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Nanostructured materials for photocatalysis

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Abstract

Photocatalysis is a green technology which converts abundantly available photonic energy into useful chemical energy. With a rapid rise of flow photoreactors in the last decade, the design and development of novel semiconductor photocatalysts is happening at a blistering rate. Currently, developed synthetic approaches have allowed the design of diverse modified/unmodified semiconductor materials exhibiting enhanced performances in heterogeneous photocatalysis. In this review, we have classified the so far reported highly efficient modified/unmodified semiconductor photocatalysts into four different categories based on the elemental composition, band gap engineering and charge carrier migration mechanism in composite photocatalysts. The recent synthetic developments are reported for each novel semiconductor photocatalyst within the four different categories, namely: pure semiconductors, solid solutions, type-II heterojunction nanocomposites and Z-scheme. The motivation behind the synthetic upgrading of modified/unmodified (pure) semiconductor photocatalysts along with their particular photochemical applications and photoreactor systems have been thoroughly reviewed.

Introduction

Photocatalysis is a multidisciplinary field combining physics, chemistry, materials science and chemical engineering pushing the boundaries for many decades for more effective light utilisation and harvesting to promote chemical reactions. The amalgamation of various research fields within photocatalysis spurred exponentially due to the synthesis of multitudinous products, environmental reforming and production of clean hydrogen (H₂) fuel and CO₂ conversion among others.^{1–4} Photocatalysis exhibits several key advantages over conventional catalytic processes involving tedious steps, high temperatures and pressures and using transition metal catalysts.^{5,6}

The implementation of process engineering strategies into photocatalysis pointed out that heterogeneous photocatalysis is preferable to homogeneous in most cases, due to the facile separation of products, higher stability and easier recyclability of photocatalysts. Heterogeneous photocatalysis typically involves photoinduced chemical reaction on the surface of semiconductor materials (photocatalyst) upon exposure to photons.

Titanium dioxide (TiO₂) stood as the main facet of heterogeneous photocatalysis during the past decades, among all semiconductor photocatalysts available.^{7,8} This was mainly due to its high photocatalytic efficiency and several advantages: stability under extreme conditions, appropriate band edge potentials, *etc.* The pivotal motive of our work is to provide an overview of various alternative semiconductor heterogeneous photocatalysts, considered so far and to categorize them into groups depending on their composition and methodology of application. The different classes of photocatalysts – along with their applications in photochemical reactions – mentioned in this review start with binary and ternary semiconductors (SCs) and end up in detailing more prominent aspects of nanocomposite photocatalysts, generated by the conjunction of binary, ternary and other semiconductor materials. This contribution aims to consolidate the different semiconductor systems, the rationale behind their design or

indicate the particular photochemical reactions for which their performances were higher compared to other SCs, along with the type of photoreactor systems in which the photocatalysts were implemented. Obviously, we did not aim to address all the semiconductors here, but a rather intensive focus is made on the most efficient systems. Alternatively, this review addresses the current updated synthetic routes for obtaining such requisite SCs and furnish the description of the polyphasic photocatalytic reactors in which they have been studied. In the past decades manifold of synthetic strategies were reported in modifying the photocatalysts for performing different photochemistries or rendering them active in visible region/any part of the electromagnetic spectrum. These novel attempts emanated into wide range of photocatalytic systems. The progressive motif in the development of semiconductor materials in the regime of sensitized photoreactions has led us to classify them into four different families or categories (see Fig. 1).

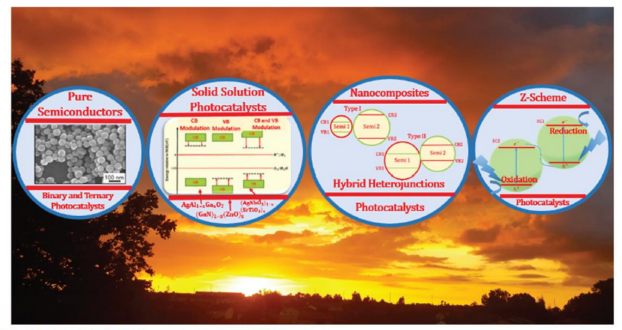


Fig. 1 Classification of semiconductor photocatalysts into four different families.

Binary semiconductor photocatalysts

Synthetic methods of binary SCs have enhanced their photocatalytic performances by achieving monodispersity and unique morphologies with high specific surface area. Diverse nanomaterials with well-defined surface properties are present in different binary SCs, as described in Fig. 2. The intrinsic parameters of the synthesis, nature of the nanomaterials synthesized as result of such reactive domains along with their corresponding photocatalytic applications have been summarized in Table 1.

Oxides-based binary photocatalysts

Teoh *et al.* synthesized TiO₂ nanoparticles by Flame Spray Pyrolysis (FSP) method (Fig. 3a and b), which displayed higher activity for sucrose photo-mineralization than the commercially available Degussa P25.⁹

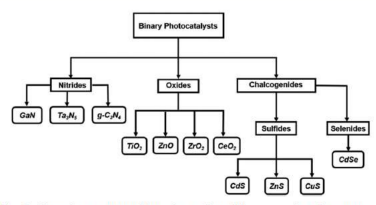


Fig. 2 Flowchart summarizing the various binary semiconductor nano photocatalysts reported in this review.

Direct FSP technique yielded nanomaterials with well-regulated specific surface area and morphology; high purity of crystallite phases without the requirement of post treatment processes (precipitations and centrifugations, *etc.*). Different particle sizes were obtained by varying the flow rate of precursors to the flame spray reactor, portrayed in Fig. 3a. Subsequently, Teoh *et al.* published a detailed review on the successful attempt of FSP for synthesizing diverse binary nanophotocatalysts (ZnO, WO₃, *etc.*), doped binary SCs, noble metal loaded systems and mixed complex metal oxide nanocomposites (NCs).¹⁰

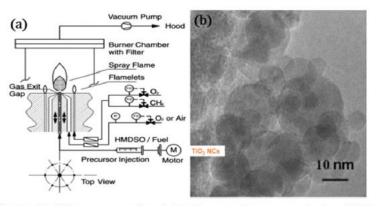


Fig. 3 (a) FSP reactor utilized for the continuous synthesis of TiO₂ photocatalyst,¹¹ more active than the commercial Degussa P25 synthesized by flame aerosol technique. (b) HR-TEM image of TiO₂ Nanocrystals (NCs) synthesized using the FSP reactor.⁹ ((a) Reprint with permission from L. Mädler, H. K. Kammler, R. Mueller, and S. E. Pratsinis, Controlled synthesis of nanostructured particles by flame spray pyrolysis, *J. Aerosol Sci.*, 2002, **33**, 369–389 Copyright 2018 Royal Society of Chemistry. (b) Reprint with permission from W. Y. Teoh, L. Mädler, D. Beydoun, S. E. Pratsinis and R. Amal, Direct (one-step) synthesis of TiO₂ and Pt/TiO₂ nanoparticles for photocatalytic mineralisation of sucrose, *Chem. Eng. Sci.*, 2005, **60**, 5852–5861, Copyright 2018 Royal Society of Chemistry).

Zhanxia *et al.* synthesized peculiar nano-flower (stratified 3D-flower like nanostructures with 2D building block) shaped zirconia (ZrO₂). Their precursor selection (ZOS – zirconium oxide sulphate & sodium acetate) and their synthetic strategy (hydrothermal + calcination) furnished the required 3D-nanoflowers. This peculiar ZrO₂ nanoflower (with a lower specific surface area than the ZrO₂ nanoparticles) discloses enhanced photocatalytic performance for dye degradation than the tetragonal ZrO₂ nanoparticles (NPs) with higher surface area. The higher

activity observed was due to their superior absorbance of photons, as depicted in the absorption spectrum of Fig. 4b.

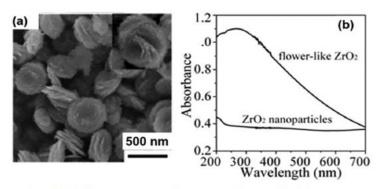


Fig. 4 (a) FE-SEM image of nanoflower shaped ZrO_2 . (b) UV-vis absorption spectrum depicting the enhanced absorption behaviour of the flower shaped nanostructures of ZrO_2 compared to the tetragonal ZrO_2 NPs.¹² ((a and b) Reprint with permission from Z. Shu, X. Jiao and D. Chen, Synthesis and photocatalytic properties of flower-like zirconia nanostructures, *CrystEngComm.*, 2012, **14**, 1122–1127, Copyright 2018 Royal Society of Chemistry).

The enhanced absorption behaviour stems from the specially exposed facets (100) of the petals present in the graded nanoflower structures.¹² Nano-ZrO₂ outperforms several binary nanophotocatalysts (zinc oxide (ZnO) and cerium oxide (CeO₂)) for disparate chemical reactions due to its high negative conduction band (CB) edge potential and wider optical band gaps for different crystalline phases, cubic (3.8 eV), tetragonal (4.11 eV) and monoclinic (4.51 eV).^{13–15} The photochemistry explored so far on both ZnO and CeO₂ were carried out on nanomaterials synthesized in batch mode by hydrothermal process or by simple precipitation techniques.^{16–18} However, due to the rapid development of supercritical fluid flow processes, the synthesis yielded ZnO and CeO₂ nanomaterials with well-defined properties for photocatalysis.^{19,20}

Chalcogenides-based binary photocatalysts

Across the realm of heterogeneous photocatalysts, binary chalcogenides were assessed on equal intensity as binary oxide SCs. Zinc sulfide (ZnS) and cadmium sulfide (CdS) nanophotocatalysts were tremendously researched in the last four decades for their successful applications, not only in environmental purification reactions but also for the reduction of carbon dioxide (CO₂), aldehydes, water splitting and reductive dehalogenation of benzene derivatives. Especially, ZnS was focused a lot, due to their wider energy band gap of about 3.2–4.4 eV²¹ (largest among the binary chalcogenides of group II–VI²²) and its facile phase control of ZnS nanocrystals.²³ A simple, robust production of ZnS nanoporous nanoparticles (NPNPs) *via* the solution phase thermal decomposition technique in the presence of *N*-vinylpyrrolidinone (PVP) was reported by Hu *et al.* These ZnS NPNPs photocatalysts outperformed commercial TiO₂ in degrading eosin B dye.²⁴ They do not require any post treatment process as required by ZnS photocatalyst (synthesized by continuous process) for improving their activity.²⁵ The higher photocatalytic activity of ZnS NPNPs, stems from their high monodispersity and spherically uniform shape. Each sphere of ZnS is an aggregate of hexagonal ZnS nanocrystals (NCs) of 3–5 nm formed as a result of self-assembly process as depicted in Fig. 5.

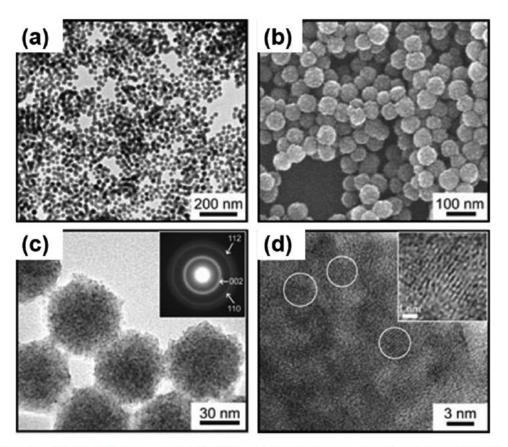


Fig. 5 (a) TEM image of ZnS NPNPs. (b) SEM image of spherical nanoporous ZnS. (c) HR-TEM of ZnS NPNPs with electron diffraction of nanostructures in the inset of the figure. (d) HR-TEM depicting the lattice fringes of ZnS NCs, which undergoes self-assembly process to form the nanoporous ZnS spheres.²⁴ ((a, b, c and d) Reprint with permission from J.-S. Hu, *et al.*, Mass Production and High Photocatalytic Activity of ZnS Nanoporous Nanoparticles, *Angew. Chem., Int. Ed.*, 2005, **44**, 1269– 1273, Copyright 2018 John Wiley and Son).

Identically, the synthesis of hollow spherical CdS and hollow nanoporous copper sulfide (CuS) were reported by Luo et al.²⁶ and Xu et al.,²⁷ respectively. Jing et al. introduced the thermal sulfidation technique replacing the traditional synthetic methodology of precipitation and subsequent heat treatment for obtaining nanosized CdS photocatalysts. The thermal sulfidation technique lowers the phase transformation and crystallization temperature of CdS compared to the other synthetic routes.²⁸ CdS from this technique possesses nano-step arrangement on its surface, exhibited enhanced stability against both air oxidation and photocorrosion during the photocatalytic process. It manifested larger activity for H2 generation than the conventionally synthesized CdS nanomaterial. CdS-based photocatalytic systems profoundly displayed exalted performances for the production of solar-fuel and environmental remediation processes because of its narrow band gap (2.38 eV) and higher negative conduction potential than the reduction potential of H2O/H2.^{26,28,29} Likewise, cadmium selenide (CdSe) based nanophotocatalytic system was also acutely implemented in water splitting for hydrogen (H2) production, inspite of its narrow bulk band gap (1.74 eV). Enhanced photocatalysis of nano CdSe for H2 generation stems from its high magnitude of surface electron charge density and strong quantum confinement behaviour. Wherein precise variation of size of CdSe NCs tunes the energy band gaps providing the appropriate band edge (redox) potentials for photocatalysis.³⁰ A strong relationship

between the photocatalytic H₂ production and various size of CdSe NCs (1.75–4.81 nm) synthesized by solvothermal method was reported by Holmes *et al.*³¹

Nitrides-based binary photocatalysts

Compared to binary oxides/chalcogenides, the exploration of binary nitrides as photocatalysts emerged only in the last two decades. Gallium nitride (GaN) inspection as a photocatalyst started after its immense success in sensing fields due to high mechanical strength and chemical stability (at various pH) compared to other binary SCs.³² The first case of GaN photocatalysis for water splitting reaction implementing powdered GaN was reported by Kida *et al.*³³ Following which, the synthesis of GaN SC in the form of nanowires exhibiting higher activity for photodegradation of organic dyes compared to the submicron GaN dot arrays and thin films was reported by Jung *et al.*³⁴ Tantalum nitride (Ta₃N₅) is an interesting, auspicious, transition metal binary nitride SC used for solar water splitting, due to its narrow direct band gap (2.1 eV) and suitable band edge potentials.^{35–38} For the degradation of dye (methylene blue) under visible light, Ta₃N₅ exhibits higher photocatalytic activity than visible light active nitrogen doped titania nanomaterials.³⁹ Ma *et al.* developed a synthetic strategy where they coupled alkali metal treated tantalum oxide precursor with ammonolysis process yielding highly crystallized Ta₃N₅ nanomaterials, which evinces six times higher photocatalytic activity for water oxidation reaction than Ta₃N₅ nanomaterials synthesized by other techniques.⁴⁰

Graphite-like carbon nitride (g-C₃N₄) is also a direct visible light active SC (2.7 eV).⁴¹ Unlike other SCs, g-C₃N₄ is a nonmetallic conjugated stable polymer, n-type organic SC composed of earth abundant compounds (carbon and nitrogen). It possesses distinctive optical, structural and electronic properties optimum for photocatalytic processes.⁴² g-C₃N₄ implementation in photocatalysis was first demonstrated by Wang et al. for water splitting under visible light irradiation.⁴³ A number of additional studies were subsequently carried out on the synthesis and development of g-C₃N₄-based photocatalysts for applications in solar fuel generation, bacterial disinfection, pollutants degradation and organic transformations.^{44–53} Dong et al. recently reported the synthesis of novel g-C₃N₄ via thermal polymerization using sodium nitrate as oxidizing agent. As-synthesized g-C₃N₄ were able to degrade tylosin (TYL) (antibiotics) within 30 min under simulated sunlight irradiation. The activity was several times higher as compared to conventional g-C₃N₄ photocatalysts. The increase in activity was attributed to the alteration of the g- C_3N_4 crystal structure (induced by sodium nitrate) and a red shift in the visible light absorption range.⁵⁴ To conclude, we have briefly exposed in this section some notable binary photocatalysts and their recent synthetic improvements. For traditional photocatalysts, the synthetic methodologies have been developed rapidly and now continuous synthetic techniques are available to synthesize photocatalysts at very high yield, as seen in the cases of TiO₂ or ZnO. Concerning the unconventional binary photocatalysts like GaN or CuS, the synthetic methodologies⁵⁵ are specially studied to tune their photocatalytic activities not as deeply investigated to date as compared to conventionally synthesized systems.

Ternary photocatalysts

Ternary oxide/chalcogenide SCs comprise two different metallic cations and a single anion. These have been significantly studied in the heterogeneous photocatalytic field because of their stable nature and ability to promote various chemistries as compared to binary SCs. The photocatalysts reviewed in this section can be grouped into different categories according to their composition and crystalline structure as depicted in Fig. 6. Table 2 summarizes the synthesis and details of photocatalytic process of ternary SCs which are reported here.

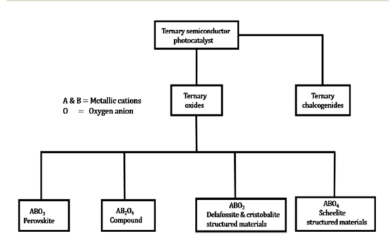


Fig. 6 Flow chart of the diverse ternary semiconductor photocatalysts reviewed in this article.

