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Review on nanoscale Bi-based photocatalysts

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Nanoscale Bi-based photocatalysts are promising candidates for visible-light-driven photocatalytic environmental remediation and energy conversion. However, the performance of bulk bismuthal semiconductors is unsatisfactory. Increasing efforts have been focused on enhancing the performance of this photocatalyst family. Many studies have reported on component adjustment, morphology control, heterojunction construction, and surface modification. Herein, recent topics in these fields, including doping, changing stoichiometry, solid solutions, ultrathin nanosheets, hierarchical and hollow architectures, conventional heterojunctions, direct Z-scheme junctions, and surface modification of conductive materials and semiconductors, are reviewed. The progress in the enhancement mechanism involving light absorption, band structure tailoring, and separation and utilization of excited carriers, is also introduced. The challenges and tendencies in the studies of nanoscale Bi-based photocatalysts are discussed and summarized.

Conceptual insights

Photocatalytic technology is effective for environmental remediation and energy conversion. Nano Bi-based photocatalysts are an important family of visiblelight-excitable photocatalysts. Therefore, enhancing the activity of these photocatalysts by adjusting their nanostructure is one of the hot topics in photocatalysis. In this paper, we have reviewed the recent development on the improvement of the performance of nanoscale Bi-based photocatalysts to provide general and new insights into this family.

Introduction

Escalating serious environment and energy crises have resulted in a growing demand for effective environmental remediation and energy conversion techniques. These needs can be addressed using photocatalysis, which is an increasingly developed photochemical strategy. In a photocatalytic procedure, electrons and holes are generated from semiconductors under light irradiation and participate in reduction and oxidization reactions, respectively. Thus, photocatalysis promotes redox reactions, such as pollutant degradation, water oxidization, H2 evolution, CO2 reduction and N2 fixation.¹⁻¹⁹ Consequently, photocatalysis is regarded as a green technique for removing organic pollutants and producing fuel or electricity. Given that visible-light energy constitutes about 43% of solar energy, visible-light-responsive photocatalysts are preferred in photocatalysis and photoelectrocatalysis.

Up to now, many types of semiconductors, including metal oxides (TiO₂, ZnO, Ag₂O, Fe₂O₃, Cu₂O, Ta₂O₅), metal sulfides (ZnS, CdS, MoS₂, Bi₂S₃), multi-component oxides (Bi₂WO₆, InTaO₄, BiVO₄, Ag₃VO₄, SrTiO₃), metal selenides (MoSe₂, CdSe), metal phosphides (Ni₂P), metal phosphates (Ag₃PO₄), metal halides and oxyhalides (AgBr, BiOBr) and metal-free materials (SiC, g-C₃N₄, and Si), have been employed as photocatalysts.3,4,20-24 Among them, those that possess a band gap of $E_{\rm g} > 3$ eV are called wide-band-gap photocatalysts, such as TiO2, ZnO, ZnS, SrTiO3, and KTaO₃. In contrast, those with a band gap of $E_g \leq 3$ eV are called visible-light-responsive photocatalysts, such as Ag₂O, Bi₂WO₆, InTaO₄, CoO, BiVO₄, Fe₂O₃, Cu₂O, Ag₃VO₄, TaON, CdS, Ta₃N₅, CdSe, Bi₂S₃, SiC, g-C₃N₄, and Si. Bi-Based photocatalysts are important visible-light-responsive photocatalysts and have recently attracted rapidly increasing attention, shown not only from the number of publications, but also from the ratio of publications in the field of photocatalysis (see Fig. 1a and b). Considering the stability of Bi³⁺, most studies have focused on Bi³⁺-containing compounds, such as Bi_2O_3 , 25 BiVO_4 , 26 $\text{Bi}_4\text{Ti}_3\text{O}_{12}$, 27 $\text{Bi}_{12}\text{TiO}_{20}$, 28 $\text{Bi}_2\text{O}_2\text{CO}_3$, 29 Bi_2WO_6 , 30 BiPO_4 , 31 BiFeO_3 , 32 BiOX (X = Cl, Br, I), 33 Bi₃TiNbO₉,³⁴ and Bi_{0.5}K_{0.5}TiO₃.³⁵ The majority of these compounds possess a layered structure and plate-like appearance. Bi⁵⁺-Containing compounds, such as LiBiO₃, NaBiO₃, and KBiO₃,

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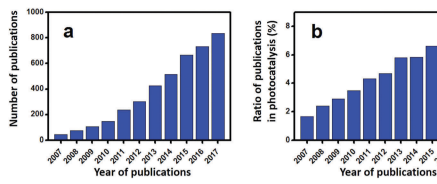


Fig. 1 (a) Number of publications of Bi-based photocatalysts during the last decade (source: web of science; date: 8th April 2018; key word: bismuth and photocatalytic), (b) the ratio of publications of Bi-based photocatalysts in the field of photocatalysis (source: Web of science; date: 8th April 2018; key word: photocatalytic)

can also be excited by visible light. However, such types are less reported than the others because of the instability of Bi⁵⁺. In Bi(III) compounds, the hybridized O 2p and Bi 6s2 orbitals may cause an upshift of the valence bands (VBs).33 Therefore, the band gaps of Bi compounds are usually smaller than 3.0 eV and can be excited by visible light. Bi-based photocatalysts are regarded as promising candidates for removing toxins from water and air^{3,36-38} as well as for the production of fuel. 2,39-41 However, the photocatalytic performance of bulk Bi-based semiconductors is not as high as those of nanoscale Bi-based photocatalysts, because the photogenerated electrons and holes of these materials have not been easily exploited and utilized. Also, bulk photocatalysts have weaker light absorption and smaller surface areas than nanoscale photocatalysts (see Fig. 2). Numerous attempts have been made to enhance bulk Bi-based semiconductors to achieve ideal photocatalytic activity. These studies have focused on nanoscale component adjustment, morphology control, heterojunction construction, and surface modification^{2,37,38,42} (Fig. 3).

Component adjustment

The band structure parameters, such as VB, conduction band (CB), and band gap (E_{φ}) , are crucial to the photocatalyst activity.

Furthermore, component adjustments, such as doping, solidsolution preparation and stoichiometry alteration, are effective methods that can be used to tune the band structures of bismuthal semiconductors. Therefore, suitable component adjustment is favorable for bismuthal photocatalysts.

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Doping

Heteroatom doping is widely adopted to increase the visible-light absorption of photocatalysts, because this process can generate a doping level between the CB and VB (Fig. 4). Consequently, the energy required to excite electrons decreases, and the light response of semiconductors increases. 43-47 Doping can also improve the charge transmission properties of semiconductors and lead to the augmented transfer efficiency of carriers. Doping is commonly employed to enhance the activity of Bi-based semiconductors.

Many bismuthal compounds are visible-light responsive. Thus, doping has been performed more frequently to enhance the visible-light absorption of bismuthal compounds with a wider band-gap, such as in Bi₂O₂CO₃, 48 BiPO₄, 49 BiOCl, 50 and Bi₃TiNbO₉. 34 For example, the band gap of Bi₃TiNbO₉ can be reduced from 3.1 eV to 2.6 eV when it is doped with a 10% molar ratio of Ni²⁺. ³⁴ Zheng et al. ⁵¹ found that the band gap of

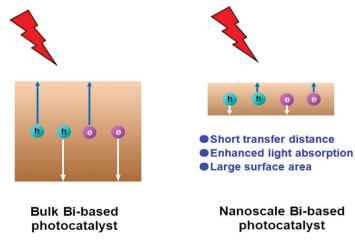


Fig. 2 Schematic illustration showing the superiority of nanoscale photocatalysts.

