponding to $C=N$ progressively increases with the extension of the

Fig. 5 (a) Control experiments and quenching experiments with varying reagents over 3%-PdS-In₂S₃. (b) and (c) Time-dependent in situ FTIR spectra of 3%-PdS-In₂S₃. (d) Hammett analysis of electronically disparate BA over 3%-PdS-In₂S₃. (e) The quantitative analysis result of EPR spectrums of 3%-PdS-In₂S₃ in Ar-saturated CH₃CN solution in the presence of BA and DMPO. (f) Proposed mechanism of H₂ evolution and BA transformation by visible photocatalysis on PdS –In₂S₃.

lighting duration, indicating the formation of N-BDBA. In addition, the characteristic peaks at 3025 cm^{-1} , 3061 cm^{-1} and 3084 cm−¹ belong to the stretching vibration of the C–H bond of the benzene ring, and the increase in the intensity of the peaks can be attributed to the generation of N -BDBA.⁵⁵

Clarifying the structure of the reaction intermediates is of major importance, for which further radical-quenching tests and electron paramagnetic resonance (EPR) were performed. When the radical trapping agent of 5,5-dimethyl-1-pyrroline-N-oxide (DMPO) is added, the photocatalytic activity of the materials is markedly weakened, which indicates the forming of reactive radical intermediates in the process of the reaction. For identifying the vital intermediate of BA oxidation, the Hammett plot for photocatalytic selective oxidation of various para-substituted BA combined with H_2 evolution over 3% -PdS-In₂S₃ was obtained, as shown in Fig. 5d.⁵⁶ The outcomes demonstrate that the Brown-Okamoto constants (σ^{\dagger}) of the side groups (F, Me, MeO, Cl, and H) have a well-reasoned linear correlation with the log(k_x/k_H) values (reaction constant of $\rho = -0.967$), which not only indicates that the cationic species $(PhCH_2NH_2^{\bullet+})$ are generated during BA oxidation, but also confirms that the cleavage of α -C–H bonds is the rate-limiting step for the formation of N -BDBA.^{3,57} Furthermore, to gain further insight into the details of the reactive intermediates in this photocatalytic system, the EPR technique was employed to characterize the radical intermediate in the reaction with DMPO as the scavenger. As portrayed in Fig. 5e, upon light illumination, six characteristic peaks with similar intensity are observed on both 3%-PdS-In₂S₃ and blank In₂S₃, which can be ascribed to the DMPO-PhCHNH₂ signal, indicating the generation of Ph(·CH)NH₂ free radical ($\alpha_{\rm N}$ = 16.03 and $\alpha_{\rm H}$)

 $= 22.97$).⁵⁸ In addition, the signal belonging to DMPO– PhCHNH₂ on the blank $In₂S₃$ is weaker than that of 3%-PdS– In_2S_3 , which is consistent with the above photoactivity trend.

Given the foregoing discussion, an underlying mechanism of dehydrocoupling of BA to N-BDBA on 3% -PdS–In₂S₃ is put forward as follows: as shown in Fig. 5f, upon irradiation by visible light, the In_2S_3 nanospheres are optically excited to produce the hole–electron pairs. Subsequently, as the IEF is formed by the tight interfacial contact and matched energy level arrangement between In_2S_3 and PdS in the binary composite the photogenerated holes are trapped expeditiously by PdS, while the photogenerated electrons are blocked back to the CB of In_2S_3 . The holes trapped by PdS first attack BA to form the cationic species of $PhCH_2NH_2^{\ast+}$, which is then converted to $Ph(·CH)NH₂$ radicals through deprotonation. Subsequently, the generated radicals of $Ph(\cdot CH)NH_2$ are further attacked by the photogenerated holes to form the imine intermediate of (phenyl)methenamine, then eventually, the (phenyl)methanimine interacts with another BA molecule to form N-benzyl-1 phenylmethylenediamine,⁵⁹ which readily undergoes an addition–elimination reaction, thus ensuring the elimination of ammonia in the resulting N-BDBA (Fig. S19†). Meanwhile, the protons generated in the process are reduced to H_2 by photogenerated electrons on the CB of In_2S_3 .