Ternary oxide photocatalysts

Perovskite photocatalysts (ABO₃ types). Perovskite oxide (cubic crystal structure) SCs, with the general formula ABO₃ are most propitious photocatalysts studied in the field. Here the terms A & B refer to different metallic cations. Predominantly, the 'A' cation is either alkali or alkaline or rare earth metal element whereas the 'B' cation is regularly a transition metal element, whose size is smaller than the A cation. Strontium titanate (SrTiO₃) finds immense application in heterogeneous photocatalysis & thermoelectric fields due to their thermal stability, high photocorrosion resistance and facile capacity to precisely tune its physical & chemical properties depending on its composition.⁵⁶ Such tunability makes SrTiO₃ a nice n-type semiconductor at room temperature via simple electrodonation or doping strategies. SrTiO₃ (3.2 eV)⁵⁷ photocatalyst works effectively for absolute mineralization or degradation of organic pollutants under UV irradiation,^{58–61} enhanced water splitting reaction with a metal cocatalyst loaded on its surface^{57,62} and ammonia production/decomposition reactions.⁶³ The standard solid state synthetic method of nano SrTiO₃ photocatalysts requires elevated temperature but it suffers from polydispersed size distribution and reduced specific surface area. In the last decade, this synthetic process was replaced by hydrothermal,⁵⁹ sol–gel⁶⁴ and solvothermal method.⁶⁵ Mariko et al. reported that SrTiO₃ NPs synthesized by the lower temperature hydrothermal technique exhibited enhanced degradation of nitric oxide (NO) gas (58.70% in 22 min) compared to the SrTiO₃ photocatalysts synthesized by conventional solid state method (56% in 22 minutes) due to the high specific area of (33.1 m² g⁻¹) in the former and lower value (2.88 m² g⁻¹) in the latter case under UV irradiation.66

Unlike titanates, tantalates exhibit photocatalytic water splitting activity without the addition of reduction (NiO or Pt) or oxidation (IrO₂ or MnO_x) co-catalysts due to the higher CB and VB edge of tantalates as compared to the redox potential of water (CB consists of Ta 5d orbital and VB consists of O 2p orbital). CB and VB edge potentials are more negative and positive with respect to reductive and oxidative band edge potentials of the co-catalysts, respectively. The high delocalization of excitons energy is due to corner sharing crystal arrangement.⁶⁷ Lithium tantalate (LiTaO₃) exhibits the highest activity for water splitting under UV light without co-catalyst due to its wide band gap of 4.6–4.7 eV.⁶⁸ The common synthetic methods of LiTaO₃ (solid state synthesis, spray drying methods and polymerizable complex route) yielding large particles with small surface area were replaced by solvothermal and hydrothermal routes. Takasugi *et al.* synthesized LiTaO₃ by both solvothermal and hydrothermal routes. LiTaO₃ obtained from the hydrothermal route which gave very low H₂ production rate (8 µmol h⁻¹). Significant specific surface area (23.7 m² g⁻¹) and very small particle size (10–30 nm) were the reasons for better performance of LiTaO₃ obtained from the solvothermal route.⁶⁹

Sodium tantalate (NaTaO₃), contrary to LiTaO₃, exhibits low activity for water splitting without co-catalyst loading.²⁰ The study of co-catalyst loaded NaTaO₃ system led to the development of lanthanide (La) doped NiO/NaTaO₃ highly active photocatalytic system which yielded tremendous H₂ evolution at a rate of 5.9 mmol h^{-1} under UV irradiation without any sacrificial agent(s).⁷¹ Currently, La-doped NaTaO₃ photocatalysts has been successfully employed for the decomposition of refractory organic compounds, ^{72,73} reduction of CO₂, ⁷⁴ photocatalytic steam reforming of methane⁷⁴ and ammonia synthesis.⁷⁵ Regarding the synthetic prospective of NaTaO₃ photocatalysts, direct band gap NaTaO₃ (4.1 eV) with orthorhombic structure are obtained from the traditional solid state route displaying low activity for water splitting. Replacing the typical solid state by sol-gel technique, NaTaO₃ with a monoclinic structure possessing an indirect band gap (3.9 eV) with high density of states near the band edges are produced, evincing a higher activity compared to solid state synthesized NaTaO₃.⁷⁶ He *et al.* synthesized NaTaO₃ (3.96 eV) nanopowders through a hydrothermal route exhibiting high photoactivity under UV light for both gas (complete photodegradation of formaldehyde, within 35 minutes of irradiation) and liquid (complete photodegradation of rhodamine B (RhB), within 3 hours of irradiation) phase reaction.⁷⁷ Recently, Nguyen *et al.* extended the optical absorption of NaTaO₃ photocatalyst into visible region by co-doping it with La/Bi (Bismuth) through hydrothermal synthesis.⁷⁸ Within the alkali tantalates (ATaO₃, A = Li, Na or K), potassium tantalate (KTaO₃) can be a fascinating alternative for binary TiO₂ photocatalyst due to its low band gap (3.6 eV), appropriate band edge potentials (bottom of CB at -0.7 eV and top of VB at 2.9 eV vs. NHE at pH 0) and proportionately low excitation energy to tap solar energy.^{79,80} Like the above mentioned tantalates, the synthesis of nano KTaO₃ photocatalysts through conventional techniques (solid state or alkoxide method) suffers from large size and defect structure resulting from the hindrance in meticulous control of morphology or size of the nanoparticles due to very high temperature (>1000 °C).^{81–83}

Li *et al.* reported that the size and morphology of KTaO₃ have strong influence on the photoreduction of CO₂ and protons in water. Here, KTaO₃ synthesized by solvothermal method exhibited conjoined nanoflake morphology with low size (15–50 nm) and large surface area (43 m² g⁻¹), displayed an outstanding 20 and 7 times exalted yield for H₂ and CO compared to the conventional cubic morphology attained in solid state synthesis with large size (200–500 nm) and low surface area (<5 m² g⁻¹); without additional sacrificial electron donors and electrical bias.⁸⁴ To conclude, tantalate perovskites have received considerable focus compared to titanate and niobate active oxides due to the appropriate requisite conduction band edge (consisting of Ta 5d orbitals) for the generation of solar fuels.^{85–87} However, the compounds of above mentioned ABO₃ types are active under UV light and Silver Niobate (AgNbO₃) (2.8 eV) has drawn some attention in photocatalysis due to its visible light response.⁸⁸ Such perovskite structured photocatalysts are composed of Ag 4d orbitals, far below the oxidation potential of water and therefore employed to split water under visible light in the presence of sacrificial agents.^{89–92} Arney *et al.* reported the replacement of the synthetic solid state route of AgNbO₃ by means of a molten salt flux synthetic route. This newly developed flux method resulted in materials which exhibited enhanced photocatalytic activity for H₂ generation (5.9 µmol h⁻¹ g⁻¹) from aqueous methanol solution under visible light with respect to the activity displayed by AgNbO₃ synthesized *via* solid state route (3.4 µmol h⁻¹ g⁻¹).⁸⁹

Ternary oxide (AB₂O₄ types) photocatalysts

The second set of ternary oxide photocatalysts elucidated in this review are AB₂O₄ type like calcium indate (Caln₂O₄). Which contain p-block indium (In) metal with a d¹⁰ configuration. Such SC have evinced to form a new set of photocatalysts compared to the platitudinous binary/ternary systems containing d⁰ configuration transition metal ions (Ta, Nb and Ti). Caln₂O₄ have demonstrated the photodegradation of organic pollutants under visible light.⁹³⁻⁹⁵ Similarly, the synthesis of nano Caln₂O₄ *via* conventional Solid State reaction (SSR) suffers from high temperature (>1000 °C) and long calcination time (up to 12 h).⁹³⁻⁹⁶ In 2009, Ding *et al.* reported a solution-combustion synthetic route replacing the traditional route for producing Caln₂O₄ nanorods with large specific surface area and high crystallinity. As-synthesized Caln₂O₄ photocatalyst exhibits 66% mineralization of toluene in gas phase within 360 min of irradiation under visible light, which is four times higher than the activity disclosed by Caln₂O₄ nanomaterials synthesized by traditional SSR route. Similarly, platinum (Pt) dispersed Caln₂O₄ nanorods for water splitting, disclosed a hydrogen generation rate of 1.23 µmol h⁻¹ g⁻¹, which is 24 times higher than the photocatalytic activity displayed by Pt dispersed Caln₂O₄ (synthesized by SSR route).⁹⁷ Still, the solution-combustion methodology suffered

from high temperature annealing and calcination (>1000 °C) and varying calcination time (12–24 h). To eliminate such difficulty in the synthesis of Caln₂O₄, several techniques were rapidly developed in the subsequent years: optimized solid-state involving sequential calcination,⁹⁸ solvothermal,⁹⁹ co-precipitation¹⁰⁰ and sol–gel.^{101,102} Newly, Tavares *et al.* promulgated the synthesis of Caln₂O₄ NCs continuously through ultrasonic spray pyrolysis technique, utilizing laminar flow aerosol reactor as depicted in Fig. 7; exhibiting a precise control of mean particle size. The asdeveloped continuous system yielded Caln₂O₄ NCs in short production time (residence time of 1 min) with well resolved spherical morphology (avg. diameter 511 nm and surface area of 24 m² g⁻¹) and good crystallinity, which exhibited enhanced photocatalytic activity for MB degradation under UV-visible irradiation compared to Caln₂O₄ nanophotocatalysts produced by SSR and solution-combustion routes.¹⁰³

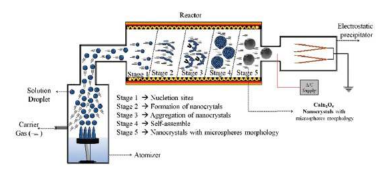


Fig. 7 Pictorial representation of the experimental setup and continuous formation of Caln₂O₄ NCs *via* one pot ultrasonic spray pyrolysis.¹⁰³ (Reprint with permission from M. T. S. Tavares, M. M. Melo, V. D. Araujo, R. L. Tranquilin, C. R. R. Almeida, C. A. Paskocimas, M. R. D. Bomio, E. Longo and V. F. Motta, Enhancement of the photocatalytic activity and white emission of Caln₂O₄ nanocrystals, *J. Alloys Compd.*, 2016, **658**, 316–323, Copyright 2019 Elsevier).

Ternary oxide (ABO₂ types) photocatalysts

The previously discussed ternary SCs illuminate the influence of surface area on the photocatalytic activity, whereas the SCs discussed here and in the subsequent section details the effect of electronic structure on the photocatalytic activity. The third set of ternary oxide SCs discussed here are visible light active, silver containing group III oxides of AgMO₂ types (M = Al, Ga, In).¹⁰⁴⁻¹⁰⁷ This family includes: α -AgGaO₂ (2.38 eV), α -AgInO₂ (1.90 eV), β -AgGaO₂ (2.18 eV) and β -AgAlO₂ (2.95 eV) which are indirect SCs. ' α ' refers to the delafossite structure and ' β ' to the cristobalite related structure. Maruyama *et al.* reported the use of a cation-exchange technique for the synthesis of this AgMO₂ (M = Al, Ga, In) SCs. Ouyang *et al.* performed the photocatalytic evaluation of α -AgGaO₂, α -AgInO₂ β -AgGaO₂ and β -AgAlO₂ SCs by studying the gas phase decomposition of isopropanol (IPA) under visible light.^{107,108} The hierarchy of photocatalytic activity was found to be α -AgGaO₂ > β -AgAlO₂ > β -AgGaO₂ > α -AgInO₂. AgGaO₂ possessing delafossite crystal structure impart the highest activity (88.8 ppm h⁻¹ of acetone produced from IPA decomposition) with respect to other members of AgMO₂ SCs.

Delafossite crystal structure exists in two different polytypes, 3-R (rhombohedral) and 2-H (hexagonal), as depicted in Fig. 8. AgGaO₂ and AgInO₂ synthesized here were crystallized in the 3-R polytype form. In both polytypes, there are two individual layers, providing a separate path for electron (BO₆ layers) and hole conduction (O–A–O dumbbell layer) upon excitation by photons. The repulsion existing between the oxygen anions in the dumbbell (O–A–O) bonds shortens the bond distance between the 'A' cation and the oxygen anion, which increases the interaction between the metallic (A) cation and the oxygen anion, resulting in the formation of highly dispersed VB. Due to such a dispersion, effective mass of the holes decreases and mobility increases leading to increased separation of charges upon excitation, resulting in superior photocatalytic activity. This dispersed VB characteristic is not seen in cristobalite structure (β -AgAlO₂ and β -AgGaO₂) which explains the reasons for a lower photocatalytic activity (5.3 ppm h⁻¹ and 2.2 ppm h⁻¹ of acetone produced, respectively) compared to α -AgGaO₂. The very low photocatalytic activity of α -AgInO₂ (1.3 ppm h⁻¹ of acetone production) for IPA decomposition can be solely attributed to its very low band gap (1.9 eV), not possessing appropriate band edge potentials for oxidation.¹⁰⁸ Regarding α -AgGaO₂ synthesis, Akhtar *et al.* also reported for the first time the high pressure (~10 GPa) direct synthesis of nano α -AgGaO₂ (3R-polytype) using resistively-heated diamond anvil cell, through solid state reaction for which photocatalytic applications are yet to be explored.¹⁰⁹

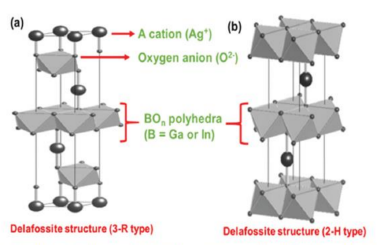


Fig. 8 (a) 3-R polytype of delafossite crystal structure. (b) 2-H polytype of delafossite crystal structure, redrawn based on the images.¹⁰⁶ ((a and b) Reprint with permission from Y. Maruyama, H. Irie and K. Hashimoto, Visible light sensitive photocatalyst, delafossite structured α -AgGaO₂, *J. Phys. Chem. B*, 2006, **110**, 23274–23278 Copyright 2018 American Chemical Society).

Ternary oxide (ABO₄ types) photocatalysts

Bismuth vanadate, BiVO₄ (2.4 eV) is a visible light active SC, of high interest due to its polymorphic nature, ferroelastic and electronic properties. Such fascinating properties strongly depend on its crystalline structure. The three polymorphic forms of BiVO₄ are scheelite-monoclinic (SM), scheelite-tetragonal (ST) and zircon-tetragonal (ZT).^{110–112} Among the crystalline forms of BiVO₄, SM exhibits the highest activity for photocatalytic O₂ evolution from aqueous silver nitrate solution and photocatalytic degradation of endocrine compounds, especially alkylphenol groups under visible light irradiation.^{113–115} SM outperforms ST in photocatalysis. In the SM–BiVO₄ structure, a variety of bond lengths between Bi–O atoms are observed, unseen in ST–BiVO₄ structures. The presence of such different bond lengths in Bi–O atoms leads to distortion of the crystal structure caused by the 6s² lone pairs of Bi³⁺ cations. Specifically, VO_4^{3-} tetrahedrons are distorted (Fig. 9a). Due to the SM structure distortion, the overlap of VB orbitals occurs. The overlapping extent of O 2p and Bi 6s orbitals in VB increases with the increase of the extent of distortion in the crystal structure and results in the increase of charge separation upon photoexcitation, due to the delocalization of electrons and holes. These resulted in enhanced migration of holes to the surface of the SC, leading to superlative photocatalytic activity. Such intensified migration of holes is absent in ST–BiVO₄ (Fig. 9b) semiconductor due to the absence of distortion in the scheelite crystal structure.^{110,116} Traditionally, SM–BiVO₄ are synthesized by solid state reaction at high temperature. Kudo *et al.* prepared BiVO₄ photocatalysts for O₂ evolution via room temperature aqueous process, which resulted in nanomaterials with well-defined crystallinity.¹¹⁵ Yu and Kudo reported the hydrothermal route to synthesize SM–BiVO₄ with enhanced surface texture and morphology as compared to nano BiVO₄ synthesized via aqueous process route. They successfully enhanced the photocatalytic activity for O_2 evolution from aqueous silver nitrate solutions by varying the pH of the synthetic medium. The variation of the pH (1, 4 and 9) of the synthetic medium successfully changed the electronic band gap (2.39, 2.46 and 2.47 eV) of the resultant nanomaterials, thus positively influencing the activity of photocatalysis.¹¹⁶ The synthesis of highly crystalline SM–BiVO₄ nanopowders (avg. size 20 nm) with spherical like morphology was more recently optimized under mechanochemical synthesis. Due to the practical issues associated with separation of photocatalysts from the reactant/product streams in large scale implementations, BiVO₄ syntheses have been

intensively focused on SM–BiVO₄ thin films in this decade. SM–BiVO₄ thin films exhibited efficient separation, stable monoclinic scheelite structure and pre-eminent light absorption capability.^{117–122} Lately in 2017, Venkatesan *et al.* combined mechanochemical synthesis and Rf-sputtering for the synthesis of BiVO₄ films with nano-structural islands which displayed good activity for the degradation (complete degradation in 240 min) of rhodamine-6G (Rh6G) under visible light.¹²³

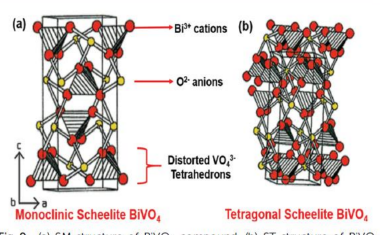


Fig. 9 (a) SM structure of $BiVO_4$ compound. (b) ST structure of $BiVO_4$ compound.¹²⁴ (Reproduced from Schwarz, *et al., Eur. Coat. J.*, 2016, 9, 26–31, Copyright 2018 Vincentz Network).

Besides the above mentioned SC oxides, aurivillius compounds, $Bi_2A_{n-1}B_nO_{3n+3}$ (A = Ca, Sr, Ba, Pb, Na, K; B = Ti, Nb, Ta, Mo, W, Fe) have also been considered. This particular group possesses a unique special layer structure where $(Bi_2O_2)^{2+}$ layers are interleaved with perovskite-like $(A_{n-1}B_nO_{3n+1})$ units.¹²⁵ Ternary SC, bismuth tungstate (Bi_2WO_6) is one of the simplest member (n = 1) of aurivillius oxides. Bi_2WO_6 possesses interesting physicochemical properties and was extensively explored in photocatalysis in the last decade. Such an intensive study is attributed to its unique structure $((Bi_2O_2)^{2+}$ layers sandwiched with perovskite-like $[WO_4]^{2-}$ layers) favouring efficient separation of photogenerated charge carriers upon excitation, leading to exalted photocatalytic activity.¹²⁶ Advanced photocatalytic systems employing Bi_2WO_6 SC are reported in the last part of this review (see section on Z-Scheme photocatalysts).

Ternary chalcogenide photocatalysts

The final set of ternary SCs addressed in this review are ternary chalcogenides of the type AB_xC_y (where A = Cu, Ag, Zn or Cd; B = Ga or In; C = S or Se or Te). These compounds exhibit high absorption coefficient over a wide spectral range and are effectively feasible for solar photon absorption. Due to their excellent optoelectronic and catalytic properties, they have been successfully implemented in solar cells, light emitting diodes, radiation detection devices and photocatalysis.^{127–137} Among these highly functional chalcogenides, SCs of the group I–III–VI (silver–indium–sulfide (AgInS₂), copper–gallium–sulfide (CuGaS₂) and copper–indium–sulfide (CuInS₂)) and group II–III–VI (cadmium–indium–sulfide (CdIn₂S₄) and zinc–indium–sulfide (ZnIn₂S₄)) have been studied in photocatalysis. Within these two groups, group I–III–VI semiconductors: AgInS₂, CuGaS₂ and CuInS₂ are highly preferred due to their negligible toxicity, high photoconductivity, high stability, environmental benign composition and visible light activity for many novel reactions.^{135–143} The course of studies on group I–III–VI photocatalysts have specifically focused on the control of morphology and physicochemical properties, thereby influencing its photocatalytic activity. In this perspective, AgInS₂ (1.87–2.14 eV)^{144,145} photocatalysts were synthesized by optimized novel techniques in recent times including hydrothermal methods,¹⁴⁶ one-step aqueous synthesis,¹⁴⁷ microwave hydrothermal techniques¹⁴⁸ and microwave heating processes with short reaction times (*ca.* 5 min).¹⁴⁹ Strikingly, Hu *et al.* synthesized AgInS₂ nanoplates and nanotubes through convenient co-precipitation low-temperature (70–190 °C) strategy where the morphology and

physicochemical properties of AgInS₂ photocatalysts were altered by varying (In/Ag) molar ratio of precursors. Synthesized AgInS₂ with (In/Ag) molar ratio of 9 exhibited a nanotube-like morphology, low photoluminescence value and maximum photocurrent density (0.011 mA cm⁻²). The maximum photocurrent implies the most efficient separation of electron–hole pairs upon excitation. This novel AgInS₂ nanotube displayed regeneration ability and highest visible light activity for complete 2-nitrophenol degradation (within 120 min) as compared to other AgInS₂ nanoplates/nanotubes due to the narrow band gap and highest charge separation efficiency.¹⁵⁰

Similar synthetic studies have been reported for CuGaS₂ photocatalysts. Classic synthetic methods (hydrothermal,¹⁵⁰ solvothermal¹⁵¹ and hot injection route¹⁵¹) suffered from drawbacks including polydispersion and limited scale-up production capabilities. Regulacio *et al.* introduced a facile non-injection route based on thermal decomposition. This synthesis presented several advantages including high reproducibility, mass production of samples and convenient control of the synthetic domain. As-synthesized nano CuGaS₂ (2.52 eV) exhibited promising photocatalytic activity for RhB dye degradation (63% of dye was degraded in 270 min) under visible light irradiation.¹⁵² Additionally, Liu *et al.* reported a non-injection route based one-step heating process for the synthesis of one-dimensional (1D) or two-dimensional (2D) CuGaS₂ NCs. Here, the morphology tuning could be achieved through the appropriate choice of ligands and precursors. These NCs displayed prominent activity for H₂ evolution in the presence of hole scavengers (Na₂S and Na₂SO₃) under simulated solar irradiation.¹⁵³ Most reported ternary SCs to date have not been synthesized under continuous conditions. In addition to this, the only ternary SCs synthesized under continuous supercritical processes have not been explored in heterogeneous photocatalysed processes to date.^{154,155}

Solid solution photocatalysts

The optical properties of SCs are a reflection of their inherent electronic structure, related to their VB to CB transition. In the case of wide bandgap SCs, their optical gap circumscribes many redox potentials but their implementation is limited due to their activity in the UV region. Certain ternary SCs possess narrow band gap (active in the visible region) but no appropriate band edge potentials to perform diverse reactions. Consequently, the inherent electronic structure in both binary and ternary SCs pure materials does not regulate an optimum balance between visible light activity and required redox potential.