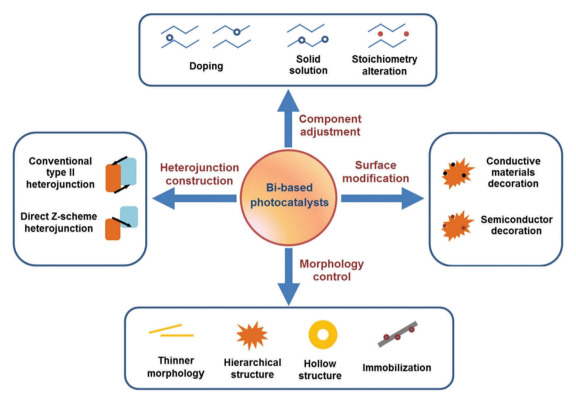


Fig. 3 Enhancement strategies for nanoscale Bi-based photocatalysts.

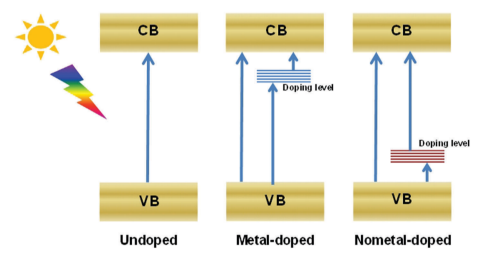


Fig. 4 The energy level mechanisms for metal-doped and nonmetal-doped photocatalysts.

Bi₂WO₆ can be narrowed by doping a small amount of Br with the surfactant cetyltrimethylammonium bromide (CTAB) during preparation. Doping by up to approximately 3% can also narrow the band gap of Bi₂MoO₆ from 2.96 eV to 2.69 eV and consequently improve the performance of Bi₂MoO₆ in the photocatalytic degradation of Rhodamine B (RhB).⁵² Zhang et al.⁵³ studied the effects of doping on the electronic structures and optical properties of BiOCl using first-principle calculations. Studies have also proposed that codoping with Sb and I can substantially narrow the band gaps and increase the light absorption of BiOCl because of the difference in the electronegativities between the Sb/I and Bi/Cl atoms.

Doping can further narrow the band gap of visible-lightresponsive bismuthal compounds.^{54,55} However, the improvement resulting from a narrowed band gap is limited, because visible light response is not the main barrier to the photocatalytic performance of bismuthal compounds. Low utilization efficiency of the generated carriers is the primary issue to be solved for these bismuthal photocatalysts. In this case, doping can also enhance the photocatalyst activity if the dopant can reduce impedance or improve charge separation. For instance, F-doped β-Bi₂O₃ has been shown to exhibit higher photocatalytic activity than that of pure β-Bi₂O₃ in the degradation of

methyl orange (MO).⁵⁶ This can be attributed to the higher separation efficiency of electron-hole pairs and lower VB position (indicating stronger oxidation capacity). B-Doping can increase the photocatalytic activity of BiOBr on the inactivity of Escherichia coli, which is attributed to the role of B as the electron acceptor for the effective separation of electrons and holes.⁵⁷ Such a role may have been caused by the empty p-orbital and unpaired electrons in B. Br doping could reduce the effective mass of electrons in Bi₂WO₆, by which the transfer and separation of the carrier can be facilitated.⁵⁸ Aurivillius-type Ce-doped SrBi₂Ta₂O₉ fabricated by Senthil and co-workers⁵⁹ exhibited a higher photocatalytic activity than SrBi₂Ta₂O₉ for H₂ generation. The improvement comes from the enhanced separation of

electrons and holes through a dipole moment derived from

the bonding of Ce⁴⁺ with the surrounding oxygen atoms. Zhang and colleagues⁶⁰ found that carbon doping increases the internal electric field by 126-fold as a result of the enlarged differences in the electrostatic potentials between the [Bi₃O₄] and [Cl] layers induced by the carbon doping (Fig. 5). Consequently, the separation of holes and electrons was improved. Er-Doped $\mathrm{Bi}_{24}\mathrm{O}_{31}\mathrm{Br}_{10}$, ⁶¹ carbon-doped $\mathrm{Bi}_{2}\mathrm{WO}_{6}$, ⁶² Mo-doped BiVO_{4} , ⁶³ and Er/Yb codoped BiVO₄⁶⁴ also showed enhanced photocatalytic activities relative to those of the original bismuthal compounds due to increased carrier transfer and separation.

Anion and mixed-valence cation dopants can also improve the photocatalytic performance of bismuthal semiconductors by facilitating the transfer and separation of carriers. For instance, doping with ${\rm IO_3}^-$ can increase the activity of BiOI in

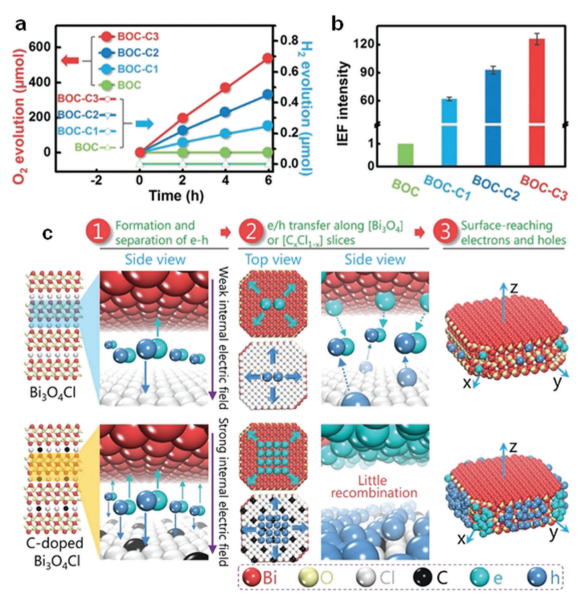


Fig. 5 (a) The performance of visible-light-driven oxygen and hydrogen evolution; (b) internal electric field intensity of C-doped Bi₃O₄Cl (assuming the intensity of BOC to be "1"); and (c) schematic diagram of the carrier migration of pure and C-doped Bi₃O₄Cl; BOC-C1, BOC-C2, and BOC-C3 present C-doped Bi₃O₄Cl with carbon concentrations of 0.92%, 1.86%, and 3.16%, respectively. Reproduced with permission from ref. 60. Copyright 2016 John Wiley and Sons.

the photocatalytic degradation of MO because of the key role played by ${\rm IO_3}^-$ in improving the transfer efficiency of the carriers.65 In-Doped BiOI exhibited augmented activity in the photocatalytic degradation of p-chloroaniline, because the introduced In resulted in the formation of a doping energy level that acted as a scavenger of photo-induced electrons and consequently prevented the recombination of charges.⁶⁶ Jiang et al.⁶⁷ fabricated Sn⁴⁺-doped Bi₂S₃ through a solvothermal method, in which Sn4+ acted as an electron scavenger by interconversion between Sn^{4+} and Sn^{2+} and facilitated O_2 reduction. Consequently, the photocatalytic performance of Sn⁴⁺-doped Bi₂S₃ was enhanced. Similarly, the dopant Tb acted as an electron-trapping agent through interconversion between Tb⁴⁺ and Tb3+ and improved the photocatalytic activity of Bi2MoO6 in RhB degradation.68

Although the above conventional doping of heteroatoms can enhance the activities of bismuthal compounds, the introduced hetero element may alter the space lattice and cause additional stress. This process suppresses the introduction of the dopant and limits the improvements of visible-light absorption and carrier transfer derived from doping. Hence, doping by substituting the original element, rather than interstitial doping, is desired.