3 Conclusion

In summary, binary $PdS-In₂S₃$ composites have been successfully fabricated for the efficient photocatalytic oxidation of amines to imines paired with $H₂$ production under visible light irradiation. The matched energy levels

together with the fabrication of an internal electric field between In_2S_3 and PdS accelerate the separation and migration of photogenerated charge carriers, leading to significantly enhanced photoactivity for amine dehydrogenation coupling. The $Ph(·CH)NH_2$ radical has been demonstrated to be the vital reactive intermediate in this photoredox-catalyzed process. Furthermore, such a reaction system not only exhibits great compatibility with a variety of amines, but also affords excellent selectivity and stability. This work is anticipated to assist in the judicious construction of photocatalysts with reasonable modification to adequately use photoexcited carriers simultaneously, offering a prospective and economic pathway for the synthesis of high-value-added chemical compounds. **Paper**
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4 Experimental section

4.1 Materials

Sodium sulfide nonahydrate (Na₂S·9H₂O), indium chloride tetrahydrate (InCl₃·4H₂O), thiosemicarbazide (CH₅N₃S), anhydrous ethanol (C_2H_6O), and acetonitrile (MeCN) were provided by Sinopharm Chemical Reagent Co., Ltd. (Shanghai, China). Benzylamine (C₇H₉N, BA), sodium tetrachloropalladate ($Na₂PdCl₄$), and Nessler's reagent were obtained from Aladdin Biochemical Technology Co. Ltd. (Shanghai, China). No further processing was done to the reagents and they were all used as presented. Deionized (DI) water was obtained from local sources.

4.2 Synthesis of In_2S_3 nanospheres

The blank $In₂S₃$ nanospheres were synthesized according to previous reports with some modifications.⁶⁰ Detailed preparation procedures are described in the ESI.†

4.3 Synthesis of PdS–In₂S₃ composites

The PdS-In₂S₃ composites were prepared via in situ chemical deposition.⁶¹ Typically, a quantity of the as-synthesized In_2S_3 nanospheres was ultrasonically dispersed into 50 mL of 100 mM Na₂S·9H₂O aqueous solution for some minutes, then stirred vigorously at room temperature for a moment. Subsequently, a calculated volume of 10 mM $Na₂PdCl₄$ aqueous solution was rapidly injected under intense agitation. Then, the suspension was exposed to light and stirred vigorously for 0.5 h. The resultant composites were filtered and washed with absolute ethanol and finally dried overnight in a vacuum oven. The obtained catalysts with different mass ratios were named $x\%$ -PdS–In₂S₃ ($x = 0.2$, 0.5, 1, 3, 5, and 7), where the x represents the coefficient of the theoretical mass ratio of PdS to $In₂S₃$.

4.4 Photocatalytic performance testing

The photoactivity testing was carried out based on previous research with some adaptations. In a typical process, 5 mg of photocatalyst, 10 mL of MeCN, and 0.1 mmol of BA were placed into a double-layer quartz reactor. Prior to illumination, argon (Ar) gas was used to purge the reactor for 0.5 h to completely remove air. Then, the suspension was exposed under visible light ($\lambda > 420$ nm) by a 300 W Xe lamp (CEL-HXF300-T3, Beijing China Education Au-light Co., Ltd., China). Finally, the reactor was connected to circulating condensate for the reaction. A gas chromatograph (Shimadzu GC-8A 2014C, MS-5A column) was employed to quantify the generated H_2 . Liquid products were detected through GC-MS (Shimadzu GC-MS QP 2020, Q-Exactive).

Author contributions

Yu Chen: investigation, confirmation, writing-original draft, data analysis, and writing-review & editing. Chang-Long Tan: formal analysis and writing-review & editing. Jing-Yu Li: writing-review & editing. Ming-Yu Qi: writing-review & editing. Zi-Rong Tang: funding acquisition, formal analysis, conceptualization, resources, and supervision. Yi-Jun Xu: funding acquisition, conceptualization, resources, project administration, and supervision.

Conflicts of interest

The authors declare no conflicting interests.

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