In the heterogeneous photocatalytic field, meticulous modulation of band gap *via* preparation of solid solution SC are generally more beneficial as compared to doped or loaded SCs (binary or ternary). Optimum formulations of solid solution systems can be successfully accomplished by combining a wide band gap binary/ternary SC with very narrow band gap binary/ternary SC, *i.e.* ZnS–CdS solid solution.¹⁵⁶ The resultant solid solution $(Zn_{1-x}Cd_xS (0.2 \le x \le 1))$ does not only exhibit activity under visible light but also maintains its suitable band edge potentials to perform the required reactions. Apart from the band gap position, other properties, which have to be considered while synthesizing the solid solutions include similarities in the crystal structure, electronegativity and valence state of the two end members with which the solid solution is fabricated.¹⁵⁷ Generally, the hybridized CB or VB promotes faster charge transfer due to its dispersed nature and increases the photocatalytic activity. This phenomenon is more pronounced in solid solutions than in ternary or binary photocatalysts. According to IUPAC, solid solution is defined as, "a crystal containing a second constituent, which fits into and is distributed in the lattice of the host crystal. (The term 'solid solution' is not referred for amorphous materials)".¹⁵⁸ The diverse solid solution photocatalysts reviewed here (see Fig. 10), can be divided into three different categories, depending on the strategy employed for engineering the band gap:

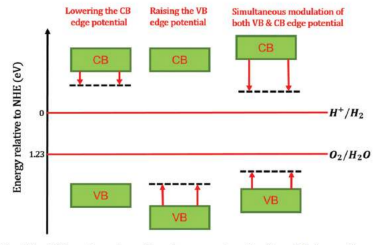


Fig. 10 Different modes of band gap engineering for attaining optimum balance between visible light activity and redox potential of semiconductor band edge.

- 1. Solid solution through continuous modulation of CB,
- 2. Solid solution through continuous modulation of VB,
- 3. Solid solution through continuous modulation of both CB and VB.

Solid solutions through continuous modulation of CB

Ternary SCs of the family AgMO₂ (M = Al, Ga, In) have attracted significant attention in the photocatalytic field due to their dispersed VB (see section ternary oxides (ABO₂ types) photocatalysts). The solid solution fabricated with $AgMO_2$ SCs can effortlessly extend into the entire spectrum of the visible light and still retains the VB potential. It displays photocatalytic decomposition of various organic compounds due to the strong oxidizing potential. Shuxin et al. improved the visible light activity of β -AgAlO₂ (2.83 eV) for IPA decomposition through continuous tuning of the CB edge of β -AgAlO₂, by combining it with narrow band gap β -AgGaO₂ (2.18 eV) SC, resulting in a β -AgAl_{1-x}Ga_xO₂ (0 ≤ x ≤ 1) solid solution.¹⁵⁹ They synthesized β -AgAl_{1-x}Ga_xO₂ (0 $\leq x \leq$ 1) through two-step (sol–gel followed by cation exchange) method. The linear band gap variation of the compound with respect to the composition (see Fig. 11a UVvis absorption spectrum) indicates that continuous solid solution through precise tuning of CB was obtained in this system. Among the various compositions, β -AgAl_{0.6}Ga_{0.4}O₂ (2.58 eV) exhibited highest photocatalytic activity for IPA decomposition. The rate of acetone production (photo-decomposed product of IPA) by β -AgAl_{0.6}Ga_{0.4}O₂ photocatalyst was 35 and 63 times higher than its end components β -AgAlO₂ and β -AgGaO₂, respectively. The high activity is attributed to the novel band gap which customizes an optimum balance between the visible light activity and appropriate redox potential. The colour of the solid solutions with different compositions and acetone evolution rates with respect to band gaps of continuous CB modulated solid solution is depicted in Fig. 11b.¹⁵⁹ A similar CB modulation was achieved recently in the family of AM₂O₆ (A = Sn and M = Nb or Ta) ternary SCs by fabricating the series of $Sn(Nb_{1-x}Ta_x)_2O_6$ ($0 \le x \le 1$) solid solutions. $Sn(Nb_{0.8}Ta_{0.2})_2O_6$ photocatalyst was proved to provide excellent activity and stability for the complete mineralization of MB dye (98.9% mineralization efficiency in 120 min) under visible light irradatiation as compared to other members of the $Sn(Nb_{1-x}Ta_x)_2O_6$ ($0 \le x \le 1$) solid solutions.¹⁶⁰

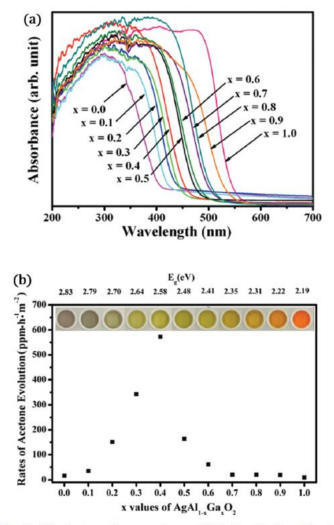


Fig. 11 (a) UV-vis absorption spectrum of β -AgAl_{1-x}Ga_xO solid solution fabricated from β -AgAlO₂ and β -AgGaO₂ ternary oxide SCs by precise tuning of CB from 2.83 eV to 2.19 eV.¹⁵⁹ (b) Photocatalytic activity for acetone production.¹⁵⁹ ((a and b) Reprint with permission from S. Ouyang and J. Ye, β -AgAl_{1-x}Ga_xO₂ Solid-Solution Photocatalysts: Continuous Modulation of Electronic Structure Toward High-Performance Visible-Light Photoactivity, *J. Am. Chem. Soc.*, 2011, **133**, 7757–7763, Copyright 2018 American Chemical Society).

Solid solutions through continuous modulation of VB

Niobium-based perovskite SCs were extensively explored in photocatalysis, since they suffice the requirement of intensified stability. As their CB edge potential is far more negative than the potential of O_2/O_2 ^{--.90,161} Silver niobate (AgNbO₃) (2.8 eV) is one such SC with enhanced stability. It possesses narrow band gap (2.8 eV) due to its hybridized VB, composed of Ag 4d and O 2p orbitals and exhibits visible light active activity for producing O_2 from silver nitrate solution. Its oxidation potential was however not sufficient to decompose several organic compounds.¹⁶²

Li *et al.* improved the photo oxidizing ability of AgNbO₃ by combination with NaNbO₃ (3.4 eV) perovskite SC through solid state synthetic method, resulting in Ag_xNa_{1-x}NbO₃ ($0 \le x \le 1$) solid solution. The CB edge potential of both AgNbO₃ and NaNbO₃ are pinned at the same level (-0.4 eV vs. NHE).⁹⁰ A decrease in lattice parameters and Nb–O bond length in NbO₆ occurs by increasing NaNbO₃ content in the solid solution, resulting a decrease of particle size. Simultaneously, the hybridization ratio of Ag 4d and O 2p decreases resulting in the continuous modulation of VB, shifting the VB top from AgNbO₃ (+2.39 eV) to VB top of NaNbO₃ (+3 eV). These findings indicate that the oxidation potential of the solid solution system is comparatively more positive to that of AgNbO₃. Among the continuous range

of Ag_xNa_{1-x}NbO₃ ($0 \le x \le 1$) solid solutions, Ag_{0.6}Na_{0.4}NbO₃ photocatalyst evidenced enhanced activity for gaseous IPA decomposition under visible light (activity three times higher with respect to AgNbO₃) due to an optimum state achieved between oxidation potential, visible light absorption ability and particle size.¹⁶³ Continuous VB modulation without CB alteration was also reported on d^{10} -(oxy) nitride $(Ga_{1-x}Zn_x)(N_{1-x}O_x)$ ($0 \le x \le 0.25$) solid solution by Maeda et al.¹⁶⁴ fabricated from d¹⁰-ZnO (3.2 eV) whose CB bottom is composed of hybridized s and p metallic cation. The resultant solid solution has heavily dispersed CB with increased mobility from photogenerated electrons, thus intensifying the photocatalytic activity. GaN (3.4 eV) was chosen as another constituent here, as N 2p orbitals possess higher potential energy than O 2p orbitals and similar CB with respect to ZnO. $(Ga_{1-x}Zn_x)(N_{1-x}O_x)$ (0 $\leq x \leq 0.25$) was designed for overall water splitting under visible light irradiation. Such solid solution formation featuring requirements of narrow band gap below 3 eV, appropriate band edge potential and stability for overall water splitting could provide an upliftment of the VB, leading to narrow band gap SCs (see Fig. 12b, decrease of band gap with increase of ZnO content). The upraise of VB potential to CB (without any changes) was hypothesized due to the repulsion existing between O 2p and Zn 3d orbitals¹⁶⁵ (Fig. 12a). The authors performed the synthesis of this solid solution through nitridation of oxide precursors (Ga₂O₃ & ZnO) at high temperatures (>850 °C) under ammonia flow. As-synthesized solid solution exhibited negligible activity for overall water splitting with co-catalysts including 1 wt% Rh (rhodium) and 1.5 wt% Cr (chromium) on $(Ga_{1-x}Zn_x)(N_{1-x}O_x)$ ($0 \le x \le 0.25$) system. Solid solution with a very low Zn quantity (Zn/Ga = 0.14) exhibited enhanced photocatalytic H₂ and O₂ production of 360 μ mol h⁻¹ and 162 μ mol h⁻¹, respectively.¹⁶⁵ The above conventional nitridation method suffered from several drawbacks including long nitridation times, high temperature (>850 °C) and large (micron) sized particles. Recently, Menon et al. replaced the conventional nitridation by solution combustion method,¹⁶⁶ which yields particles in the nanoscale (40 nm), the resultant $(Ga_{1-x}Zn_x)(N_{1-x}O_x)$ possessed high zinc content (Zn/Ga = 20, 30 and 40). The solid solution with (Zn/Ga = 20)exhibited overall water splitting under visible light without the aid of a co-catalyst, with apparent quantum yield of H₂ production at 0.048%.¹⁶⁶ Dharmagunawardhane et al. developed a high pressure (1 GPa) and high temperature (1150–1200 °C) synthetic methodology to synthesize $(Ga_{1-x}Zn_x)(N_{1-x}O_x)$ solid solutions over the entire compositional range (x = 0.07 to 0.9) using a talc pressure cell via solid state reaction. They observed the smallest band gap (2.65 eV) and highest photocatalytic activity of H₂ evolution (2.3 µmol h⁻¹) from water in the absence of co-catalysts, pHmodifiers and sacrificial agents for (Ga_{0.49}Zn_{0.51})(N_{0.49}O_{0.51}) solid solution under visible light.¹⁶⁷

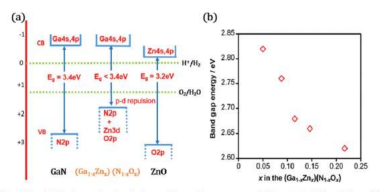


Fig. 12 (a) Schematic depiction of band structure of $(Ga_{1-x}Zn_x)(N_{1-x}O_x)$ solid solution (redrawn based on the article¹⁶⁵). (b) Variation of band gap with respect to ZnO content in the solid solution.¹⁶⁵ Fig. 11(a and b) (Reproduced with permission from ref. 165. Copyright 2018 American Chemical Society).

Solid solutions via continuous modulation of both CB and VB

Simultaneous regulation of equilibrium between absorption extension into visible region and redox abilities of photocatalysts was achieved in this type of solid solutions prepared *via* synchronous precise tuning of both VB and CB. Due to the favourable inclination of partial substitution of comprehensive metallic cations and valences at the position of A and B elements in ABO₃ perovskites, these materials stand as highly promising photocatalysts. Wang *et al.* performed the precise tuning of both CB and VB in $Ag_{1-x}Sr_x(NbO_3)_{1-x}(TiO_3)_x$ ($0 \le x \le 1$) solid solutions, formed by combining visible active AgNbO₃ (450 nm) and UV active SrTiO₃ (380 nm) perovskites.¹⁶⁸ The solid solution was Chunping Xu, Prasaanth Ravi Anusuyadevi, Cyril Aymonier, Rafael Luque, Samuel Marre. Nanostructured materials for photocatalysis. *Chemical Society Reviews*, Royal Society of Chemistry, 2019, 48 (14), pp.3868-3902. (10.1039/C9CS00102F). (hal-02185643)

synthesized *via* solid state reaction method. The resultant solid solution's CB is composed of hybridized Nb 4d and Ti 3d orbitals. As Nb 4d is more positive than Ti 3d, an increase of AgNbO₃ content brings the bottom of CB band downwards. Similarly, VB of this system was composed of hybridized Ag 4d and O 2p orbitals where Ag 4d is more positive than O 2p and it pushes the top VB upwards thus favouring perfectly engineered SC active in the visible region with an optimum balance between redox potentials. The presence of such hybridized CB and VB in this solid solution with continuous dispersion favors enhanced charge carrier transportation upon excitation, leading to increased photocatalytic activity. The solid solution obtained here exhibited continuous modulated CB and VB for the composition ($0 \le x < 0.95$) and discontinuous CB and VB for the composition ($0.95 \le x \le 1$).¹⁶⁸ Discontinuous interbands (dopant levels) are due to very low concentration AgNbO₃ in the solid solution. Among the continuous series of solid solutions, Ag_{0.75}Sr_{0.25}(NbO₃)_{0.75}(TiO₃)_{0.25} evidenced the highest photocatalytic activity for O₂ production from silver nitrate solution (162μ mol h⁻¹), with an apparent quantum efficiency of about 16.4% under visible light, compared to its two end members. The same photocatalyst displayed the highest activity for IPA decomposition, where it completely mineralize IPA to CO₂ within 22 h of visible light irradiation.¹⁶⁸

As previously mentioned for perovskite oxides ABO₃ type (see section on perovskite photocatalysts), 'A' site is occupied by a cation larger in size than the cation at the 'B' site. This smaller cation on the 'B' site has a stronger interaction with oxygen than the larger cation at 'A' site. Possibly, perovskites may manifest different optical and electronic properties depending on the electronegativity and ionic radius of the smaller cations at the 'B' site.¹⁶⁹ The confirmation reported by Grinberg et al. that narrow band gap perovskites can be achieved by using two different transition-metal cations on the 'B' site¹⁷⁰ led to the synthesis of perovskites like Ba₂FeNbO₆ (BFNO) with mixed Fe³⁺ and Nb⁵⁺ cations on the 'B' site. In Ba₂FeNbO₆ (2.29 eV), the VBM composed of Ni 3d and O 2p orbitals while CBM comprised Fe 3d and Nb 4d. This VB and CB straddles the water oxidation and reduction potential, respectively, however leading to low photocatalytic activity due to poor charge separation efficiency. The photocatalytic performance of Ba₂FeNbO₆ was improved via preparation of SrTiO₃–Ba₂FeNbO₆ solid solution NCs through a simple one-step molten salt route.¹⁷⁰ Here the band gap was precisely tuned from 3.31 eV of SrTiO₃ to 2.29 eV of Ba₂FeNbO₆ through simultaneous modulation of CB and VB edges (Fig. 13). Among the series of solid solutions synthesized, $SrTiO_3$ -Ba₂FeNbO₆-0.5 (SrTiO₃ to Ba₂FeNbO₆ optimal molar ratio is 10:0.5) exhibited the highest photocatalytic H₂ production (7.15 μmol h⁻¹) from aqueous methanol solution under visible light irradiation.¹⁷⁰ This SrTiO₃–Ba₂FeNbO₆-0.5 solid solution exhibited maximum photocurrent signal indicating that its high photocatalytic activity is attributed to efficient separation of electron-hole pairs. Simultaneous regulation of redox potential and visible light activity is achieved facilely in this system.¹⁷⁰

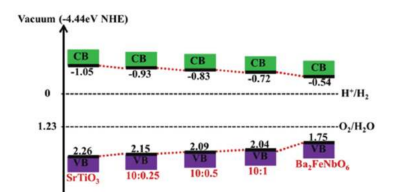


Fig. 13 Depiction of energy level diagram of $SrTiO_3$, $SrTiO_3-Ba_2FeNbO_6$ solid solutions (ratio 10:0.25, 10:0.5 and 10:1 corresponds to the molar ratio of $SrTiO_3$ to Ba_2FeNbO_6).¹⁷⁰ (Reprint with permission from G. Zhang, S. Sun, W. Jiang, X. Miao, Z. Zhao, X. Zhang, D. Qu, D. Zhang, D. Li, Z. Sun, *Adv. Energy Mater.*, 2017, **7**(2), 1600932. Copyright 2019 John Wiley and Son).

Sulfide-based solid solution photocatalysts have been extensively reported to be synthesized by continuous tuning of both CB and VB for efficient hydrogen production.^{171–174} Binary sulfides including CdS found to be moderately active photocatalysts for H_2 production (see section on binary photocatalysts), with inherent issues related to quick recombinations of photogenerated charge carriers and photo-corrosion under visible light (essentially limiting their use at a large scale^{175–178}).