For example, when Bi and Ti in Bi₄Ti₃O₁₂ are partially replaced by Cr, the doped Bi₄Ti₃O₁₂ exhibits a higher activity in H₂ generation because of the considerably narrowed band gap and enlarged light absorption of Bi₄Ti₃O₁₂.⁶⁹ The cosubstitution of Gd and Mn in BiFeO₃ nanoparticles (NPs) reduces the band gap and electrical resistivity of BiFeO₃.⁷⁰ The band gap of BiOCl was narrowed when Bi3+ was partially substituted by Sn²⁺, and this process consequently endowed BiOCl with increased photocatalytic activity for degrading RhB.⁷¹ When O in BiVO₄ was partially substituted by F through a solid-vapor reaction, the photoelectrochemical water oxidation on BiVO₄ was strongly promoted.⁷² This effect may be attributed to the more negative flat band potential (an upshift of approximately 0.1 V) and increased carrier density resulting from the doping of F. The moderate upshift of the flat band potential makes the consumption of electrons easier, which is favorable to the holes in the VB for oxidation reactions. Bi₃FeMo₂O₁₂ was found to be superior to Bi₃GaMo₂O₁₂ in the photocatalytic degradation of RhB under visible-light irradiation. This result was due to the narrower band gap of Bi₃FeMo₂O₁₂ (2.3 eV) than that of Bi₃GaMo₂O₁₂ (2.95 eV) as a result of the contribution of Fe 3d orbitals to the CB.73 Nevertheless, not all elemental substitutions lead to a reduced band gap in bismuthal compounds. If Bi and Fe in BiFeO₃ NPs are cosubstituted with Ca and Ti using a sol-gel method, the band gap will increase upon an increase in the amount of Ca and Ti.74

Elemental substitution can also improve the separation of carriers and thus boost the performance of photocatalysts. Gu et al. 75 obtained Ce-doped BiVO₄ by substituting Bi³⁺ with Ce ions in a homogeneous precipitation synthesis. The introduction of Ce enhanced the photocatalytic performance of MB and phenol removal by effectively hindering the recombination of photo-generated carriers. BiV_{0.8}Mo_{0.2}O₄ exhibited an improved

photoelectrochemical activity in water oxidation because of the more efficient charge transport and electron-hole separation derived from the formation of cation vacancies (Bi5+)/oxygen interstitials as a result of high Mo doping.⁷⁶

Solid solution

Fabricating solid solutions is an effective technique to adjust the band structure of photocatalysts by hybridizing two or more crystalline phases at the atomic scale. A solid solution can be regarded as a special doped semiconductor, in which the anions or cations of the host semiconductor are selectively replaced by introduced ones over the whole range of the composition.⁷⁷ By adjusting the concentration of cation and/or anion substitution (by controlling the components and composition), the band structure of a solid solution can be tuned. 78,79 Although the aforementioned doping methods can enhance the performance of bismuthal photocatalysts to some extent, the shortcomings of doping are obvious. The formed doping levels are generally discrete (Fig. 3), and the dopant may also act as the recombination center of carriers and increase the number of recombination sites for electrons and holes. 78,79 Contrary to doping, solid solutions contain no discrete energy levels. The band structures of solid solutions are uniform, and their band gaps usually fall in the region between those of original semiconductors (Fig. 6). Consequently, the electron excitation can be improved with a lower increase in carrier recombination.

Bismuth oxyhalide solid solutions are the most frequently studied among Bi-based solid solutions because of the similar structures of BiOX (X = Cl, Br, I) and easy fabrication of solid solutions comprising these compounds. Hence, several studies have also considered BiOX as a kind of heterooxyhalidesubstituted bismuth oxyhalide.80 Furthermore, along with the ratio of oxyhalides, the absorption edges of bismuth oxyhalide solid solutions usually vary gradually between the absorption edges of the pristine compounds (Fig. 7).81 This phenomenon renders the tailoring of the band structure as highly convenient. For example, hierarchical Bi-rich Bi₄O₅Br_xI_{2-x} solid-solution nanosheets can be prepared via a precursor method and show improved photocatalytic activity for CO₂ conversion and Cr(v₁) removal.⁸² Flower-like BiOCl_{0.5}Br_{0.5} hierarchical solid solutions can be fabricated through reacting Bi₂O₃ with a mixed solution of KCl and KBr at room temperature in the presence of glacial acetic acid and exhibit enhanced photocatalytic activity for RhB degradation under visible-light irradiation, compared to pure BiOCl and BiOBr. 83 Similarly, 3D flower-like BiOCl_xBr_{1-x}, 84 $BiOCl_xI_y$, 85 and $BiOCl_{1-x}Br_x$ hierarchical microspheres 86 have also been prepared by simple one-pot methods and have exhibited better activities than pure BiOCl, BiOBr, and BiOI. When the three latter halogens were introduced during fabrication, ternary solid solutions BiOCl_xBr_yI_z (x + y + z = 1) could be obtained, and the as-prepared solid solutions generally possessed a controllable band gap and highly enhanced visible-light photocatalytic activity. 87 F-Containing bismuth oxyhalide solid solutions are less reported in solid-solution preparations because of the partial miscibility of $BiOF_{1-x}Y_x$, as proposed by Zhao et al.⁸⁸ They examined the properties of $BiOX_{1-x}Y_x$ (X, Y = F, Cl, Br, I)

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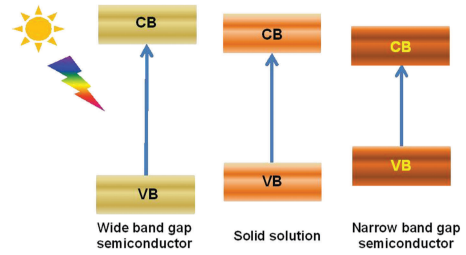


Fig. 6 Schematic diagram of the band structure tuning through solid-solution fabrication.