The formulation of highly crystalline solid solutions combining wide band gap ZnS with CdS can solve the photostability issues of CdS under visible light irradiation. The synthesis of $Cd_xZn_{1-x}S$ solid solutions was achieved employing several techniques such as co-precipitation,^{179,180} cation-exchange¹⁸¹ and hydrothermal methods.^{182,183} Such conventional techniques possessed several drawbacks including high deviation of elemental composition of solid solutions compared to theoretical synthesis values, minimum control over band structure and long reaction times to obtain highly crystallized solid solutions. Li *et al.* circumvented the above mentioned drawbacks *via* facile thermolysis of complex zinc–cadmium–thiourea to synthesize $Cd_xZn_{1-x}S$ (0 < x < 1) solid solutions were found to be aggregates of small crystallized NPs (avg. size = 10 nm) (Fig. 14a and b). The continuous red shift of steep absorption band edge with increasing CdS content confirmed the precise engineering of both CB and VB for these solid solutions (Fig. 14d).¹⁸⁴

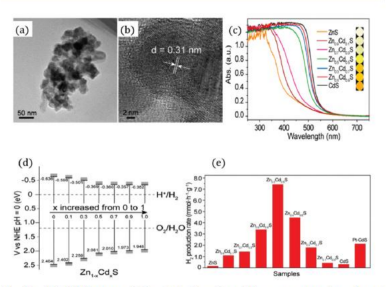


Fig. 14 (a) TEM image of $Zn_{0.5}Cd_{0.5}S$ solid solution, average diameter of aggregates (50–100 nm). (b) HR-TEM image of $Zn_{0.5}Cd_{0.5}S$ solid solution, diameter of each NP is about 10 nm. (c) UV-vis diffuse reflectance spectra of $Cd_xZn_{1-x}S$ solid solutions (x = 0, 0.3, 0.5, 0.7 and 0.9). (d) Band gap variation with respect to the composition. (e) H₂ production rate with respect to the composition in $Cd_xZn_{1-x}S$ solid solution.¹⁸⁴ Fig. 13(a–d) (Reproduced with permission from ref. 184. Copyright 2018 American Chemical Society).

Among the different ranges of solid solution, $Cd_{0.5}Zn_{0.5}S$ (2.45 eV) exhibited a superior balance between CB and VB potentials as well as visible light absorption. This photocatalyst exhibited maximum H₂ production (7.42 mmol h⁻¹ g⁻¹) from water in the presence of sacrificial agents (Na₂S & Na₂SO₃) in the absence of a co-catalysts (Pt).¹⁸⁴ This photocatalytic activity exceeded the activities of CdS and ZnS by 24 and 54 times, respectively, being much higher than the activity displayed by co-catalyst (Pt) loaded CdS photocatalysts (Fig. 14e).¹⁸⁴

Fu *et al.* successfully incorporated the Cd_{0.5}Zn_{0.5}S solid solution (synthesized by the previously disclosed thermolysis route¹⁸⁴) into PAN (polyacrylonitrile) nanofibers and developed leaf-like, bio-inspired flexible mat-shaped Cd_{0.5}Zn_{0.5}S@PAN photocatalysts.¹⁸⁵ This hierarchically porous leaf-like solid solution exhibited a remarkable photocatalytic activity for H₂ production from water splitting (475 µmol h⁻¹ of H₂ per 50 mg) at an apparent quantum efficiency of 27.4% at 420 nm. Cd_{0.5}Zn_{0.5}S@PAN photocatalytic activity was reported to be superior to that of Cd_{0.5}Zn_{0.5}S solid solution NPs due to the enhancement of mass transfer kinetics, multiple reflection and scattering effect of incident light as result of the porous structure.¹⁸⁵

Ternary chalcogenide SCs of the group I–III–VI (I = Cu, Ag; III = Al, Ga, In; VI = S, Se, Te) have received increasing attention as a new class of materials in the field of solar cells and also for solar driven H₂ production (water splitting).¹⁸⁶ In ternary sulfide SCs including AgInS₂ or CuInS₂, the VB is composed of d-states of Ag or Cu assorted with p-states of sulfur,^{187,188} resulting in a VB far more negative to that of ZnS or CdS.

In these ternary compounds, the VB is precisely tuned by monovalent cations of Ag or Cu and the CB is modulated *via* Indium doping (Fig. 15a and b). These ternary SCs were combined with wide band gap ZnS for the synthesis of solid solution photocatalysts for optimized H₂ evolution.

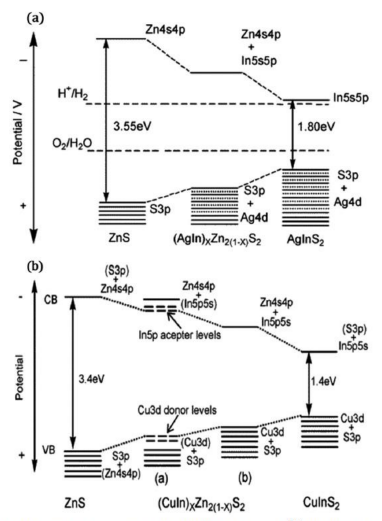


Fig. 15 (a) Band gap modulation in $(AgIn)_{x}Zn_{2(1-x)}S_{2}$.¹⁸⁹ (b) Band gap engineering attained in $(CuIn)_{x}Zn_{2(1-x)}S_{2}$ solid solutions¹⁹⁰ Fig. 14. (a) Reproduced with permission from ref. 189. Copyright 2018 American Chemical Society. Fig. 14(b) (Reproduced with permission from ref. 190. Copyright 2018 American Chemical Society).

Tsuji *et al.* reported the synthesis of $(AgIn)_x Zn_{2(1-x)}S_2$ ($0 \le x \le 1$) (3.55–1.8 eV) solid solution through the heat treatment of complex greyish Ag–In–Zn sulfide precursor.¹⁸⁹ Here, they attempted several different heat treatment strategies, from which an improvement of the photocatalytic activity for the resultant solid solutions was observed. $(AgIn)_{0.22}Zn_{1.58}S_2$ (2.33 eV) synthesized *via* heat treatment under N₂ flow exhibited a relevant H₂ production (72 µmol h⁻¹) from aqueous solutions of K₂SO₃ and Na₂S under visible light irradiation.¹⁸⁹

The same photocatalyst when synthesized *via* heat treatment, under sealed quartz ampule tube without N₂ flow, provided a remarkably superior activity for H₂ production (328 µmol h⁻¹).¹⁸⁹ The photocatalytic activity was 4.5 times higher than the material synthesized under N₂ flow, as the newly proposed heat treatment methodology prevents the volatilization of metallic cations during synthesis. On loading 3 wt% of co-catalyst (Pt) on the (AgIn)_{0.22}Zn_{1.58}S₂ photocatalyst (synthesized by heat treatment in sealed quartz tube), the rate for H₂ production under visible light was further enhanced three times (944 µmol h⁻¹) with a highest apparent quantum yield of 20%.¹⁸⁹ Subsequently, the same authors employed the above newly modified heat treatment process (850 °C for 5 h) in sealed quartz tube for the synthesis of (CuIn)_xZn_{2(1-x)}S₂ (0 ≤ *x* ≤ 1) (3.5–1.4 eV) solid solutions.¹⁹⁰ For very low CuInS₂ content (*x* < 0.05), discontinuous discrete energy levels were formed in the system (Fig. 15b). Above this composition, continuous VB is formed and monotonous red shift to visible region occurs.¹⁹⁰ Among the solid solutions synthesized, (CuIn)_{0.09}Zn_{1.82}S₂ evidenced moderate photocatalytic activity for H₂ production (7 µmol h⁻¹) from aqueous solutions containing sacrificial agents (K₂SO₃) under visible light irradiation.¹⁹⁰ The same photocatalyst with 0.5 wt% of co-catalysts (Pt) loading displayed an enhanced activity for H₂ production (684 µmol h⁻¹) with an apparent quantum yield of 12.5%.¹⁹⁰

Following the successive visible light activity for H₂ production using the above two different chalcopyrite (Ag or CuInS₂)–ZnS photocatalysts, Tsuji *et al.* developed further the heat treatment technique for solid solution synthesis.¹³⁸ They reduced the heat treatment temperature from 850 °C down to 600 °C and increased the heat treatment duration of complex single source solid solution precursor from 5 h to 20 h.¹³⁸ ZnS–CuInS₂–AgInS₂ solid solutions were synthesised through improved heat treatment (reduced temperature and prolonged heat treatment) and compared in terms of photocatalytic activity to ZnS–CuInS₂ and ZnS–AgInS₂ solid solution photocatalysts. Prior to photocatalytic studies, all three different solid solutions were loaded with 0.75 wt% of ruthenium (Ru) co-catalysts on their surfaces. The (CuAg)_{0.15}In_{0.3}Zn_{1.4}S₂ solid solution with the aid of co-catalysts (Ru) exhibited the maximum photocatalytic activity for H₂ production (2.3 mmol h⁻¹) from aqueous solution containing sacrificial agents (K₂SO₃ & Na₂S).¹³⁸

Most of the solid solution nanomaterials synthesized so far have specifically targeted the control of the physicochemical properties by the variation of chemical composition. More specifically, the photoinduced electron transfer rate was heavily altered in these ternary-based solid solution by the electronic energy levels through the composition of the solid solution. Concurrent control of size and composition is a preferable strategy to obtain optimum photochemical properties in ternary based solid solutions. The first synthesis of size controlled solid solution (ZnS–AgInS₂) was reported by Gabka *et al.* for the production of NCs of ZnS–AgInS₂ solid solution photocatalysts of *ca.* 2–10 nm in size. These NCs displayed color tunable photoluminescence (PL) by simultaneous modulation of NC's size, shape and composition.¹⁹¹ Subsequently, Tatsuya *et al.* reported the size controlled synthesis of ZnS–AgInS₂ solid solution NCs with a fixed composition without changing their particle shape *via* thermal reaction of the metal acetate with thiourea.¹⁹² Due to quantum size effect, the difference in H₂ evolution potential (*E*[H⁺/H₂], –0.76 V *vs.* NHE) and the CB edge potential of NCs increased with a decrease in particle size.¹⁹² Each composition of ZnS–AgInS₂ solid solution NCs with decreased particle size provided an enhanced H₂ evolution rate. However, the H₂ production rate decreased with a further decrease in particle size due to the formation of defect sites/deeper trap sites.¹⁹² The highest photocatalytic activity was observed for (AgIn)_xZn_{2(1-x)}S₂ ($0 \le x \le 1$) solid solutions having both optimum sizes of about 4.2–5.5 nm and energy gap of 2.3–2.4 eV.¹⁹²

In conclusion, solid solution photocatalysts can exhibit very high activity under visible/UV irradiation as optimum balance between required redox potentials and absorption band edge. Simultaneous modulation of size and composition has now gained importance for a more precise tuning of the electronic levels depending on the Chunping Xu, Prasaanth Ravi Anusuyadevi, Cyril Aymonier, Rafael Luque, Samuel Marre. Nanostructured materials for photocatalysis. *Chemical Society Reviews*, Royal Society of Chemistry, 2019, 48 (14), pp.3868-3902. (10.1039/C9CS00102F). (hal-02185643)

required photochemical properties. The main drawbacks or limiting steps in the synthesis of solid solution photocatalysts include (i) the precursor preparation step and (ii) the cost of the single source precursor. An overview of the synthetic methodologies and synthesis conditions of solid solution photocatalysts along with heterogeneous photochemical systems is reported in Table 3.

Nanocomposite photocatalysts

An advanced photocatalyst must exhibit two critical features: (i) an ameliorated light absorption capacity and (ii) an enhanced photocatalytic performance. These crucial features provide insights for design of composite photocatalysts by coupling two or three different SCs together, the coupling based on the energy exchange between SCs with different electronic structures. Unlike SC–metal composite systems, SC–SC coupled systems feature a composite in which both materials individually absorb the photonic radiation, resulting in harvesting more photonic energy. The first crucial requirement of enhanced light absorption is satisfied here.

Whenever a SC is coupled to a different SC, it results in a heterostructure with three possible different types of heterojunctions depending on the respective position of CB & VB levels of each SC: (i) straddling gap (type I) (ii) staggered gap (type II) and (iii) broken gap (type III) (Fig. 16).

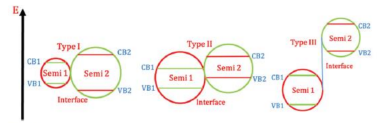


Fig. 16 Schematic representation of energy gap of three types of SC–SC heterojunctions.

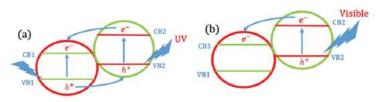


Fig. 17 Band gap diagram of type-II heterojunctions illustrating the photo-generation of charge carriers and efficient separation of them upon (a) UV excitation and (b) visible light excitation.

The vectorial transfer of electrons and holes in these SC–SC coupled heterostructured junctions are determined by the band gap, the electron affinity (bottom edge of CB) and the work function (top edge of the VB) of the coupled semiconductors.

Within the three types of heterojunctions SCs, type-II hetero-composite system is most desirable for photocatalysis. The difference in chemical potential (arising from the appropriate position of band edge potentials) (VB & CB of semiconductor 2 > VB & CB of semiconductor 1) between the two different SCs in type II system results in band bending at the interface of the SCs. Band bending phenomenon instigates an electric field, which drives the electrons and holes of SCs upon photoexcitation in the opposite direction, leading to spatial separation of photon generated charge carriers consequently enhancing their photocatalytic activity.^{193,194}

Regularly, such type-II systems are designed in such a way that it results in photocatalyst possessing enhanced light absorption and efficient charge separation. To have an enhanced light absorption, heterojunction composite

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materials are generally composed of one SC active in the UV region (TiO₂ or ZnO) while the other material is active in the visible region (CdS, CdSe, *etc.*). When type-II composite materials are activated by UV radiation, both the band gaps are excited, electrons migrate from the CB2 to CB1, holes migrate from the VB1 to VB2, thus resulting in efficient separation of charge carriers (Fig. 17a).

In the case of visible light excitation, only the narrow band gap material is excited, the electrons from CB2 migrate to CB1 and holes remain within the VB2, resulting in efficient charge separation (see Fig. 17b). Thus in these SC–SC coupled systems having type-II heterojunction, two critical conditions are satisfied, enhanced light absorption and efficient charge separation resulting in higher photocatalytic performance.

Considerations of type-II nanocomposite photocatalytic systems

Importance of SC 1/SC 2 ratio. The light absorption limit of type-II composites is determined by the band gap of the narrower SC, active in the visible region. The absorption intensity of the composite can be increased by increasing the amount of narrow SC. Nevertheless, this does not necessarily enhance the photocatalytic performance of the composite system, as it has been seen in examples, including AgI–BiOI composite systems.¹⁹⁵ Consequently, determining the optimal composition of both SCs in type-II systems is crucial to reach an appropriate balance between the intensity of light absorption and photocatalytic performance.

Importance of interfaces. In SC–SC nanocomposite systems formed between diverse SCs, there are possibilities for the presence of defects at the material interface due to lattice mismatch. These defects can trap the photogenerated electrons or holes leading to recombination. This prevents the optimum use of photogenerated electrons and holes for redox reactions. In contrast, the lattice mismatch can lead to an electric field at the interface between the two materials, which separates the charges further leading to higher photocatalytic activity. The quality of the heterogeneous interface stands to be another crucial factor influencing the photocatalytic performance.¹⁹³ In order to obtain high quality interfaces with less number of defects, high temperature annealing is usually employed in the synthetic methodologies. The annealing temperature is critical and determines the quality of the interface, which ultimately affects the photocatalytic performance, as exemplified in the case of Bi₂O₃–SrTiO₃.¹⁹⁶ The quality of the heterojunction interfaces in various nanocomposites can also be improved by careful selection of appropriate synthetic methods.

p-n type junction composite systems

Apart from conventional type-II heterojunctions, basically made up of n-type SCs, another type of type-II system has been extensively reported formed by coupling n-type and p-type SCs, resulting in a heterojunction called p–n type junction. This is an exceptional type of heterojunction which possesses the previously mentioned advantages of composite systems (*i.e.* increased light absorption limit and effective charge carrier separation). Additionally, these p–n type junction composites possess an internal electric field at the interface formed between the SCs. This field further intensifies the charge separation process, leading to an enhancement of photocatalytic performance. Specifically, reported p–n junction type-II photocatalysts exhibit a higher reductive activity as compared to its individual constituents, with a lower oxidative activity. This is because under equilibrium conditions, when an internal electric field is formed at the interface of the p–n junction composite material, the p-type becomes negatively charged and the n-type becomes positively charged. Upon excitation, the holes start moving towards the negative side and electrons towards the positive side, where the p-type semiconductor behaves as a hole trap and scavengers, unfavourable for photocatalytic oxidation reactions.

These NCs include Cu₂O/TiO₂, NiO/TiO₂, p-ZnO/n-TiO₂, CuBi₂O₄/Bi₂WO₆, Co₃O₄/ZnO and Cu₂O/In₂O₃.¹⁹⁷⁻²⁰¹

TiO₂-Based nanocomposite photocatalysts

TiO₂ photocatalysts have been tremendously studied for several decades due to their very high oxidation ability, high photostability and low cost.202,203 Still, the use of titania systems is hindered due to its wide band gap (3.2 eV) of anatase phase, requiring UV irradiation (λ < 380 nm) for photocatalytic activation.204 On other hand, CdS is a relatively narrow band gap SC (2.4 eV), extensively researched and modified (see section on solid solution photocatalysts) for H2 production from water splitting reactions under visible light irradiation. The growth of CdS on

Chunping Xu, Prasaanth Ravi Anusuyadevi, Cyril Aymonier, Rafael Luque, Samuel Marre. Nanostructured materials for photocatalysis. *Chemical Society Reviews*, Royal Society of Chemistry, 2019, 48 (14), pp.3868-3902. (10.1039/C9CS00102F). (hal-02185643)

 TiO_2 results in a TiO_2/CdS type II heterojunction, where upon illumination by visible light, photo-generated electrons from CB of CdS are transferred to CB of TiO₂ and holes remain within CdS (Fig. 18b), resulting in a more efficient charge separation as compared to that of the two individual SCs, which ultimately increases photocatalytic efficiency.205 Different types of TiO₂/CdS nanocomposites have been synthesized by different techniques including (i) CdS QDs sensitized in ordered mesoporous titania through ion-exchange route205 and (ii) CdS NPs coated on TiO_2 nanotubes via wet chemical methods.²⁰⁵ Vu et al. additionally reported a non-noble co-catalyst loaded TiO₂/CdS NCs (Ni–TiO₂/CdS).²⁰⁶ This system comprised TiO₂ nanorods, NPs of CdS and Ni clusters. Capped TiO₂ were synthesized through a solvothermal route featuring surfactant-controllable nanorod lengths. On these nanorods, CdS NPs were grown using a colloidal synthesis, resulting in hybrid TiO₂/CdS nanocomposites. Subsequently, co-catalyst (Ni) clusters were selectively deposited on NCs via a photodepositon route. A schematic illustration of the synthetic route for Ni–TiO₂/CdS nanocomposites is shown in Fig. 18. Ni–TiO₂/CdS exhibited high photocatalytic activity for H₂ production (151 µmol g⁻¹) from aqueous solutions containing sacrificial agents (ethanol), compared to the activity of CdS (3.4 μ mol g⁻¹) under visible light irradiation. Such activity (44 times higher than CdS) was reported to be achieved due to the enhancement of charge separation upon photoexcitation by charge transfers from CdS to TiO₂ nanorods and then to the co-catalyst (Fig. 18b). Besides traditional nanocomposites prepared by coupling UV (wide band gap) and visible light (narrow band gap) active SCs, nanocomposites synthesized by combining two wide band gap SCs were also found to be active under visible light irradiation. Such visible light activity is due to the introduction of impurities and defect levels in the forbidden gap of the nanocomposite. TiO₂/ZnO is an example of such visible light active systems, formed by combining two wide band gap UV active SCs (TiO₂ (3.32 eV) and ZnO (3.37 eV)). Zhang et al. synthesized TiO₂/ZnO nanocomposites via a two-step method consisting of homogeneous hydrolysis and low temperature crystallization.²⁰⁷

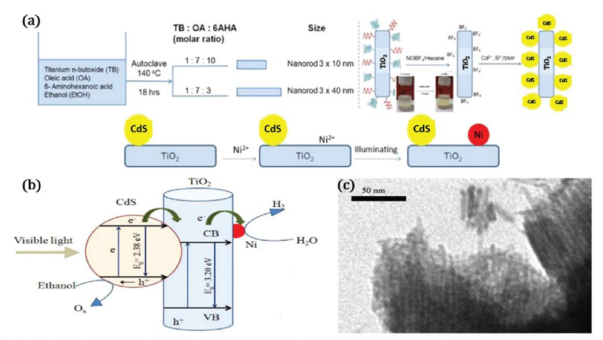


Fig. 18 (a) Illustration of synthetic methodology for obtaining Ni–TiO₂/CdS photocatalysts.²⁰⁶ (b) Charge transfer pathway in Ni–TiO₂/CdS nanocomposite upon visible light irradiation for H₂ production from aqueous ethanol solution.²⁰⁶ (c) TEM image of TiO₂ nanorods.²⁰⁶ (a, b and c) (Reproduced with permission from ref. 206. Copyright 2018 American Chemical Society).