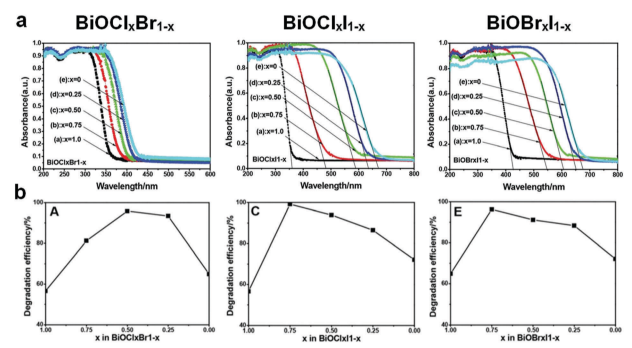


Fig. 7 (a) UV-visible light diffuse reflectance spectra and (b) RhB degradation efficiency of $BiOCl_x Br_{1-x}$. $BiOCl_x Br_{1-x}$ and $BiOBr_x l_{1-x}$ solid solutions. Reproduced with permission from ref. 81. Copyright 2013 Royal Society of Chemistry.

solid solutions through density functional theory calculations. The results showed that several properties of $BiOCl_{1-x}Br_x$, $BiOBr_{1-x}I_x$, and $BiOCl_{1-x}I_x$ solid solutions changed almost linearly with x, while the properties of $BiOF_{1-x}Y_x$ (Y = Cl, Br, I) solid solutions are partially miscible. In addition, the related calculated data were difficult to fit. Therefore, this behavior has been attributed to the partial miscibility of $BiOF_{1-x}Y_x$.

In addition to bismuth oxyhalides, other Bi-containing solid solutions were found to be more efficient than pure bismuthal compounds for visible-light-driven photocatalysis. For instance, Terebilenko et al. 89 fabricated Bi_{1-x/3}V_{1-x}Mo_xO₄ solid solutions for photocatalytic water oxidation. The resulting solid solutions possessed the same band gap of 2.25 eV (narrower than that of BiVO₄), which endowed such solid solutions with high visiblelight absorption. A Sr_{0.9}Bi_{0.1}Ti_{0.9}Fe_{0.1}O₃ solid solution synthesized by Lu et al. 90 performed more effectively than SrTiO3 in H2 production under full-range irradiation ($\lambda \geq 250$ nm). This finding was attributed to the suitable band gap of the former compound. The resulting band gap can be tuned by adjusting the ratio of Bi to Sr (see Fig. 8).

Stoichiometry alteration

The layered structures of bismuthal compounds render the component adjustment convenient. Even without introducing a foreign element, the band structures can also be tuned by changing the stoichiometry of the bismuthal semiconductor.

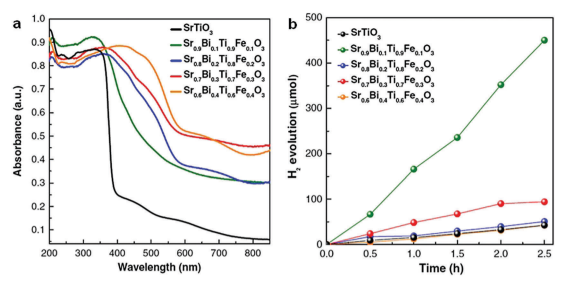


Fig. 8 (a) UV-visible diffuse reflectance spectra and (b) photocatalytic H_2 production of $Sr_{0.9}Bi_{0.1}Ti_{0.9}Fe_{0.1}O_3$ solid solutions under full-range irradiation ($\lambda \geq 250$ nm). Reproduced with permission from ref. 90. Copyright 2017 Elsevier.

In this field, similar to solid solutions, bismuth oxyhalides have been extensively studied because the band gap, VB, and CB of oxygen-rich (bismuth-rich) or oxygen-deficient compounds usually changes gradually along with the atomic ratio. Consequently, the compounds can be enhanced by stoichiometric variation. In a bismuth oxyhalide, both the O 2p and X np (n = 3, 4, and 5 for X = Cl, Br, and I, respectively) states have a major contribution to the VB, while the Bi 6p states dominate the CB. When the O content increases, the O 2p states of bismuth oxylodide become more significant than the X np state in the VB, therefore the band gap energies of O-rich bismuth oxylodides are closer to those of Bi₂O₃. 91 That is, the VB can be tuned by changing the O/X ratio.

The band gap of BiOI is very narrow, but its CB is less negative, and its VB is less positive. These characteristics result in the low reduction and oxidation capacities of BiOI. Consequently, the CB upshift and VB downshift become more favorable, but excessive shift in the CB and VB may result in a wide band gap, which is unfavorable for photocatalysis. That is, a moderate upshift of CB and downshift of VB are required. Generally, the VB of bismuth oxylodides downshifts to a more positive level because of the increased contribution of the O 2p states, while the CB upshifts to a more negative position owing to the additional contribution of I 5p, thus the band gap of the bismuth oxylodides increases with an increase in the bismuth and oxygen content. Hence, bismuth oxylodides with moderate overdoses of bismuth and oxygen, such as Bi₄O₅I₂ or Bi₆O₉I₃, are more advisable. For instance, Bi₄O₅I₂ nanosheets (fabricated using a solvothermal method) exhibited a higher activity than BiOI in the photocatalytic degradation of RhB because of the more positive VB of the former. 92 For BiOCl, narrowing the wide band gap is the primary aim of the atomic ratio adjustment. Oxygen-rich bismuth oxychlorides, such as $Bi_{24}O_{31}Cl_{10}$ ($E_g = 2.71$ eV, ref. 93) and $Bi_{12}O_{17}Cl_2$ ($E_g = 2.57$ eV, ref. 94), possess narrower band gaps and can be excited by visible light. Therefore, the visible-light

photocatalytic performances of these bismuth oxychlorides are better than that of BiOCl. Moreover, Cl-rich bismuth oxychloride was also found to possess a narrower band gap. For instance, the band gap of Bi₁₂O₁₅Cl_{l6} nanosheets fabricated by Wang et al.95 was estimated to be 2.36 eV, and the photocatalytic performance of Bi₁₂O₁₅Cl_{l6} nanosheets for bisphenol A (BPA) removal was superior to that of BiOCl nanosheets. Studies on oxygen-rich bismuth oxybromides have mainly focused on Bi₄O₅Br₂ because of its narrow band gap. In reports by Xia and co-workers, 96 Bi₄O₅Br₂ nanosheets fabricated using a solvothermal method were shown to possess a narrower band gap (2.37 eV) and lower carrier transfer resistance than those of BiOBr (2.82 eV) and were more effective than BiOBr in the photocatalytic degradation of BPA. For similar reasons, Bi₄O₅Br₂ showed much higher activity than BiOBr in the photocatalytic degradation of resorcinol97 and ciprofloxacin (CIP)98 under visible light. Furthermore, Bi₄O₅Br₂ nanosheets were also discovered to be more suitable than BiOBr nanosheets in the photoreduction of CO₂. 99 These results are due to the CB position of Bi₄O₅Br₂ being higher than that of BiOBr and thus more favorable for the reduction reaction of CO₂.

The band structures of other Bi compounds, such as BiVO₄, Bi₂SiO₅, and Bi₂O₃, can also be altered by varying the stoichiometry for activity enhancement. Batool et al. 100 found that Bi₄(SiO₄)₃ nanofibers possess lower impedance and exhibit higher photocatalytic performance than Bi₂SiO₅ nanofibers in the degradation of MO. Huang and co-workers¹⁰¹ prepared Bi₄V₂O₁₁ through a solvothermal method. The synthesized Bi₄V₂O₁₁ had a narrower band gap (2.15 eV) than that of BiVO₄ (2.4 eV) and showed high water oxidation activity under 300 W Xe lamp irradiation. In Kalyan's research, a BiO_{2-x} phase with a narrow band gap (1.84 eV) formed on the surface of a Bi₂O₃ photoanode during photoelectrochemical testing in the presence of KOH and H₂O₂. The formed BiO_{2-x} phase improved the photocurrent because of its narrow band gap.