The observed absorption red shift of the TiO_2/ZnO nanocomposite system into the visible region was claimed to be due to the introduction of dopant energy levels in the TiO_2 forbidden band gaps by the Zn^{2+} cations.²⁰⁷ Importantly, TiO_2/ZnO composite exhibited an improved photocatalytic efficiency with respect to TiO_2/ZnO nanocomposites successfully synthesized through hydrothermal, spin coating and atomic layer deposition techniques (where these strategies were also found to be economically unfavourable in the long run^{208–210}). Hussein *et al.* synthesized

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mesoporous coupled TiO₂/ZnO composite with varying composition (TiO₂: ZnO) employing a facile and cheap sol–gel route. As synthesized materials with 30 and 50 wt% of ZnO coupled with TiO₂ exhibited high photocatalytic activity for H₂ production (770 μ mol g⁻¹ and 739 μ mol g⁻¹, respectively) from aqueous solutions containing methanol as an electron acceptor, under UV irradiation as compared to the moderate activity of commercially available TiO₂, Degussa P25 (356 μ mol g⁻¹).²¹¹ Prasannalakshmi *et al.* reported the synthesis of rod shaped TiO₂/ZnO NCs *via* sol–gel method, which exhibited a high activity for the photocatalytic degradation of organic dyes (brilliant green (BG) and methylene blue (MB)) under solar irradiation compared to the activity exhibited by their corresponding individual SCs.²¹² Similar to the TiO₂/ZnO system, TiO₂/ZnS photocatalyst is yet another interesting visible light active system composed of two different UV active SCs: TiO₂ (3.2 eV) and ZnS (3.6 eV). The visible light activity is due to the formation of new energy levels as a result of sulphur doping within the forbidden band gap of TiO₂.

Štengl *et al.* synthesized spherical agglomerates of TiO₂/ZnS nanocomposites consisting of alternate nanosized titania and sphalerite crystalline islands *via* homogeneous hydrolysis route.²¹³ The resultant TiO₂/ZnS (2.9 eV) nanocomposite displayed the highest activity for photocatalytic degradation of orange II dye under visible irradiation ($\lambda = 400 \text{ nm}$) (degradation constant for nanocomposite, $k = 0.0069 \text{ min}^{-1}$) as compared to the activities of commercially available Degussa P25 (3.1 eV) ($k = 0.0022 \text{ min}^{-1}$) and pure anatase (3.2 eV) ($k = 0.0009 \text{ min}^{-1}$). The enhanced photocatalytic activity was attributed to the reduced charge transfer resistance and increased surface area for the reactant adsorption.²¹³ NPs of TiO₂ and TiO₂/ZnS nanocomposites were also recently reported to be synthesized *via* sol–gel method, where the TiO₂/ZnS system exhibited enhanced activity for the degradation of organic dyes (BG and MB) under solar irradiation compared to TiO₂ NPs.²¹⁴

ZnO-Based nanocomposite photocatalysts

ZnO SCs in nanosize regime possesses certain advantageous features including (i) a high exciton binding energy of 60 meV (as comparison TiO₂ has 4 meV), (ii) a high electron mobility of 200 cm² V⁻¹ s⁻¹ (TiO₂ has 30 cm² V⁻¹ s⁻¹), (iii) a facile adaptation/tuneability of the structure, (iv) an anisotropic growth and (v) an simplicity of crystallization.^{215–220}

In spite of such distinct features, both ZnO and TiO₂ suffer from low photocatalytic efficiency due to higher rate of recombination of photoinduced electrons and holes compared to the rate of charge transfer to the surface bound redox reactions.²²¹ Additionally, they exhibit very low visible light activity due to their wide band gaps. Coupling SC to ZnO resulting in type-II band offset is generally of two types. ZnO can either be coupled to similar binary oxide wide band gap SC (as seen in the case of TiO_2/ZnO NC). The second type of coupling involves growing of lattice matched photosensitizer inorganic crystals on ZnO surface, which ensures a maximum harvesting of visible light and consequently an enhanced photocatalytic activity.²²² The sensitizers should possess a narrow band gap and appropriate band edge potentials for the transfer of the photogenerated electrons from its CB to ZnO CB under visible light activity, leading to enhanced separation of charge carriers.²²³ Among various inorganic sensitizers, the coupling of In_2S_3 to ZnO has been recently reported of interest as In_2S_3 possesses a narrow band gap (2–2.3 eV) with higher carrier mobility and good stability in photocatalysis.²²⁴ Khanchandani et al. reported the synthesis of ZnO/In₂S₃ core/shell nanorods array employing a soft chemistry approach consisting of surface functionalization route by using citric acid as functionalization agent (Fig. 19).²²⁵ ZnO nanorods and In₂S₃ NPs were synthesized *via* solvothermal method. ZnO nanorods were subsequently successfully coated with In₂S₃ employing a surface functionalization route (see Fig. 19). The one-dimensional ZnO nanorods structure is preferred here as it provides an optimum geometrical structure facilitating effective transport of charge carries and reduces the charge recombination to a certain extent within the ZnO nanostructures.²²⁶

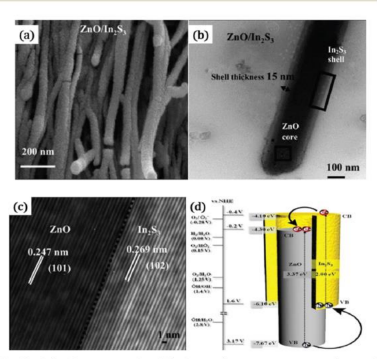


Fig. 19 (a) FESEM image of ZnO/In_2S_3 core/shell nanorod arrays. (b and c) TEM and HRTEM of ZnO/In_2S_3 core/shell NCs. (d) Illustration of charge transfer and separation in NC systems.²²⁵ (a, b, c and d) (Reproduced with permission ref. 225. Copyright 2018 American Chemical Society).

As synthesized core/shell composites exhibited a type-II band offset as its absorption band edge shifted to the visible region (due to the incorporation of In_2S_3) and exhibited high photocatalytic activity (degradation constant, k = 0.0174 min⁻¹) for RhB dye degradation under visible light irradiation, compared to ZnO nanorods (k = 0.00039 min⁻¹) and In_2S_3 NPs (k = 0.0042 min⁻¹). The enhanced photocatalytic activity of ZnO/In₂S₃ core/shell NCs (56 times higher than that of ZnO nanorods and 5 times higher with respect to In_2S_3 NPs) was due to the effectual separation of photoinduced charge carries, attributed to the type-II heterojunction formed between the two different SCs.²²⁵

Similarly to In₂S₃, CdS was used to form a type-II heterojunction nanocomposite with ZnO. As CdS possesses a narrow band (2.4 eV) gap for visible light activation, similar lattice constant to ZnO and high optical absorption coefficient. Khanchandani *et al.* extended the same surface functionalization route employed for ZnO/In₂S₃ nanocomposites to the synthesis of ZnO/CdS nanorods core/shell array systems. Citric acid (CA) was used as the surface functionalization agent (Fig. 20).²²⁷ By varying the CA amount, the shell thickness of CdS was successfully tuned. The influence of the shell thickness on the photocatalytic activity of the nanocomposite was also reported. The authors synthesized ZnO/CdS (core/shell) materials with three different shell thickness: 10 nm, 20 nm and 30 nm. The absorption band edge of ZnO nanorods was red shifted to visible region when they are coated with CdS shell. The monotonous red shift to the visible region increased with an increase in the shell thickness. As compared to ZnO nanorods and CdS NPs, the core/shell nanocomposites displayed high activity for the photocatalytic degradation of RhB under simulated solar radiation.²²⁷ Within all nanocomposites with varying shell thickness, ZnO/CdS with maximum shell thickness exhibited optimum photocatalytic activity due to an extended absorption in the visible region.

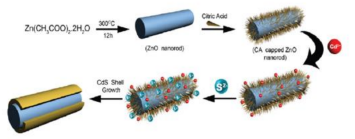


Fig. 20 Schematic depiction of the surface functionalization route for the synthesis of ZnO/CdS core/shell NCs.²²⁷ (Reproduced with permission from ref. 227 (2012) Copyright 2018 American Chemical Society).

The degradation rate constant (k (min⁻¹)) of ZnO nanorods, CdS NPs, ZnO/CdS (10 nm shell thickness), ZnO/CdS (20 nm shell thickness) and ZnO/CdS (30 nm shell thickness) are 0.011, 0.0035, 0.019, 0.025 and 0.039 min⁻¹, respectively.

SnO₂-Based nanocomposite photocatalysts

Tin(iv) oxide (SnO₂) is also an ideal constituent in nanocomposite systems due to its wide band gap (3.6 eV) and CB edge potential (E_{CB} = 0 V vs. NHE, at pH = 7), lower than the CB potential of most SCs implemented in photocatalytic organic mineralization/degradation reactions. Additionally, SnO₂ is largely available at a low cost, possesses excellent chemical stability, high optical transparency in the visible range (97%) and good electrical properties.^{228–232}

Wang *et al.* fabricated SnO₂/TiO₂ nanocomposites by hydrothermally growing SnO₂ on TiO₂ nanofibers (synthesized by electrospinning technique).²²⁸ In contrast to this two-step synthesis (electrospinning + hydrothermal), Zhang *et al.* synthesized SnO₂/ZnO nanocomposites *via* direct electrospinning technique.²³³ Both SnO₂/TiO₂ and SnO₂/ZnO nanocomposites exhibited high photocatalytic activity for the degradation of RhB under UV light as compared to the reduced activities of their individual constituent SCs.

Iron oxide (α -Fe₂O₃) phases are frequently coupled to SnO₂ and studied in the heterojunction domain for applications in photocatalysis, gas and liquid sensors.^{234–237} α -Fe₂O₃ is a n-type semiconductor with a narrow band gap (2.2 eV) exhibiting poor photocatalytic performance due to small diffusion length of holes (2–4 nm). The efficient separation of photogenerated charges are observed when α -Fe₂O₃ is coupled to SnO₂.

Kang *et al.* synthesized necklace-like hierarchical SnO_2/Fe_2O_3 nanocomposites employing a chemical vapour deposition route (CVD), growing hierarchically Fe_2O_3 hexagonal disks on nanocables consisting of SnO_2 as core and epitaxial Fe_2O_3 as shell (Fig. 21a and b). As synthesized novel hierarchical SnO_2/Fe_2O_3 system exhibited high photocatalytic activity (complete degradation in 2.5 h) for the degradation of MB under visible light as compared to pure SnO_2 (synthesized by CVD, exhibited 83% degradation within 2.5 h) and commercially available Fe_2O_3 (58% degradation of MB within 2.5 h).²³⁸

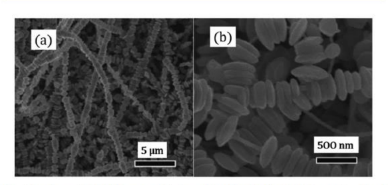


Fig. 21 (a and b) SEM image of hierarchical SnO₂/Fe₂O₃ NCs synthesized by CVD technique.²³⁸ Reprint with permission from J. Kang, Q. Kuang, Z. X. Xie and L. S. Zheng, Fabrication of the SnO₂/ α -Fe₂O₃ hierarchical heterostructure and its enhanced photocatalytic property, *J. Phys. Chem. C*, 2011, **115**, 7874–7879, Copyright 2018 American Chemical Society.

Xia *et al.* synthesized a non-core/shell SnO_2/Fe_2O_3 nanocomposite through a co-precipitation technique, where the resultant composite reveals exalted activity for the degradation of acid blue 62 dye under visible light. These SnO_2/Fe_2O_3 NCs degraded 98% of the dye within one hour, which is 3.6 times higher than the activity revealed (27% degradation of dye in one hour) by commercially available titania (Degussa P25) photocatalyst.²³⁴

SnO₂/SnS₂ is yet another visible light active nanocomposite system developed and studied especially for photocatalytically mediated environmental remediation applications. Tin sulfide (SnS₂) is a n-type narrow band gap SC (2.18–2.44 eV), standing as an ideal compound for dye sensitization studies due to the absence of surface states associated with dangling bonds. These dangling bonds act as traps for the photogenerated electrons.²³⁹ Tin sulphide has been therefore employed as photocatalyst in the visible light region due to its thermal, oxidative stability in air and stable existence without corrosion in acidic and neutral solutions.²⁴⁰ Zhang *et al.* synthesized non-core/shell type SnO₂/SnS₂ nanocomposites with a firm interface contact *via in situ* oxidation of SnS₂ under hydrothermal (in the presence of H₂O₂) conditions.²⁴¹ By varying the duration of oxidation and the amount of H₂O₂, the SnO₂ content in the nanocomposite was successfully varied (see HR-TEM image, Fig. 22). As-resultant SnO₂/SnS₂ (18.1 mass% of SnO₂) exhibited high photocatalytic activity (degradation constant, *k* = 9.260 min⁻¹) for MO degradation under visible light as compared to the activity of SnS₂ (*k* = 5.941 min⁻¹) (synthesized by hydrothermal reactions) and physical mixture of SnO₂/SnS₂ (7.094 min⁻¹).²⁴¹

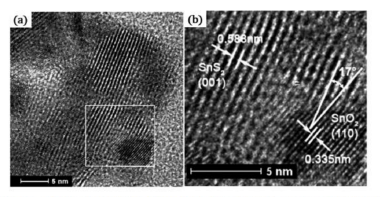


Fig. 22 (a) HR-TEM of SnS_2/SnO_2 (18.1 wt% of SnO_2). (b) Enlargement of the interface observed in figure.²⁴¹ (a and b) (Reproduced with permission from ref. 241. Copyright 2018 American Chemical Society).

Magnetically separable nanocomposite photocatalysts (Fe3O4-based systems)

Photocatalysts with magnetic supports or magnetically separable materials (Fe₃O₄) coupled to SCs forming a nanocomposite can be easily separated from the liquid phase by applying an external magnetic field. This allows overcoming the limitations associated with separation techniques, resulting in high recovery and reusability of the photocatalysts.²⁴² Importantly, iron oxide (Fe₃O₄) undergoes phase transition, when subjected to heat treatment at higher temperature (400 °C). This phase transition leads to a loss of the magnetic properties. Since most of the magnetic nanocomposites were synthesized *via* sol–gel process (where heat treatment is crucial) the preparation of highly and stable photoactive titania–magnetic nanocomposite with enhanced magnetic properties is rather challenging.²⁴³ The essential criteria to design highly photocatalytic active magnetic nanocomposite system using Fe₃O₄ (0.1 eV) SC involves the construction of nanocomposites with special structures or the development of special types of interface phase junction (*e.g.* insertion of an insulation layer) between the magnetic materials and the active photocatalysts.

In the perspective of active SC photocatalysts, bismuth oxyhalides BiOX (X = Cl, Br or I) of the group V–VI–VII ternary SCs have attracted significant attention in recent times due to their optical and crystalline properties. BiOX SCs possess tetragonal matlockite structures in which $[Bi_2O_2]$ layers are interleaved between halogen layers, resulting in a unique open layered structure and complex band structure. Such crystalline structure possesses large space between the layers ($[Bi_2O_2]$ and halogen layers), which facilitates the separation of photo-induced electron–hole pairs. Additionally, it exhibits low recombination probability of photo induced electron–hole pairs due to its indirect band gap, thus resulting in a significantly improved photocatalytic activity.^{244–246} Extensive reviews have been published on BiOX (X = Cl, BR, I) SCs, detailing their recent synthetic developments and photocatalytic applications: organic pollutants degradation, fuel production and removal of heavy metals.^{247–250}

Amongst BiOX (X = Cl, BR, I) SCs, BiOCl exhibits the highest photocatalytic activity under UV irradiation, with comparably higher activity to that of TiO₂ for the degradation of dyes. Its activity under visible light irradiation can be additionally improved *via* preparation of heterojunction nanocomposites.²⁵¹ Zhang *et al.* synthesized a Fe₃O₄/BiOCl nanocomposite *via* colloidal synthesis, where BiOCl (bismuth oxychloride) flakes are inlaid with Fe₃O₄ NPs (see Fig. 23b and TEM image in Fig. 24b).

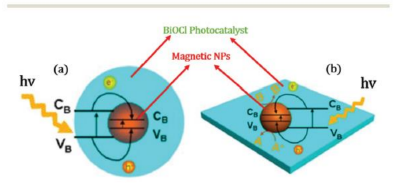


Fig. 23 (a) Illustration of conventional core shell $Fe_3O_4/BiOCl NCs$ with charge transfer route. (b) Coupled structure of $Fe_3O_4/BiOCl NCs$, where Fe_3O_4 NPs are inlaid in the BiOCl nanoflakes.²⁵² Reproduced with permission from ref. 252. (Copyright 2018 Elsevier).

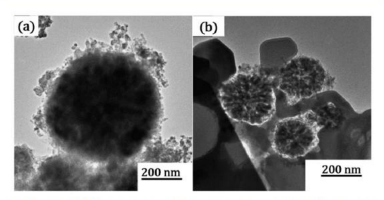


Fig. 24 (a) TEM image of conventional core/shell $Fe_3O_4/BiOCl$ NCs. (b) TEM image of coupled $Fe_3O_4/BiOCl$ NCs.²⁵² (Reproduced with permission from ref. 252. Copyright 2018 Elsevier).