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Morphology control

The photocatalytic properties of semiconductors are highly dependent on their morphologies and sizes. 103 High specific surface area, low thickness, and hierarchical and hollow structures can increase light absorption and accessibility of photocatalysts. Such properties enable the accelerated migration of carriers to the surface and reduce the dosage of photocatalysts. Accordingly, the performance of Bi-based photocatalysts can be enhanced by building ultrathin and hierarchical and hollow structures or by immobilization on substrates with specific morphologies.

Thin morphologies

Thickness greatly affects the activities of bismuthal compounds because of the influence of such a variable on the efficiency of light absorption and the distance of the photo-generated carriers from the surfaces. A smaller sized bismuthal compound usually leads to a higher specific surface area and better photocatalytic and photoelectrochemical performance. 104,105 Considering unique layered plate-like structures, the exposed-facet ratio of bismuthal compounds usually changes along with the thickness of the compounds. Therefore, the thickness effect has often been discussed in related reports. Thinner structures can endow bismuthal compounds with a higher specific surface area, increased light absorption efficiency, and enabled utilization of photogenerated electrons and holes. Ultimately, such properties may enhance photocatalytic activity. With a decrease in the thickness, the surface energy of the nanoplates would also largely increase, single plates become thermodynamically unstable and these nanoplates prefer to stack together into one larger particle to reduce surface energy during growth. As a result, many Bi-based photocatalysts usually possess a granular appearance comprising nanoplates. The thin morphology discussed in this section refers to these nanoplates.

Chen and colleagues 106 found that reducing the thickness of BiOBr nanosheets can significantly increase the exposed (001) facets and the photocatalytic activity of the BiOBr nanosheets. The enhanced photocatalytic activity of the BiOBr nanosheets was ascribed mainly to the enhanced absorption of visible light and improved separation efficiency of charge carriers due to the ultrathin structure. Li et al. 107 proposed that the photoactivity of the BiOBr nanosheets depends on nanosheet thickness. Similarly, reducing the thickness of Bi₂O₂CO₃

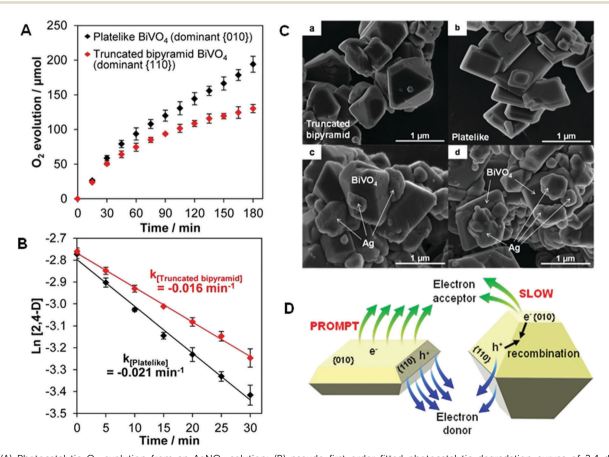


Fig. 9 (A) Photocatalytic O2 evolution from an AgNO3 solution; (B) pseudo-first-order-fitted photocatalytic degradation curves of 2,4-dichlorophenoxyacetic acid (2,4-D) on truncated bipyramid (red) and plate-like (black) BiVO₄ under visible light ($\lambda > 420$ nm); SEM images of bare (C-a) truncated bipyramid and (C-b) plate-like BiVO₄ and the corresponding Ag-deposited (C-c) truncated bipyramid and (C-d) plate-like BiVO₄. (D) Schematic diagram of the promotion induced by increasing the size of the (010) facets in BiVO₄. Reproduced with permission from ref. 111. Copyright 2016 American Chemical Society

[accompanied by an increase in the (001) facets] improved its photocatalytic activity in the degradation of RhB, 108 and the activities of Bi₂WO₆ were augmented along with a reduction in thickness. 109

Thinner structures sometimes result in additional facet and structure differences, which affects the performance of bismuthal photocatalysts. For example, increasing the exposure of (001) facets (accompanied by a reduction in the thickness) can improve the efficiency of RhB degradation on Bi₂Fe₄O₉ nanoplates, because a larger exposed (001) surface can provide more Fe³⁺ cations to generate more hydroxyl radicals.¹¹⁰ A reduction in the thickness can also improve the photocatalytic performance of BiVO₄ in O₂ evolution, owing to the increased ratio of reduction functional (010) facets versus dominating oxidation functional (110) facets (Fig. 9). 111 Hu et al. 112 noted that reducing the thickness of BiTaO4 not only increased the specific surface area but also caused upshifts in the VB and CB. Density of states (DOS) calculations showed that the energy levels of the VB and CB of the (010) facets of BiTaO4 are more negative than those of bulk BiTaO4. The (010) facets in BiTaO4 increase along with a reduction in the thickness, therefore, thinner BiTaO₄ single-crystal nanoplates possess more {010} facets and higher VB and CB positions.

When the thickness of a bismuthal semiconductor is reduced to below 10 nm, the specific surface area, light absorption,

and carrier migration will be further enhanced. Sometimes, new characteristics might also arise. Ultrathin square-like BiOCl nanosheets (3-7 nm in thickness) have been shown to exhibit a higher activity than thicker nanosheets in the photocatalytic degradation of RhB due to a larger specific surface area. 113 Di et al. 114 reported the preparation of ultrathin BiOI nanosheets and their composites, which displayed a uniform thickness of approximately 0.9 nm (Fig. 10), approaching the thickness of one [I-Bi-O-Bi-I] unit (0.9149 nm). The CB position of the BiOI nanosheets was determined to be -0.95 eV (NHE), negative to reduce O₂ to O₂. This characteristic is beneficial for H2 generation or CO2 reduction. However, the researchers did not provide results on the reduction application, although their reports on the surface modification of BiOI nanosheets still exhibited improved activity for the photocatalytic degradation of RhB. The ultrathin H_{1.78}Sr_{0.78}Bi_{0.22}Nb₂O₇ nanosheets (Fig. 11) prepared by calcination exhibited 5.5 and 26.2 times higher activity than H_{1.78}Sr_{0.78}Bi_{0.22}Nb₂O₇ plates and SrBi₂Nb₂O₉ platelets for photocatalytic H₂ generation, respectively. This result can be ascribed to the higher separation efficiency of the photo-generated carriers and specific surface area. 115

Preparation of fabric or quantum sized materials is another method for obtaining thin morphologies and high specific surface areas. Bharathkumar et al. 116 fabricated BiFeO3 fibers

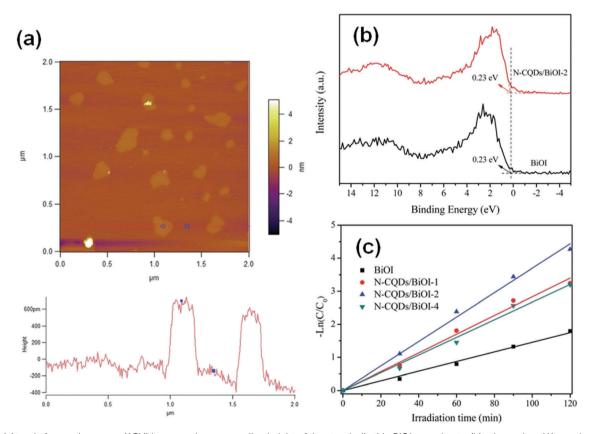


Fig. 10 (a) Atomic force microscopy (AFM) image and corresponding height of the atomically thin BiOI nanosheets; (b) valence-band X-ray photoelectron spectra and (c) pseudo-first-order-fitted kinetic curves (for the RhB degradation) of pure BiOI and N-carbon quantum dot (CQD) decorated BiOI. Reproduced with permission from ref. 114. Copyright 2016 Royal Society of Chemistry

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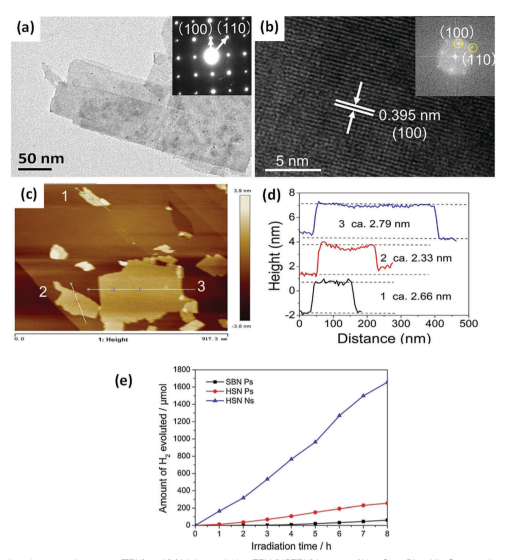


Fig. 11 (a) Transmission electron microscopy (TEM) and (b) high-resolution TEM (HRTEM) images of H_{1.78}Sr_{0.78}Bi_{0.22}Nb₂O₇ nanosheets (HSN Ns); (c) AFM image; (d) corresponding height of the HSN Ns; and (e) the H2 evolution curves of the SrBi₂Nb₂O₉ platelets (SBN Ps), H_{1.78}Sr_{0.78}Bi_{0.22}Nb₂O₇ platelets (HSN Ps), and HSN Ns. Reproduced with permission from ref. 115. Copyright 2017 Elsevier

using an electrospinning method. The obtained BiFeO₃ fibers then exhibited improved relative photocatalytic activity compared with that of BiFeO₃ particles for degrading RhB, due to their high specific surface area. Zhao and co-workers¹¹⁷ prepared metal Bi nanofibers using an aqueous reduction procedure and found that Bi nanofibers exhibited a higher photocatalytic activity than Bi nanoplates and NPs in degrading RhB. Yang et al. 118 fabricated bismuth oxide formate (HCOOBiO) nanowires (Fig. 12) with a diameter of 20 nm and a length of up to several hundreds of micrometers in N,N-dimethylformamide (DMF) solution via a solvothermal route. These HCOOBiO nanowires showed a higher activity than HCOOBiO microspheres in the photocatalytic degradation of MO. In the studies of Lu and colleagues, 119 BiVO4 nanofibers were prepared using an electrospinning and calcination method and showed a higher efficiency than BiVO4 rods in the photocatalytic removal of MB under visible-light irradiation. Additionally, Bi₂S₃ with a high length-to-width ratio was found to

be highly effective in the photocatalytic reduction of CO2 to CH₃OH. 120 The high length-to-width ratio not only increases the specific surface area but also enlarges the band gap (by a downshift of the VB and an upshift of the CB) owing to the increased influence of the surface state resulting from the smaller size. And as it happens, the CB is upshifted to a level sufficient enough for CO₂ reduction. Bismuth monoxide quantum dots (QDs) obtained using a hydrothermal method exhibited high efficiency in the photocatalytic reduction of N_2 . The ammonia production rate reached 1226 μmol g⁻¹ h⁻¹ in the absence of a sacrificial agent and a cocatalyst. As Bi₅O₇Br was tailored into nanotubes with a diameter of 5 nm, its CB (-1.14 eV) was more negative than that of BiOBr. Though such a CB is still not thermodynamically accessible for direct free N2 fixation, N2 fixation was achieved because the formed oxygen vacancy in samples injects trapped photogenerated electrons directly into the chemically adsorbed N2 and weakens the N≡N triple bond. 122

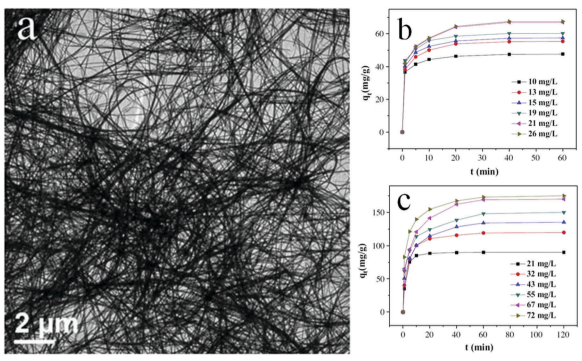


Fig. 12 Scanning electron microscopy (SEM) images (a) of ultra-long HCOOBiO nanowires. Removal of (b) MO and (c) Cr(VI) at various initial concentrations from ultra-long HCOOBiO nanowires. Reproduced with permission from ref. 118. Copyright 2015 Elsevier.

Hierarchical structures

Photocatalysts with hierarchical structures usually exhibit better performances than bulk photocatalysts. Such superiority is due to the high specific surface area, efficient light harvesting, accessibility, and easy transport of the reactants of the former catalysts. Yang et al. 123 accomplished the early work on hierarchical ordered oxides. Since then, the design and synthesis of hierarchical-structured nanomaterials have attracted significant attention. 124-126

Specific surface area is one of the most important factors that results in the higher activity of bismuthal compounds. Generally, photocatalysts with a hierarchical structure possess relative higher specific surface areas than bulk photocatalysts or individual rods and plates, and work better in photocatalytic reactions. For example, grain-like Bi₂₄O₃₁Br₁₀ hierarchical architectures (Fig. 13) fabricated using a solvothermal method in the presence of starch also showed improved photocatalytic activity in the degradation of RhB.¹²⁷ Qin and co-workers¹²⁸ prepared BiYO₃ with a high specific surface area using a hydrothermal method with soft templates. The BiYO₃ prepared using disodium ethylenediaminetetraacetate (EDTA) as a template possessed the highest specific surface area and photocatalytic activity for reducing CO₂. Jia et al. 129 found that hierarchical Bi₂S₃ produced using a hydrothermal method [with cetyltrimethylammonium bromide (CTAB) as a surfactant] exhibited higher photocatalytic activity than Bi₂S₃ nanorods in the degradation of RhB. This performance was ascribed to the synergetic effect of the shape, surface area, band gap, crystallinity, and size resulting from the 3D hierarchical structure. Han's group 130 fabricated flower-like