The as-obtained Fe₃O₄/BiOCl structure exhibited similar photocatalytic activity for the degradation of RhB dye under visible light as compared to the activity of BiOCl nanoflakes. Here, facile separation of photocatalysts could be achieved using an external magnetic field and the recycled catalysts maintained their photocatalytic activity even after five consecutive runs.²⁵² In this particular system, the photocatalytic activity of BiOCl was preserved upon combination with the magnetic semiconductor due to the special type of coupled heterostructure, where the rate of charge transfer to redox reactions is higher than the rate of recombination, despite this being a type-I heterojunction.²⁵²

Chalcogenide nanocomposite photocatalysts (CdS/ZnS nanocomposites)

Similar to TiO₂/Fe₃O₄ and Fe₃O₄/BiOCl possessing type-I heterojunction, CdS/ZnS nanocomposites are another type of hybrid heterostructures with type-I phase junction extensively employed in heterogeneous photocatalysis.

CdS/ZnS nanocomposite was successfully synthesized using metal organic chemical vapour deposition (MOCVD)^{253,254} and wet chemical synthetic methods consisting of surface functionalization routes.^{255,256} Wang *et al.* synthesized 1-D (dimensional) CdS/ZnS nanocomposite with nanowire morphologies through a two-step mild solution method consisting in a solvothermal reaction.²⁵⁷ This synthetic strategy did not require any pre-treatment of the core's surface (CdS) to introduce any functional groups prior to shell growth. As synthesized CdS/ZnS core/shell nanowires (Fig. 25b) exhibited good photocatalytic activities for the degradation of 4-chlorophenol (4CP) (89% of 4CP was degraded in 12 h) under visible light compared to the activity of CdS nanowires (74% degradation in 12 h) and anatase-TiO₂ (30% degradation in 12 h of irradiation).²⁵⁷ Additionally, CdS/ZnS core/shell exhibited maximum photocatalytic activity for the degradation of MB dye under visible light irradiation (complete degradation in 6 h of irradiation) with respect to uncoated CdS nanowires (63% of degradation in 6 h) and anatase-TiO₂ (68% degradation in 6 h).²⁵⁷ Despite being a type-I heterojunction, the present CdS/ZnS core/shell system exhibited enhanced photocatalytic activity because of the passivation of surface states in the CdS cores by the ZnS shell, which acts as surface traps for the photogenerated charge carriers.²⁵⁷

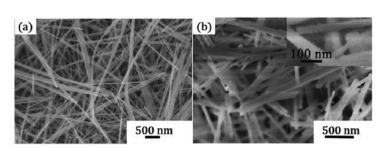
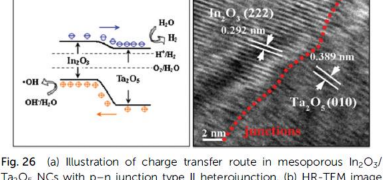


Fig. 25 (a) SEM image of CdS nanowires with 45 nm diameter. (b) SEM of CdS/ZnS core/shell NCs with 60 nm diameter and shell thickness of 5–10 nm. The inset figure depicts the surface of the shell, which consists of ZnS NPs.²⁵⁷ (a and b) (Reproduced with permission from ref. 257 OPEN ACCESS, Copyright 2018 Springer).

Mesoporous In₂O₃/Ta₂O₅ nanocomposite systems

Mesoporous tantalum oxide derived photocatalysts can exhibit fascinating mass transport properties. Intrinsically, they possess large surface, thin walls and good light harvesting capabilities. Potentially, they stand as ideal candidates for photocatalytic H₂ production from water splitting reactions. Takahara *et al.* synthesized mesoporous Ta_2O_5 through a ligand-assisted templating method, which unveils photocatalytic activity for overall water decomposition.²⁵⁸ In contrast to mesoporous Ta_2O_5 materials, highly crystallized Ta_2O_5 materials provided negligible H₂ production and no O₂ production. The high photocatalytic activity of the mesoporous Ta_2O_5 , where the rate of migration of photogenerated charge carriers is higher in mesoporous Ta_2O_5 than the migration occurring in highly crystallized Ta_2O_5 samples, from the lattice to the surface.²⁵⁸ Even though the mesoporous structure possesses such advantageous features, it is highly limited because of the collapse of the porous structure and the decrease of the specific surface area during the template removal process, due to its very low thermal stability.²⁵⁸ Such limitation could be resolved by the introduction of a second component into the mesoporous structure.^{259,260}

Xu et al. synthesized mesoporous heterostructured In₂O₃/Ta₂O₅ NCs via one-step sol-gel process using pluronic (P123), triblock copolymer surfactant, as structural directing agent. The incorporation of In₂O₃ (2.7 eV)²⁶¹ into the mesoporous Ta_2O_5 (3.9–4 eV)²⁶¹ resulted in the formation of a type-II system with reduced band gap energy, rendering the photocatalyst active in the visible region. Subsequently, the thermal stability of mesoporous Ta₂O₅ was greatly enhanced after the formation of the nanocomposite. The as synthesized mesoporous In₂O₃/Ta₂O₅ exhibited a relevant photocatalytic activity for H₂ generation (189 µmol h⁻¹ g⁻¹) from aqueous methanol solution under simulated solar irradiation as compared to the activity of non-porous In₂O₃/Ta₂O₅ NCs synthesized without template (54 μ mol h⁻¹ g⁻¹) and pure crystalline Ta₂O₅ (84 μ mol h⁻¹ g⁻¹).²⁶¹ Compared to non-mesoporous In₂O₃/Ta₂O₅ nanocomposites, mesoporous systems exhibited high activity due to their large specific surface area, which provides more active sites for H₂ production. The photogenerated electrons and holes have to travel short distance in mesoporous structure due to very thin walls, which is not observed in the other two photocatalysts.²⁶¹ The same authors further investigated the three dimensional mesoporous In_2O_3/Ta_2O_5 NC systems as a function of the wt% of In₂O₃ incorporation and the calcination temperature employed in the heat treatment.²⁶² A thin doping layer was found to be formed at the interface of the mesoporous In_2O_3/Ta_2O , due to the substitution of In^{3+} by Ta^{5+} , resulting in the formation of p-type conduction. Contrastingly the doping of Ta^{5+} in In_2O_3 results in the formation of n-type conduction.²⁶² A p-n junction type-II heterostructure is thus formed in this mesoporous composite system (Fig. 26a). The heterojunction area in the nanocomposite increased with an increasing amount of In₂O₃ (wt%).²⁶² The maximum area of heterojunction was observed for 20 wt% of In₂O₃, beyond which the heterojunction area decreased together with the photocatalytic performance in the materials. Regardless, the mesoporous nanocomposite with varying In_2O_3 (wt%) still provided a comparatively higher activity to that of pure crystalline Ta₂O₅ photocatalysts.²⁶² The mesoporous In_2O_3/Ta_2O_5 NCs with 20 wt% of In_2O_3 was synthesized at varying calcination temperature (450, 550, 650, 750 °C).²⁶² An increase in calcination temperature from 450 °C resulted in a decrease in specific surface area and pore volume, with a complete destruction of the mesoporous structure observed at 750 °C leading to the formation of crystallized NCs with very low photocatalytic activity (Fig. 26b).²⁶²



(b)

Fig. 26 (a) Illustration of charge transfer route in mesoporous ln_2O_3/Ta_2O_5 NCs with p-n junction type II heterojunction. (b) HR-TEM image of the interface region between ln_2O_3 and Ta_2O_5 materials in ln_2O_3/Ta_2O_5 NCs (with 20 wt% ln_2O_3) calcined at 750 °C.²⁶² (a and b) (Reproduced from ref. 262. Copyright 2018 Elsevier).

Results confirmed that the onset of destruction of the mesoporous structure started from 650 °C, below which the three dimensional novel structure was retained.²⁶²

To conclude, differing from other families of photocatalysts, the synthesis of nanocomposite generally takes place in two consecutive steps. The first step involves the synthesis of base SC materials as "support" and the second step involves the growth of a different SC material on the denoted support material. The transformation of two consecutive steps into a direct single step method is now under intensive research in the field of photocatalysis, in order to tune the interface of two different semiconductor materials to enhance the photocatalytic activity. Nanocomposite system fabrication can be considered as essential in the future design of advanced photocatalytic systems. Such design mostly involves surface passivation (elimination of trap states) of core and creation of type-II heterojunction for efficient charge separation. Table 4 summarizes a number of relevant photocatalytic nanocomposite systems with their synthetic approaches along with their photochemical applications. The diverse modified/unmodified SCs reported in the field of heterogeneous photocatalysis have been segregated and the synthetic development within each family was specifically rationalized.

Z-Scheme photocatalysts

1st generation Z-scheme photocatalytic systems

(a)

Heterogeneous photocatalysis for overall water splitting process can be achieved *via* two different systems, namely (1) a conventional type consisting of a single SC (modified/unmodified) material which carries out the overall water splitting to produce H_2 and O_2via one-step excitation upon exposure to photons (Fig. 27) and (2) two different photocatalytic active SCs activated *via* a two-step photoexcitation mechanism under light irradiation. The latter system is denoted as Z-scheme photocatalytic system as it mimics the naturally occurring Z-scheme process in photosynthesis of plants, where H_2O and CO_2 are converted to carbohydrates and O_2 under solar irradiation.

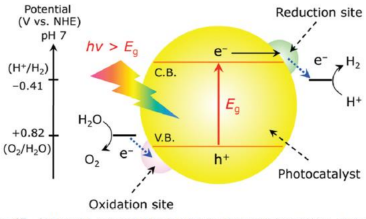


Fig. 27 Schematic representation of one-step photoexcitation system using single semiconductor material for overall water splitting²⁶⁶ (Reprint with permission from K. Maeda, K. Domen, Photocatalytic water splitting: recent progress and future challenges, *J. Phys. Chem. Lett.*, 2010, **1**, 2655–2661. Copyright 2019 American Chemical Society).

From the one-step photoexcitation system (Fig. 27), it is obvious that reduction and oxidation sites on the SC refers to the active sites generated by specifically co-catalysts loaded to perform the water redox reactions. Irrespective of the system employed for overall water splitting, the loading of co-catalysts (Pt, NiO_x or Rh) on the SC is the main pre-requisite to promote water splitting even in the presence of SCs able to perform overall water splitting without co-catalysts.²⁶⁷

The two-step photoexcitation system (Z-scheme) was first instigated and explored by Bard *et al.* in 1979²⁶⁸ in which a dual photocatalytic system was reported where one SC is used specifically for H₂ generation and the other one for O₂ generation, simultaneously, from water splitting. The utility of photons is more efficient in Z-scheme systems as the Gibbs free energy change required to activate the different SCs is reduced with respect to the energy required for a one-step photoexcitation system.²⁶⁸ This provides highly efficient photocatalysts with stronger oxidation and reduction capabilities on either side of the system.^{266,269} Upon irradiation to photons, both SCs are excited resulting in the generation of electrons and holes (Fig. 28). The photogenerated electrons remain on the H₂ photocatalyst with higher reduction potential and the holes remain on the O₂ photocatalyst with higher oxidation potential, resulting in the optimization of the redox potential of the system. The interaction between these two SCs takes place *via* transfer of electrons from the CB of the O₂ photocatalyst to VB of the H₂ photocatalyst through a pair of reversible redox mediators (A and D).²⁷⁰ The photocatalysis in such system takes place in the liquid phase and it is designated as liquid-phase Z-scheme photocatalytic system, belonging to the 1st generation of Z-scheme systems.^{271,272}

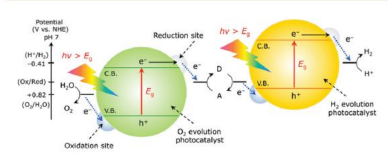


Fig. 28 Schematic depiction of Z-scheme two-step photoexcitation route for overall water splitting reaction; E_g is the energy band gap; *D* is the electron donating species and *A* is the electron accepting species.²⁷⁰ (reprint with permission from K. Maeda, K. Domen, Photocatalytic water splitting: recent progress and future challenges, *J. Phys. Chem. Lett.*, 2010, **1**, 2655–2661. Copyright 2019 American Chemical Society).

The main downside of this Z-scheme is the occurrence of the reduction of 'A' and oxidation of 'D' molecules on the surface of the photocatalyst where the oxidation and the reduction of water is taking place, respectively. Optimum selectivity control is the key factor for these systems due to which the number of liquid phase Z-scheme systems displaying optimum photocatalytic activity is highly limited.

Thermodynamically, the reaction of electron donor (A) and electron acceptor (D) with the photogenerated electrons in the CB of the H₂ photocatalyst and holes in the VB of the O₂ photocatalyst is more favourable to occur as compared to the water splitting reaction.²⁷³ These backward reactions must be suppressed in order to increase the effective number of photogenerated electrons and holes for water splitting reaction. Domen *et al.* reported a Z-scheme system consisting of Pt-loaded ZrO₂/TaON and Pt-loaded WO₃ as H₂ and O₂ photocatalysts respectively, with IO_3^{-}/I^{-} as redox mediator.²⁷⁴ In this system, the backward reactions with redox mediators were significantly minimized displaying a high quantum efficiency of 6.3% at 420 nm.²⁷⁴ The photocatalytic activity of the 1st generation Z-scheme system is predominantly dependent on the concentration of the redox mediators, ^{270,275} the side reactions of the redox system (*i.e.* pH-dependent of the reactant solution)²⁷⁶ and the interactions of the redox mediators with co-catalysts.²⁷⁷ Additionally, the redox mediators absorb the visible light and reduce the light absorption of the SCs. Besides, the liquid redox mediators suffer from lack of long-term stability and activity.

In order to eliminate the drawbacks associated with the above system and to render the Z-scheme photocatalysis viable for gas and solid-phase reaction, liquid redox mediated-free Z-scheme systems became an important prerequisite for an efficient system in the photocatalytic field. This led to the beginning of 2nd generation Z-scheme photocatalytic systems.

2nd generation Z-scheme photocatalytic systems

Z-Scheme systems without liquid redox mediators (A and D) and with spatial fixation of SCs form all-solid-state (ASS) Z-scheme photocatalytic systems which constitute 2nd generation Z-scheme systems. These systems' applications are viable for gas-phase as they eliminate the drawbacks of the 1st generation Z-scheme systems. Noble metal nanoparticles (Ag or Au) are generally employed as electron mediators for an efficient charge-carrier separation by transport through the interface between SC₂ and SC₁^{278,279} (Fig. 29). Primarily, these noble metal nanoparticles (Ag or Au) act as photosensitizers due to localized surface plasmon resonance (SPR) effect. This system was firstly introduced by Tada *et al.* which reported the design of anisotropic CdS–Au–TiO₂ nanojunctions by a facile photodeposition technique for the reduction of methylviolgen (MV²⁺).²⁸⁰ Both SCs (CdS and TiO₂) were photoexcited upon exposure to UV irradiation. The electrons from the CB of TiO₂ were transferred to Au then to the VB of CdS and subsequently recombined with photogenerated holes in the VB of CdS. This results in the generation of electrons and holes with high reduction and oxidation potential on CdS and TiO₂ photocatalyst, respectively.²⁸⁰ The vectorial

Chunping Xu, Prasaanth Ravi Anusuyadevi, Cyril Aymonier, Rafael Luque, Samuel Marre. Nanostructured materials for photocatalysis. Chemical

Society Reviews, Royal Society of Chemistry, 2019, 48 (14), pp.3868-3902. (10.1039/C9CS00102F). (hal-02185643)

electron transfer (TiO₂ \rightarrow Au \rightarrow CdS, similar to the electron pathway in Fig. 29) not only improved the photocatalytic activity but also the photostability of CdS SC.²⁸⁰

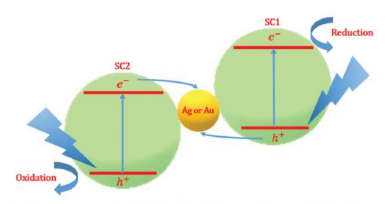


Fig. 29 Schematic depiction of charge carrier pathway in ASS-Z-scheme photocatalytic system (2nd generation Z-scheme photocatalytic systems).

Cheng *et al.* synthesized Ag₃PO₄/Ag/SiC ASS Z-scheme photocatalysts by a simple *in situ* precipitation technique at room temperature.²⁸¹ As synthesized photocatalysts with Ag NPs as charge transmission bridges (electron mediators) exhibited a similar vectorial electron transfer (Ag₃PO₄ \rightarrow Ag \rightarrow SiC) as compared to that reported in the previous system and provided a higher photocatalytic activity for the degradation methyl orange and phenol under visible light irradiation with respect to Ag₃PO₄/Ag and SiC/Ag nanocomposites due to the efficient separation of electron–hole pairs in the Z-scheme system.²⁸¹ The localized surface plasmon resonance (SPR) effect of Au/Ag on these systems led to the development of plasmonic ASS Z-scheme photocatalytic systems under visible light irradiation.^{282–}²⁸⁴

Cheng *et al.* synthesized plasmonic Z-scheme α/β -Bi₂O₃–Ag–AgCl photocatalysts utilising a deposition–precipitation technique coupled with a photo-reduction process.²⁸⁵ In the case of this plasmonic photocatalyst, only Bi₂O₃ (*E*_g of α -Bi₂O₃ = 2.71 eV and β -Bi₂O₃ = 2.48 eV) was excited, while AgCl (3.25 eV) is not activated under visible light irradiation ($\lambda > 420$ nm). At the same time, the photogenerated electron–hole pairs appeared in Ag NPs due to the plasmonic effect. Here, the vectorial electron transfer was reported to take place from the CB of Ag NPs to the CB of AgCl, where the reduction reaction occurs and the photogenerated electrons from the CB of Bi₂O₃ recombined with the plasmon-induced holes of Ag NPs.²⁸⁵ α/β -Bi₂O₃–Ag–AgCl exhibited enhanced photocatalytic activity as compared to α/β -Bi₂O₃ nanowires for the degradation of Rhodamine B and acid orange 7 dyes. The higher photodegradation activity was attributed to the hierarchical nanostructures with large surface area, the profuse interfacial active sites, the enhanced absorption of visible light due to SPR effect of Ag NPs and the efficient separation of electron–hole pairs.²⁸⁵

In essence, only one SC is generally activated in plasmonic Z-scheme systems, while the SPR effect of the noble metal NPs always plays a significant role. When both SCs are excited, the noble metal particles primarily act as electron mediators. However, ASS Z-scheme systems are composed of costly noble metals which also reduce the light absorption capability of SCs. These drawbacks have limited their large scale implementation and alternatives for redox mediators-free Z-scheme systems have been sought, resulting in the development of third-generation Z-scheme systems, detailed hereafter.

3rd generation Z-scheme photocatalytic systems

3rd generation of Z-scheme photocatalysts (direct Z-scheme photocatalysts) do not require any redox medium for charge migration, generally comprising two different SCs. Structurally, a direct Z-scheme photocatalyst is analogous

to a type-II SC–SC coupled heterojunction system (see section on nanocomposites), but the charge-carrier migration mechanism is rather different to that disclosed in type-II systems.

In a classical type-II heterojunction photocatalyst, the CB and the VB levels of SC₂ are comparatively higher to the CB and VB levels of SC₁. Upon exposure to photons irradiation, the photogenerated electrons transfer from the CB of SC₂ to the CB of SC₁, while the holes move from the VB of SC₁ to the VB of SC₂ (Fig. 30b). This results in a spatial separation of electron–hole pairs. However, the electrons and holes accumulated in the CB of SC₁ and the VB of SC₂, respectively, possess low reduction and oxidation potentials. In this regard, the redox ability of the overall photocatalyst is largely reduced.