δ-Bi₂O₃ by thermally treating bismuthyl acetate CH₃COO(BiO), which was prepared from Bi₂O₃ and glacial acetic acid (HAc) or Bi(NO₃)₃·5H₂O and HAc. The obtained δ-Bi₂O₃ showed superior activity over that of δ -Bi₂O₃ sheets in the photocatalytic degradation of RhB. Zhou et al. 131 prepared uniform cylindrical BiFeO₃ samples by adjusting the ratio of HNO3: NaOH to 8:12 in the hydrothermal process. The uniform cylindrical BiFeO₃ exhibited enhanced photocatalytic activities for removing RhB under visible-light irradiation, because the uniform structures favored charge transport and increased the accessibility to organic matter. Li et al. 132 prepared BiOBr strips accumulated from tiny pieces using a solvothermal method with bismuth subsalicylate as a template. They found that the BiOBr strips showed an enhanced performance for the photocatalytic degradation of RhB. Furthermore, the hierarchical structure of bismuthal oxyiodide obviously promoted the performance of the compound in phenol removal.133

Even in the construction of a hierarchical structure, the morphologies of bismuthal photocatalysts can be further tuned to different forms by adjusting the solvent, pH, and surfactants, as well as by adding capping agents. For instance, Wang et al. 134 synthesized 3D hierarchical flower-like α-Bi₂O₃ microspheres with various morphologies (Fig. 14) in an ethanol-water mixture. They heated the mixture in an oil bath at 95 °C, and the morphology was controlled by adjusting the amount of added glycerol and oleic acid (capping agents). Sarka et al. 135 fabricated Bi₂S₃ in different shapes (Fig. 15) using a solvothermal method with various solvents. The performance of Bi2S3 nanocrystals in removing MB was better than those of the Bi2S3 nanorods and nanoplates due to the higher specific surface

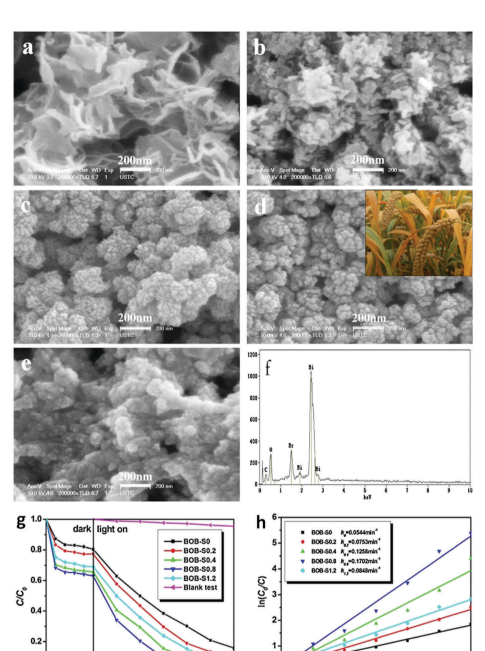


Fig. 13 SEM images of Bi₂₄O₃₁Br₁₀ hierarchical architectures (BOB) (a) BOB-S0, (b) BOB-S0.2, (c) BOB-S0.4, (d) BOB-S0.8 (grain-like), and (e) BOB-S1.2 (e.g., -\$1.2 indicates that the mass of starch used was 1.2 g); (f) EDS pattern of BOB-\$0.8; (g) photocatalytic degradation curves of RhB; and (h) pseudofirst-order-fitted kinetic curves over Bi₂₄O₃₁Br₁₀ samples under visible-light irradiation. Reproduced with permission from ref. 127. Copyright 2016 Elsevier.

area and suitable band structure of the Bi₂S₃ nanocrystals. Zhao et al. 136 altered the morphologies of BiVO₄ from olive-shaped to primrose-like, leaf-shaped, or strip-like morphologies (Fig. 16), by adjusting the concentration of glycerol in the aqueous solution during template-free solvothermal synthesis. In addition to solvent control, the morphologies of BiVO4 can also be modified by pH adjustment. Besides this, the hierarchical morphology alteration of BiVO₄ can also be achieved through

Time(min)

0.0-

adjusting the pH.137 The relationship between pH and the morphologies of BiVO4 is illustrated in Fig. 17. If NaHCO3 is added in the solid-state synthesis of BiVO₄, the morphology can be turned from polyhedral to a rice-like shape, by increasing the nucleation rate and hindering the continual growth of the original product particles. 138 In the preparation of Bi₂WO₆, its morphology can be tuned from accumulated sheets to a complex morphology by adjusting the molecular weight of the

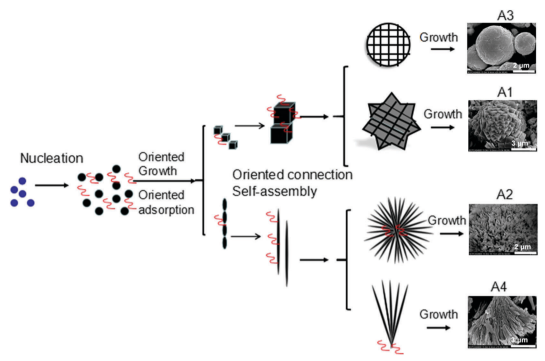


Fig. 14 3D hierarchical flower-like α-Bi₂O₃ microspheres fabricated using different solvents: A1 (0.5 mL of glycerol), A2 (1 mL of glycerol), A3 (1 mL of glycerol and 1 mL of oleic acid), and A4 (1 mL of glycerol and 2 mL of oleic acid). Reproduced with permission from ref. 134. Open Access 2016 World Scientific.

added polyvinylpyrrolidone (PVP). 139 Examples of such complex morphology include flower-like, red blood cell-like, and square pillar-like morphologies. Bi₂WO₆ with a thin sheet-like appearance showed the best photocatalytic performance in the decomposition of RhB under visible-light irradiation.

Hollow and porous structures

The hierarchical structure of Bi compounds can be transformed to a hollow structure, which is more preferable for improving the photocatalytic performance, using a suitable method. For instance, Qian et al. 140 fabricated mesoporous Bi₄Ti₃O₁₂ by calcinating a mixture of Bi(NO₃)₃ and tetrabutyl titanate. The product had a markedly higher specific surface area (109.4 m² g⁻¹) and exhibited a better performance than that of the bulk compound. Gong and co-workers¹⁴¹ prepared porous BiVO₄ by electrospinning and calcination in the presence of PVP. The obtained porous BiVO₄ had a larger specific surface area and enhanced photocatalytic activity for the degradation of MB. Kashfi-Sadabad et al. 142 synthesized Sm-doped Bi₂MoO₆ hollow spheres by developing a solvothermal pathway in the presence of pluronic P123 copolymer. During the synthesis, no spherical structures formed in the absence of P123 (Fig. 18b). At a low dosage of P123, the products became solid spheres without hollow structures (Fig. 18c). Hollow spheres formed gradually and became enlarged upon further P123 addition. Finally, Bi₂MoO₆ hollow spheres with diameters of 1-1.5 µm and a thickness of approximately 100 nm (Fig. 18d) were obtained when the P123 amount reached 3 g. The formation of these spheres is illustrated in Fig. 18a. Hollow spherical bismuthal photocatalysts can also be fabricated using

a solvothermal method in the absence of a template. Zhou and colleagues¹⁴³ fabricated hollow Bi₂WO₆ microspheres (Fig. 19) using a template-free solvothermal route with a mixed solvent composed of ethylene glycol (EG) and ethanol. The well-defined hollow Bi₂WO₆ microspheres exhibited considerably better photocatalytic activity in the degradation of RhB than Bi₂WO₆ with other morphologies under visible-light irradiation. Li et al. 144 synthesized hollow Bi₂MoO₆ spheres using a solvothermal route in a Bi₂MoO₆ precursor solution. The obtained yolk shell-like Bi₂MoO₆ displayed markedly higher efficiency in removing RhB. Such improvement was attributed to the hollow structure accompanied by enhanced light absorption, increased specific surface area, and augmented charge transfer. If a mixed solution containing glycerol, ethanol, and deionized water was adopted, hollow Bi₂MoO₆ was still obtained during the solvothermal synthesis. 145 Multishell hollow structures are more attractive than other structural types because of their higher surface area and unique nested structure, which ensure highly efficient light utilization. Zong et al. 146 achieved Bi-V-O heterostructured multishell hollow spheres (composed of interconnected BiVO₄ and Bi₄V₂O₁₁) (Fig. 20) using NaOH-treated carbonaceous microsphere templates and a suitable heating rate. The double-shell Bi-V-O hollow spheres were more effective than their single-shell counterparts in removing MB under visible-light irradiation because of the multiple light reflections and effective light utilization achieved through the multishell hollow structure.