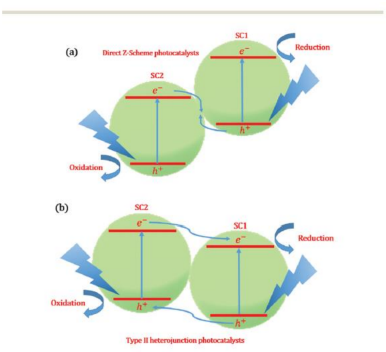


Fig. 30 (a) Schematic illustration of charge carrier separation in direct Z-scheme photocatalytic system (3rd generation Z-scheme system). (b) Charge carrier migration mechanism in type-II heterojunction photocatalysts.

In contrast, the photogenerated electrons in SC_2 with low reduction potential combine with the photogenerated holes in the VB of SC_1 with low oxidation potential in direct Z-scheme photocatalytic systems. This results in the accumulation of electrons and holes with higher reduction and oxidation potentials, respectively, in SC_1 and SC_2 (Fig. 30a). The spatial separation of charge carriers takes place in direct Z-scheme photocatalysts, facilitating the system to achieve optimum redox capabilities.^{286–288}

The charge carrier's migration mechanism in direct Z-scheme is homogeneous (similarly to both ASS Z-scheme and liquid-phase Z-scheme systems) but in the absence or need for any redox mediator (liquid or solid NPs). Direct Z-scheme systems hence avoid the backward reactions taking place in 1st generation Z-scheme photocatalysts. Additionally, direct Z-scheme systems facilitate the migration of the charge carriers directly across the interface of two SCs, thereby reducing the transformation distance with respect to 2nd generation Z-scheme photocatalysts, eventually ending up in avoiding unfavourable reactions. Direct Z-scheme systems also avoid the light shielding effect of noble metal NPs-based redox mediators.

Due to structural similarities between Z-scheme and type-II heterojunctions photocatalysts, the charge transfer mechanism in SC–SC coupled systems has been investigated in detail to differentiate between direct Z-scheme and type-II heterojunctions. Currently, several characterization techniques have been employed to obtain deep insights

into the charge carrier migration process, namely internal electric field simulation, effective-mass calculation, X-ray photoelectron spectroscopy, radical species trapping test and photocatalytic reduction tests.²⁷²

Direct Z-scheme photocatalysts were firstly reported by Yu *et al.* by demonstrating the high photocatalytic activity of g-C₃N₄/TiO₂ nanocomposites under UV irradiation for the degradation of formaldehyde in air.²⁸⁹ The contact interface between SCs (g-C₃N₄ and TiO₂) was directly achieved by coupling g-C₃N₄ with TiO₂ NPs by calcination (550 °C for 60 min) of mixtures of TiO₂ NPs and urea. After this seminal study, additional research efforts were reported on direct Z-scheme photocatalysts by several researchers across the globe. Li *et al.* employed a similar calcination technique (600 °C, 120 min) on the mixture of bulk g-C₃N₄ and nanotube titanic acid (NTA) to successfully synthesize visible light active direct Z-scheme g-C₃N₄/TiO₂ nanocomposites.²⁹⁰ During thermal treatment, bulk g-C₃N₄ was oxidized to g-C₃N₄ nanosheets and NTA got dehydrated, resulting in the formation of novel-anatase TiO₂ NPs. The thin nanosheets of g-C₃N₄ provided high surface area and short bulk diffusion length for the spatial separation of charge carriers.²⁹⁰ The visible light activity originated from the doping of TiO₂ NPs during the heat treatment. As synthesized direct Z-scheme photocatalyst 30%-g-C₃N₄/TiO₂ (prepared from the precursor mixture containing 30 wt% of bulk g-C₃N₄) exhibited high activity for the photo-oxidation of propylene under visible light ($\lambda \ge 420$ nm). This activity was attributed to the large surface area of the nanocomposite, the effective spatial separation of charge carriers and the morphology of photocatalysts (conjunction of thin nanosheets g-C₃N₄ and TiO₂ nanoparticles).²⁹⁰

The contact interfaces between two SCs described in the above two cases were established *via* chemical interactions, which resulted in firm interfaces. A similar type of intimate interface contact has been also achieved and reported in two-dimensional (2D) composites consisting of 2D/2D g-C₃N₄/MnO₂ heterostructured direct Z-scheme photocatalysts, directly synthesized *via* wet-chemical approach.²⁹¹ As-synthesized photocatalysts exhibited higher photocatalytic activity for the degradation of phenol and rhodamine B dye as compared to individual g-C₃N₄ and MnO₂ photocatalysts.²⁹¹ The interface formation between two different SCs in direct Z-scheme photocatalytic system can also be accomplished *via* physical interactions based on the electrostatic attractions generated by opposite charges.

Xu *et al.* constructed 2D/2D α -Fe₂O₃/g-C₃N₄ direct Z-scheme photocatalysts *via* electrostatic self-assembling of g-C₃N₄ nanosheets with hematite α -Fe₂O₃ nanoplates.²⁹² The resultant α -Fe₂O₃/g-C₃N₄ (10 wt% α -Fe₂O₃) heterostructures possessed intimate contact interfaces and displayed 13 times higher H₂ evolution rate from water containing triethanolamine (hole scavenger) with respect to g-C₃N₄ photocatalysts under visible light irradiation.²⁹² This high activity was attributed to the firm interface achieved through physical contact, where spatial separation charge carries occurred through Z-scheme mechanism.

Direct Z-scheme photocatalysts can provide three relevant advantages including (1) an efficient spatial separation of charge carriers upon excitation, (2) the generation (accumulation) of charge carriers with high redox ability and (3) and affordable and potentially cost-competitive fabrication cost. As a result, these materials display high photocatalytic activities for a wide range of applications (water splitting, degradation of pollutants, *etc.*).

With regards to the current development of direct Z-scheme photocatalytic systems, the construction of 2D/2D heterostructures is highly preferred due to the generation of contact interfaces between SCs over a large surface. Very recently, Hu *et al.* constructed a 2D/2D heterojunction of black phosphorus (BP)/Bi₂WO₆ (bismuth tungstate) monolayer nanosheets by the decoration of sandwich structure-like Bi₂WO₆ on the surface of BP nanoflakes through a simple wet chemical approach.¹²⁵ The resultant nanocomposite material possessed intimate interfaces in contact and a homogeneous distribution of Bi₂WO₆ nanosheets on the BP surface. 12 wt% BP/Bi₂WO₆ nanocomposite displayed an outstanding photocatalytic rate for H₂ evolution (21 042 μ mol g⁻¹) from water splitting and removal nitric oxide (up to 67%), comparably 9.15 times and 2.15 times to that of monolayer BP nanosheets. The high photocatalytic activity was attributed to the novel heterojunction interfaces established between monolayers of Bi₂WO₆ nanosheets and BP flakes, enhanced light absorption and the efficient separation of charge carriers.¹²⁵

Miscellaneous

Apart from the above discussed four categories of photocatalysts, there has been recently an increasing attention on piezoelectric semiconductors in photocatalysis.^{293–297} Conventionally, the term "piezoelectric" represents a group of materials (i.e. certain metal oxides) featuring non-centrosymmetric structures, that is, materials which do not possess centre of symmetry in their crystal structure. Such material possess the ability to enhance the rate of photochemical processes upon mechanical deformation by exhibiting spontaneous polarization by the induction of an electric field throughout the crystal. Due to such polarization potential, free charge carriers or photogenerated electrons or holes are spatially separated resulting in the observed enhancement in photocatalytic activity. Piezophotocatalysis (based on piezoelectric semiconductor photocatalysts) involves the synergy of photonic energy and mechanical energy to achieve enhanced photocatalytic rates.²⁹⁷ Within the perspective of synthetic approaches, such piezophotocatalysts can be designed via tailored synthesis of pure semiconductor possessing both piezoelectric and photonic properties and hybrid nanocomposites involving the conjunction of two different materials featuring piezo and photonic properties. The former case includes some recently reported materials including zinc tin oxide (ZTO–ZnSnO₃) nanowire arrays,²⁹⁸ ZnO nanowires,²⁹⁹ sodium niobate (NaNbO₃) and nanostructured films.³⁰⁰ The latter case involves hybrid system such as barium titanate/silver phosphate (BaTiO₃/Ag₃PO₄) nanocomposites where the piezo properties of $BaTiO_3$ is combined with the photochemical properties of Ag_3PO_4 , resulting in novel nanocomposite piezophotocatalysts.³⁰¹

In most of the cases, the mechanical energy required for piezophotocatalysis is provided by ultrasonic irradiation. The pros of such mechanical forces relies in the generation of periodic compression stress, which prevents the screening of piezoelectric generated charges by photogenerated charge carriers.³⁰² Even though these novel piezophotocatalysts provide enhanced separation of charge carriers leading to enhanced photocatalytic activity, the photocatalytic applications of piezophotocatalysts are mostly limited to date to pollutant removal reactions and not extended – for instance – to CO₂ reduction, novel organic molecules synthesis, *etc.* The field of piezophotocatalysis is still in its infancy and needs intensive fundamental research and engineering efforts to unleash its full potential.

Mechanical forces provided by ultrasonic waves and/or reactive extrusion (*e.g.* mechanochemistry) can result in the generation of defects on the surface of semiconductor materials.³⁰³ Nevertheless, there is still a need for more fundamental studies in such processes for elucidating the reasons behind the enhanced activity and the relationship between the mechanical force and the generated defects. Additionally, the development of reactor systems for an optimum input of mechanical energy (ultrasonic vibration, reactive extrusion-mechanochemistry-) and light energy is still far from being practical, as currently, no continuous flow photoreactor system involving both energies simultaneously has been reported to date.

Challenges & future prospects

Challenges for continuous synthesis of nanophotocatalysts

Concerning the four different families of photocatalytic materials, continuous synthetic processes have been successfully reported for conventional pure semiconductor photocatalysts (TiO₂ and ZnO). Several studies are ongoing to improve the yield of the reported process and concurrently new synthetic techniques using supercritical medium are rapidly reported for synthesizing unconventional photocatalysts (GaN and CeO₂) by several research groups. In the perspective of ternary SCs, synthetic explorations (continuous flow synthesis) has not spurred as seen in binary photocatalysts. Currently, the implementation of single source precursor and facilely operated flow systems has been studied for continuous synthesis of ternary SCs. Regarding solid solution photocatalysts, tremendous research is carried out in reducing the cost of single source precursors by elucidating or creating new synthetic strategies. Continuous synthetic shift from batch to flow, for solid solution lies completely in the dormant phase because of the non-availability of ready-made, stable and oxidation free single source precursors. Finally, for type-II and direct Z-scheme photocatalysts continuous consecutive synthesis of core (base SC) and shell using a dual reactor set up is the current area of focus.

Implementation of photocatalysts in heterogeneous photocatalysis

Heterogeneous photocatalysis involves a multiphase system involving the interaction between different phases (solid–gas, solid–liquid and solid–liquid–gas) through mass transfer (adsorption) upon mixing and charge transfer (redox reactions) upon exposure to light. Therefore, the design of a reactor for heterogeneous photocatalysis extensively depends on the photochemistry to which the system is subjected and the source of electromagnetic radiations. The design of the photoreactor is always targeted to achieve high photocatalytic activity by ensuring efficient illumination of the solid photocatalytic SC along with the synchronous enhancement of both mass transfer rate from photocatalysts surface and heat transfer rate from the reaction vessel. The composition and chemistry of phases involved in such systems determine the compatible materials for the construction of the photoreactor.^{303–308} The different multiphasic systems employed in heterogeneous photocatalysis are: (i) the solid-gas (organic/inorganic) system, (ii) the solid-liquid system and (iii) the solid-liquid-gas system. Broadly the photoreactors studied are of two types: batch and flow reactors. The batch reaction vessel used in this multiphasic systems are made up of either quartz, pyrex and sometimes stainless steel materials. The position of the irradiation source can be either internal or external irradiation.^{12,24,64,297} The effective utilization of photocatalysts within batch and flow reactors are achieved by two different methods: suspension and immobilization. In the former system, the photocatalytic material is suspended in the fluid (especially liquid) phase with the help of magnetic stirring, toroidal or non-toroidal agitation promoted by gas molecules. This suspension of the photocatalytic materials in the liquid phase (aqueous or non-aqueous) is called "slurry system". In the immobilized system, the photocatalysts are bonded to inert surfaces, either through a physical or a chemical bonding.

Strategies for efficient implementation of photocatalysts in flow reactors

Flow reactors have been developed in the last two decades due to the attainment of higher throughput, enhanced mass and photon transfer. Comparatively, flow photoreactors provide reduction of safety hazards and simplify operation compared to batch reactors.^{309,310} Flow systems based on solid–gas (diphasic) or solid–fluid (liquid and/or gas) (diphasic or triphasic) initially progressed in the direction of immobilized system due to a simple separation of photocatalysts from the reactive mixture. The solid photocatalysts immobilized/mobilized on glass beads could be packed into Pyrex or borosilicate/glass tube in case of packed bed reactors (PBR)^{310,311} or successfully fluidized in the annular space between two tubes in fluidized bed reactors³¹² (Fig. 31a). Solid photocatalysts could be perfectly immobilized through various techniques on the walls of the reactor (see Fig. 31b), in case of an annular reactor, ^{313–315} honeycomb monolith reactors²⁹⁴ and uniformly coated or dispersed in plate reactor.^{316,317} Behnajady *et al.* reported a continuous flow tubular photocatalytic reactor with immobilized photocatalysts for the degradation of acid red 27 (AR27) (an anionic monoazo dye).³¹⁸ Commercially available TiO₂ (Degussa P25) was herein immobilized on a glass plate and inserted inside the tubular reactor. The immobilization was carried out using a heat attachment methodology.

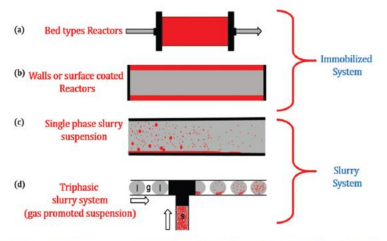


Fig. 31 Strategies applied for the efficient implementation of photocatalysts in flow reactors redrawn, based on the artice.³²⁵ (Reproduced with permission from ref. 325. Copyright 2018 John Wiley and Sons).

Li *et al.* also reported the use of double cylindrical shell (DCS) photoreactor with an immobilized monolayer of TiO₂ coated on silica gel beads.³¹⁹ The design of this photoreactor was similar to the annular reactor. Another relevant reported immobilized system was denoted as labyrinth bubble photoreactor, studied by Hao *et al.* in which TiO₂ photocatalysts was immobilized on quartz glass particles and a labyrinth fluid flow arrangement was achieved by the arrangement of parallel baffle plates within the reaction vessel.³²⁰

Effective utilization of photons under flow conditions can be achieved *via* optimum positioning of the irradiation source either internally within the reaction vessel (annular and PBR system) or externally with respect to the reaction vessel (plate reactor or fluidized bed reactor).

In both solid–liquid (diphasic) and solid–liquid–gas (triphasic) systems, slurry reactors are highly advantageous as compared to immobilized photocatalytic reactor systems due to enhanced higher photocatalytic activity. This enhanced behaviour of slurry systems is due to the intrinsic furnishing of high total surface area of photocatalysts per unit volume. The implementation of immobilized systems at large scale still remains a significant challenge. The complex design of the irradiation source, according to the requisite length of the upscale system has limited research from developing further these systems. Even though the slurry system is highly economical in terms of fabrication and operation, its major and only limitation is the requirement for downstream separation of the photocatalytic materials from the product/liquid stream.³²¹

Priya and Kanmani developed a two different novel batch slurry (BG) and batch slurry recycle reactors fitted with continuous supply of inert gas (BRRwCG) for water splitting reaction under solar irradiation.³²² Both reactor materials were made up of plexiglas material, transparent to the solar spectrum wavelength (290–700 nm). The photocatalytic material was successfully suspended in BG through magnetic stirring. Analogously, the suspension was achieved in BRRwCG through continuous bubbling of the reactant solution by an inert gas. Simultaneously, a part of the reactant solution was recirculated using a peristaltic pump (Fig. 31d). The catalyst composition in both reactors was kept identical and it was found that BRRwCG exposed higher photocatalytic activity as compared to BG due to the enhanced mass transfer and photon distribution.³²² Slurry rectors with continuous single phase (no gas promoted agitation, Fig. 31c) suffer from severe limitations of photon and mass transfer when they are upscaled to achieve higher throughput.

McCullagh *et al.* developed a novel continuous flow slurry reactor for the degradation of MB dye under UV irradiation.³²³ This novel system coupled the usage of 'Drum Reactor' design and pelletized titania photocatalytic particles. The commercially available Hombikat pellet titania catalyst was employed which avoids any downstream processing to separate the photocatalysts from the liquid phase.

Challenges for effective flow photoreactors

Research for novel photoreactors and effective utilization of photocatalysts has pushed scientists more into microreactor fabrication and design. These microreactors provide enhanced mass transfer compared to the conventionally mentioned large scale photocatalytic reactors. Due to high surface-to-volume ratio achieved in this system, as a result of their small dimensions.^{324–326} Simultaneously, they provide the opportunity for low catalyst loading, enhanced heat transfers and homogeneous photon distribution.^{326–328}

Concerning slurry reactors, there is currently a rapid research effort going on to replace the mechanically attained suspension by air or reactive gas agitation attained suspension. These systems are currently given preference because of the rapid development of microreactors which leads to low catalyst loading. Similarly, advanced progress in downstream separation equipment have also motivated researchers to switch towards continuous flow slurry reactors without any immobilization. As example, Rakhshaee *et al.* designed a plug flow reactor (PFR) in which nanohematite (α -Fe₂O₃) photocatalysts were successfully suspended in the liquid phase with air promoted suspension.³²⁹

Conclusions

Several modified/unmodified SCs studied to date in heterogeneous photocatalytic field have been described in this work. These highly efficient photocatalysts were classified into four different categories depending on their composition, band gap engineering and charge carrier migration mechanisms. Currently developed synthetic strategies for each photocatalyst have been reviewed along with the rationale behind the new synthetic approach and its advantages over conventional synthetic routes. Their photocatalytic performance was also discussed and compared with conventional systems, finding significant advantages in the interface engineered systems as compared to traditional photocatalysts. Importantly, photoreactor systems (nature of photons source used and type of reaction vessel, types of reactor) were also briefly covered in the future prospects section, with a particular emphasis on flow photochemical processes and reactors, believed to the way forward in the field. We hope the proposed overview can provide a comprehensive compilation of key studies to progress the state-of-the-art in the field and look forward to additional advances in the design of advanced versatile (nano)photocatalysts in the future with several relevant applications.

Conflicts of interest

The authors declare no conflict of interest.

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Footnote

⁺ THESE TWO AUTHORS EQUALLY CONTRIBUTED TO THIS WORK.

PC	Synthetic methods	Precursors	Size of PC	Photochemical reaction	Illumination source	Photoreactor	Ref.
TIO_2	Flame spray pyrolysis	C ₁₂ H ₂₈ O ₄ Ti, C ₈ H ₁₀ and 11–21 nm (3.1–3.3 eV) C ₅ H ₁ N.	11-21 nm (3.1-3.3 eV)	Mineralization of sucrose	Blue lamp (15 W), $(\lambda = 300-400 \text{ nm})$	Spiral slurry flow photoreactor	6
ZnO	Precipitation and calcination	ZnC ₄ H ₆ O ₄ , C _{2n} H _{4n+2} O _{n+1} and (NH ₄) ₅ CO ₃ .	20 nm	Degradation of methyl orange (MO)	UV lamp $(\lambda = 250-400 \text{ nm})$	Hollow cylindrical batch reactor	16
ZrO ₂	Hydrothermal and calcination yielding nanoflowers of ZrO ₂	Zr(SO ₄) ₂ and CH ₃ COONa	$Zr(SO_4)_2$ and CH_3COONa 400 nm and petal thickness about 20 nm.	Degradation of rhodamine B (RhB)	Degradation of rhodamine B 300 W high pressure mercury (RhB) lamp $(\lambda > 365 \text{ nm})$	Cylindrical vessel, with inner irradiation system	12
CeO ₂		CeCl ₂ and NH ₄ OH	15-36 nm (4.16 eV)	Degradation of amido black and acridine orange	250 W high pressure mercury lamp (UV range)	Glass vessel	18
ZnS	Thermal decomposition route	ZnC ₄ H ₆ O ₄ , CH ₄ N ₂ S, C ₂ H ₆ O ₂ and (C ₆ H ₉ NO) ₃₇	60 nm (3.6 eV)	Photodegradation of eosin B	125 W high pressure mercury lamp	Cylindrical pyrex cell	24
cdS	Thermal sulfidation process	Cd(CH ₃ COO) ₂ and dry H ₂ S gas.	100-300 nm (2.38 eV)	Hydrogen evolution from water splitting	350 W xenon lamp (λ > 430 nm)	Pyrex vessel with side irradiation	28
dSe	CdSe Solvothermal method	Cd ₂ Cl ₂ O ₈ , C ₂ H ₆ OS and NaHSe.	1.75-4.81 nm (3.11-1.89 eV)	Hydrogen production from water splitting	300 W xenon arc lamp $(\lambda > 320 \text{ nm})$	Borosilicate flask	31
GaN	Metal organic chemical vapour Ga(CH ₃) ₃ and NH ₃ deposition yielding nanowires	Ga(CH ₃) ₃ and NH ₃	20-50 nm in diameter, 4-6 μm in length. (3.4 eV)	Decolouration of orange II	200 W mercury lamp $(\lambda \ge 300 \text{ nm})$	Batch reactor	34
a ₃ N ₅	Ta ₃ N ₅ Alkali metal treatment coupled with ammonolysis	Ta ₂ O ₅ , NaCl, NH ₃ , KCl and K ₂ CO ₃	Average size: 80 nm (2.1 eV)	Average size: 80 nm (2.1 eV) Oxygen evolution from water 300 W xenon lamp visible oxidation $(\lambda > 420 \text{ nm})$	300 W xenon lamp visible $(\lambda > 420 \text{ nm})$	Pyrex reactor with top irradiation type arrangement	40
CuS	Chemical dealloying method yielding nanoporous globular olusters	Ti-Cu amorphous ribbon and H ₂ SO ₄	Ti-Cu amorphous ribbon Clusters of 500 nm size and Degeneration of methylene and $\rm H_2SO_4$ 40 nm thickness (1.6-1.7 eV) blue (MB) and MO	Degeneration of methylene blue (MB) and MO	500 W xenon lamp, illumination Quartz cell intensity was 0.01 W cm ⁻²	Quartz cell	27

Table 2	Summary of nano-ternary s	emiconductor photocatalysts	' synthetic strategies with the b	Table 2 Summary of nano-ternary semiconductor photocatalysts' synthetic strategies with the background of photochemical reactions studied. TS: photocatalyst (ternary semiconductor)	ictions studied. TS: photocatalyst	(ternary semiconductor)	1
IS	Synthetic methods	Precursors	Size of TS	Photochemical reaction	Illumination source	Photoreactor	Ref.
SrTiO ₃	Hydrothermal method	[CH ₃ CH(O-)CO ₂ NH ₄] ₂ Ti(OH) ₂ [Sr(OH) ₂ ·8H ₂ O] and NaOH	20-40 nm in diameter (3.2 eV)	Degradation of nitric oxide (NO) gas	UV light source: 450 W high pressure mercury lamp	Continuous flow sealed opaque reactor with olass plate	99
LiTaO ₃	Hydrothermal (nanorods) and solvothermal (nanoparticles (NPs)) methods	Hydrothermal (nanorods) TaCl ₅ , LiOH-H ₂ O, H ₂ O ₂ , NH ₃ , and solvothermal CH ₃ OH and C ₃ H ₆ O (nanoparticles (NPs))	Nanorods 20 nm diameter and length of 2-10 µm. NPs of 10-50 nm (4.6-4.7 eV)	Water splitting reaction	450 W high pressure mercury lamp, UV light source	up of quartz	69
NaTaO ₃	Hydrothermal method	Ta ₂ O ₃ , NaOH and deionized About 200 nm (3.96 eV) water		Decomposition of gaseous formaldehyde and RhB	8 W bactericidal lamp, $(\lambda > 254 \text{ nm}, 2.5 \text{ mW cm}^{-2})$	Cylindrical quartz cell	77
KTaO ₃	Solvothermal method	KOH, Ta ₂ O ₅ , C ₆ H ₁₄ , H ₂ O and C ₂ H ₂ OH	15-50 nm (3.7 eV)		300 W xenon arc lamp $(\lambda = 280-350 \text{ nm})$	Borosilicate glass reactor	84
AgNbO ₃	AgNbO ₃ Molten salt flux synthesis Ag ₂ O, Nb ₂ O ₅ and Na ₂ SO ₄ method	Ag ₂ O, Nb ₂ O ₅ and Na ₂ SO ₄	100-5000 nm (2.75-2.81 eV)	roduction from aqueous anol solution	arc-lamp	Outer irradiation type fused-silica reaction cell	89
CaIn ₂ O ₄	CaIn ₂ O ₄ Combustion synthesis	$Ca(NO_3)_2$, InN_3O_9 and $C_2H_5NO_2$	Approximately 90 nm	of MB, toluene water	out meet (< > +20 mm) 300 W xenon arc lamp fitted with UV cut off filter	with 90 mill volume	97
or AoGaO.	Cation exchange method	Cation exchange method NaGaO ₂ , AgNO ₃ and KNO ₃	200–1000 nm (2.4 eV)	decomposition Decomposition of isopropanol Xenon lamp with glass filters (IPA) under UV or visible light.	Xenon lamp with glass filters	Glass vessel with a quartz lid	108
BIVO4	Hydrothermal method	NH4VO ₃ , Bi(NO ₃) ₃ ·5H ₂ O and HNO ₃	49-490 nm (2.39-2.47 eV)		300 W xenon lamp with cut off filter for visible light [2] > 420 nm]	Closed gas circulation system fitted with pyrex made reaction cell	116
AgInS ₂	Co-precipitation strategy AgNO ₃ , C ₂ H ₆ O ₂ , Na ₂ SO ₄ , C ₃ H ₂ OH and InCl ₂ 4H ₃ O	AgNO ₃ , C ₂ H ₆ O ₂ , Na ₂ SO ₄ , C ₅ H ₅ OH and InCl ₅ 4H ₅ O	Avg. pore size 14.79 nm (1.61 eV)	Degradation of 2-nitrophenol (2NP) under visible irradiation.	$(\lambda > 420 \text{ nm})$	Batch vessel with magnetic 149 stirring	149
CuGaS ₂	Colloidal synthesis method using hot injection technique	Cu(S ₂ CNEt ₂) ₂ and Ga(S ₂ CNEt ₂) ₃	s with length 9 nm and width 8 nm (2.52 eV)		500 W halolite lamp $(\lambda > 430 \text{ nm})$	Glass vial	152

SS	Synthetic methods	Precursors	Photochemical reaction	Illumination source	Photoreactor	Ref.
β -AgAl _{1-x} Ga _x O ₂ ($0 \le x \le 1$) (2.8-2.19 eV)	Cation exchange method	CH ₃ COONa, Al(NO ₃) ₃ ·9H ₂ O, Ga(NO ₃) ₃ · <i>n</i> H ₂ O and AgNO ₃	Decomposition of IPA gas, first into acetone then to CO.	300 W xenon arc lamp and blue LEDs (400 nm $< \lambda <$ 500 nm)	500 mL reactor equipped with a pyrex glass lid window	159
$(\text{AgNbO}_3)_{1-x}(\text{NaNbO}_3)_x$ $(0 \le x \le 1)$ (3.4-2.79 eV)	Solid state reaction method $~~Ag_{2}O,Na_{2}CO_{3}$ and $Nb_{2}O_{5}$	Ag ₂ O, Na ₂ CO ₃ and Nb ₂ O ₅	Decomposition of IPA into acetone and CO ₂ under visible light	Blue-LEDs (BLEDs) with light intensity of 0.01 mW cm^{-2} ($\lambda = 400-550$ nm)	Glass reactor of volume 500 mL	163
$(Ga_{1-x}Zn_x)(N_{1-x}O_x)$ with and without RuO_2 loading (2, 6-2, 8 eV)	Nitridation of powdered precursor mixture	Ga ₂ O ₃ , ZnO and NH ₃	Water splitting under visible light	High pressure mercury lamp 450 W	Pyrex inner irradiation type reaction vessel	165
$\begin{array}{l} \mathop{\rm Ag}_{1-x} {\rm Sr}_x({\rm Nb03})_{1-x}({\rm TiO}_3)_x \\ (0 \le x \le 1) \ (3.2 - 2.7 \ {\rm eV}) \end{array}$	Solid state reaction method	Ag ₂ O, SrCO ₃ , Nb ₂ O ₅ and TiO ₂	O ₂ evolution from AgNO ₃ solution and decomposition of IPA	300 W xenon arc lamp with cut off filter $(\lambda \ge 410 \text{ nm})$	Closed gas circulation system containing pyrex vessel, with a flat side window for light irradiation	168
SrTiO ₃ -Ba ₂ FeNbO ₆ - n ($n = 0.25$, 0.5 and 1) (3.31-2.29 eV)	One-step molten salt route	TiO ₂ , BaCO ₃ , SrCO ₃ , Nb ₂ O ₅ , Fe ₂ O ₃ and NaCl	H ₂ evolution from aqueous methanol solution	300 W xenon lamp with cut off filter $(\lambda \ge 420 \text{ nm})$	Closed gas circulation system fixed batch reactor	171
$Cd_xZn_{1-x}S$ (0 < x < 1) (2.3-3.10 eV)	Thermolysis of zinc- cadmium-thiourea (Zn-Cd-Tu) complex	CH ₄ N ₂ S, C ₄ H ₆ O ₄ Zn·2H ₂ O and C ₄ H ₁₀ CdO ₆	H ₂ production from aqueous solution with sacrificial agents	350 W xenon arc lamp with UV cut off filter $(\lambda \ge$ 400 nm)	10 mL pyrex flask, with openings of flask are sealed with silicone rubber septum	184
Cd _x Cu _y Zn _{1-x-y} S Cd _{0.1} Cu _{0.01} Z _{0.89} S (2.33 eV)	Co-precipitation method	Zn(NO ₃) ₂ , Cu(NO ₃) ₂ , Cd(NO ₃) ₂ and Na ₂ S	H ₂ from aqueous solution with sacrificial agents	350 W xenon lamp with optical filter $(\lambda \ge 430 \text{ nm})$	Side window fixed pyrex cell	317
$(\operatorname{Agfn})_{x} \operatorname{Zn}_{2(1-x)} S_{2}$ $(0 \le x \le 1)$ (3.55-1.8 eV)	Heat treatment of Ag-In-Zn sulfide complex precursor	AgNO ₃ , In(NO ₃).3.9H ₂ O and Zn(NO ₃) ₂ .6H ₂ O	H ₂ evolution from aqueous solution with sacrificial agents	Solar Simulator (AM 1.5) and 300 W xenon lamp	Gas closed circulation system with a side or top window pyrex cell.	189
$(Culn)_xZn_{2(1-x)}S_2$ $(0 \le x \le 1)$ (3.55-1.8 eV) Pt or Ru loaded as co-catalyst on the solid solution	Heat treatment of Cu-In- Zn sulfide complex precur- sor and photo deposition for co-catalysts loading	Zn(NO ₃)·6H ₂ O, In(NO ₃)·3.6H ₂ O, CuCl ₂ H ₂ PtCl ₆ ·6H ₂ O and RuCl ₃ · <i>n</i> H ₂ O	H ₂ evolution form aqueous solution with sacrificial agents	300 W xenon lamp with a cut off filter ($\lambda \ge 420 \text{ nm}$)	Gas closed circulation system fitted pyrex cell with window for illumination fixed to the sides of the cell	190
ZnS-CuInS ₂ -AgInS ₂ Ru co-catalyst	Heat treatment of single source complex sulfide precursor. Loaded by photo-deposition method	Zn(NO ₃)·6H ₂ O, AgNO ₃ , In(NO ₃)·3.6H ₂ O and CuCl ₂ ·2H ₂ O (NH ₄) ₂ RuCl ₆	H ₂ evolution from an aqueous solution containing sacrificial agents.	300 W xenon lamp with cut off filter ($\lambda \ge 420 \text{ nm}$)	Closed gas circulation fixed pyrex cell	138

ocatalytic applications and their phot of solid solution (SS) photocatalysts synthesis the ed in Tue ditions methodologies cunthatin Overview of the Table 3

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N	Synthetic methods	Precursors	Size of the nanomaterials	Photochemical reaction Illumination source	Illumination source	Photoreactor	Ref.
TiO ₂ nanorods	Solvothermal method	C ₁₆ H ₄₀ O ₄ Ti, C ₁₈ H ₃₄ O ₂ , C H NO and C H OH	Nanorods: 10-40 nm	H ₂ generation from an	320 W xenon lamp with	Septum sealed gas vials 206	206
TiO ₂ /CdS	Colloidal synthesis of TíO ₂ /CdS	Gr13.VO and Carls.VC and Carls.VC C4H10C4O6 and C2H5NS C4H10C4O6 and C2H5NS	CdS nanoparticles (NPS) aggregate to form hollow nano spheres with uniform dia- meter of 150 nm on nanorods		420 nm)	filter	
Ni-TiO ₂ /CdS	Ni-TiO ₂ /CdS synthesis by photo-deposition	Ni(NO ₃) ₂					
TiO ₂ /ZnO	Sol-gel method	$C_{16}H_{40}O_4Ti$ and $Zn(NO_3)_2\cdot 6H_2O$	6-14 nm	H ₂ evolution from an aqueous methanol solution	300 W xenon lamp with UV filter, (240 nm $< \lambda < 400$ nm)	300 W xenon lamp with 5 mL semi batch reactor 211 UV filter, (240 nm < Å < 400 nm)	211
TiO ₂ /ZnS	Homogeneous hydrolysis	TiOSO4, ZnSO4 and C2,H3NS	Spherical agglomerates consisting of alternate anatase (~ 6 nm) and sphalerite crystalline islands (~ 15 nm)	tion of aqueous of Orange II	Florescent lamp (254 nm, 365 nm and 400 nm) with 8 W bower	Quartz tube with stain- less steel over	213
ZnO nanorods ZnO/in ₂ S ₃ core/shell nanorods arrays	Solvothermal method Surface functionalization route	C ₄ H ₆ O ₄ Zn-2H ₅ O, C ₆ H ₈ O, In(NO ₃) ₃ ·5H ₂ O and Na ₂ S· 9H ₃ O	Average diameter of ZnO nanorods: 100 \pm 6 nm and average length: 500-600 nm. Thickness of $\ln_2 S_3$ shell is 15 \pm 2 nm	Degradation of RhB	500 W xenon lamp with cut off filter ($\lambda \ge$ 420 nm)	500 W xenon lamp with Beaker with top covered 225 cut off filter ($\lambda \ge$ with glass plate 420 nm)	225
ZnO nanorods ZnO/CdS nanorods core/shell arrays	Solvothermal method Surface functionalization route	C ₄ H ₆ O ₄ Zn ⁻ 2H ₂ O, CdCl ₂ , CH ₄ N ₂ S, C ₆ H ₈ O ₇ , and C ₂ H ₆ O ₂	ZnO nanorods with average diameter 100 nm and CdS shell thickness 10–30 nm	Degradation of RhB, under simulated solar irradiation	No information in the article	Glass beaker immersed in ice-cold water bath	226
SnO ₂ /Fe ₂ O ₃	Co-precipitation	SnCl ₄ ·5H ₂ O, FeCl ₃ ·6H ₂ O, deionized H ₂ O and NH ₃	6-24 nm	Degradation of acid blue 62 dye	1000 W xenon lamp with cut off filter $(\lambda > 400 \text{ nm})$	Quartz jacketed reactor	234
SnS ₂ /SnO ₂	Hydrothermal synthesis <i>In situ</i> oxidation of SnS ₂ NPs under hydrothermal conditions	SnCl ₄ :5H ₂ O, C ₂ H ₅ NS, CH ₃ COOH and H ₂ O ₂	12–26 nm	Degradation of MO	1000 W xenon lamp $(\lambda > 420 \text{ nm})$	Two layer pyrex bottles with water circulation at the middle of the bottles.	241
Fe ₃ O ₄ NPs Fe ₃ O ₄ /BiOCl NCs	Colloidal synthesis	FeCl ₃ -6H ₂ O, FeCl ₃ -7H ₂ O, NH ₄ OH, deionized H ₂ O, C ₁₈ H ₃₄ O ₂ , Bi(NO ₃) ₃ , HNO ₃ , CHCl ₃ and NaC ₁₂ H ₂₅ SO ₄ .	Fe_JO4 NPs of size 7–10 nm have aggregated on large BiOGI flakes of size 200–500 nm	Photocatalytic degrada- tion of RhB dye in water under visible light.	500 W xenon lamp with cut off filter $(\lambda \ge 420 \text{ nm})$	No info mentioned in the article	252
CdS nanowires 1D CdS/ZnS core/ shell NCs	Solvothermal method	Cd(S ₂ CNEt ₃) ₂ , C ₂ H ₅ OH, Zn(CH ₃ COO) ₂ , 2H ₂ O and (NH ₂) ₂ CS	CdS nanowires diameter of 45 nm and length of several µm. Shell was composed of ZnS nanoparticles with dia- meter of about 4 nm	Degradation of MB and 4-chlorophenol (4CP)	300 W xenon lamp with cut off filter $(\lambda > 420 \text{ nm})$	Jacketed quartz reactor	257
In ₂ O ₃ /Ta ₂ O ₅	Sol-gel process	Ta(OC,Hs,Js, HCL, In(NO ₃), <i>n</i> H ₂ O and tri- block copolymer surfactant (pluronic P123)	Average pore diameter of the NCs is 5.9 nm	H ₂ production from aqueous methanol solution	300 W xenon lamp was used a simulated sunlight	External irradiated pyrex cell with closed gas circulation system	261