Hollow structures are not restricted to hollow spheres. Hollow tubes and cross-linked net-like structures are also favorable

a OAM +TOPO **OAM** Bi(ACDA)₃ **DMF** +Glucose OAM+DT Bi₂S₃ NPs Bi₂S₃ NPs HDA (Nanosphere) (Nanocrystals) Bi₂S₃ NPs Bi₂S₃ NPs (Hexagonal-nanoplate) (Nanorod) Bi₂S₃ NPs (Ultrathin-nanorod) C b Band gap; 2.17eV 0.8 In light WO, in light $\alpha h v)^2$ Bi_S_NPs(DMF+glucose) Bi_S_ NPs (OAM+ DT) Bi,S, NPs (HDA) Bi,S, NPs (OAM) 0.4 Bi S NPs (OAM + TOPO) 0.2

Fig. 15 (a) Synthesis pathways of Bi₂S₃ NPs with tuned morphologies from Bi(ACDA)₃ (ACDA = 2-aminocyclopentene-1-dithiocarboxylic acid radical) using solvent comprising glucose, oleylamine (OAM), hexadecylamine (HDA), dodecanethiol (DT), trioctylphosphine oxide (TOPO), and dimethylformamide (DMF); (b) photocatalytic degradation curves of MB under visible-light irradiation and (c) the Tauc plot of the Bi₂S₃ NPs using a mixed solvent composed of OAM and TOPO. Reproduced with permission from ref. 135. Copyright 2016 Elsevier.

morphologies for bismuthal photocatalysts. These alternative structures can be constructed with assistance from directed surfactant agents or substrates. For example, Bi₂O₂CO₃ nanotubes (Fig. 21) can grow on a graphene substrate through hydrolysis by adjusting the pH to 10.5 using urea. 147 In the presence of CTAB (as a morphology director), BiVO₄ can grow into nanotubes during hydrothermal synthesis. 148 The formed BiVO₄ nanotubes exhibit improved activity relative to that of BiVO₄ nanorods in the photocatalytic degradation of MO. When dual templates of polystyrene latex spheres (PS) and P123 are adopted in the preparation of Bi₂O₃/TiO₂ composites (sol-gel method), a series of 3D-ordered macroporous profiles are produced (Fig. 22). 149 The composites present a significantly higher activity than P25 and Bi₂O₃ in the photocatalytic removal of crystal violet. Such an advantage can be ascribed to the formed porous structure and heterojunction. Lee et al. 150 fabricated a type of porous β-Bi₂O₃ from Bi(NO₃)₃ using a rapid

20 10

50

Time (min)

phase and surface transformation with structure-guided combustion waves. The compound showed a higher photocatalytic performance in the degradation of RhB than those of α-Bi₂O₃ rods and Bi(NO₃)₃·H₂O rods.

1.2 1.4 1.6 1.8 2.0 2.2 2.4 2.6 2.8 3.0

hv (eV)

Immobilization

During an application, the undesired aggregation of NPs would seriously affect the performance of Bi-based photocatalysts, so an effective separation method should be employed to reduce the aggregation. The cyclic utilization of Bi-based photocatalysts should also be considered. Immobilization has been considered as an effective means to solve these two issues. Furthermore, with the aid of substrate materials, the profile of bismuthal photocatalysts can be tuned to achieve photocatalytic reactions. Hence, suitable immobilizations have been explored for improving the performance of Bi-based photocatalysts.

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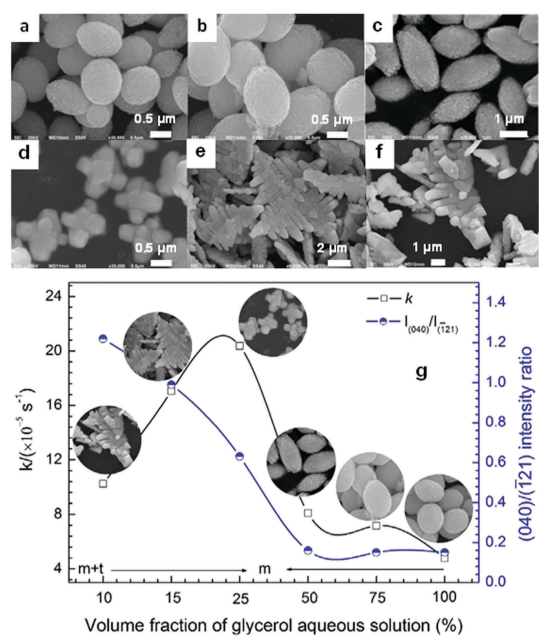


Fig. 16 SEM images of the BiVO₄ samples prepared with the following volume fractions of glycerol aqueous solution: (a) 100%, (b) 75%, (c) 50%, (d) 25%, (e) 15%, and (f) 10%; (g) relationship between the glycerol fraction, the (040)/(121) intensity ratio, the morphology, and the photocatalytic performance of the BiVO₄ samples. Reproduced with permission from ref. 136. Copyright 2016 Elsevier.

Immobilization requires the increased dispersion of bismuthal photocatalysts, which increases the availability of photocatalysts and improves the attachment between the photocatalysts and reactants. Wang and co-workers¹⁵¹ prepared a BiOCl film on a Bi plate using a room-temperature reaction, and the product exhibited a high photocatalytic activity in the degradation of RhB because of the good dispersion of the formed BiOCl nanonuclei. Sonawane and colleagues¹⁵² immobilized Bi₂O₃ on bentonite using an intercalation method. The obtained Bi₂O₃/bentonite composite exhibited improved performance relative to the separate use of Bi₂O₃ and bentonite in the photocatalytic degradation of RhB because of the increased

adsorption capacity brought about by the use of bentonite. Similarly, Wang et al. 153 immobilized phosphotungstic acidmodified BiOBr on the surface of a zeolite. This process endowed the composites with enhanced photocatalytic activity for the degradation of MO. Li et al. 154 immobilized BiOI on diatomite and obtained composites with improved photocatalytic performance in the degradation of RhB. Tong and co-workers 155 used nickel foam as a substrate to immobilize a Bi₂WO₆/TiO₂ composite and increase the efficiency of Bi₂WO₆/TiO₂ in the photocatalytic degradation of RhB. Shi et al. 156 found that the performance of BiVO4 in the photocatalytic removal of tetracycline hydrochloride could be enhanced when supported

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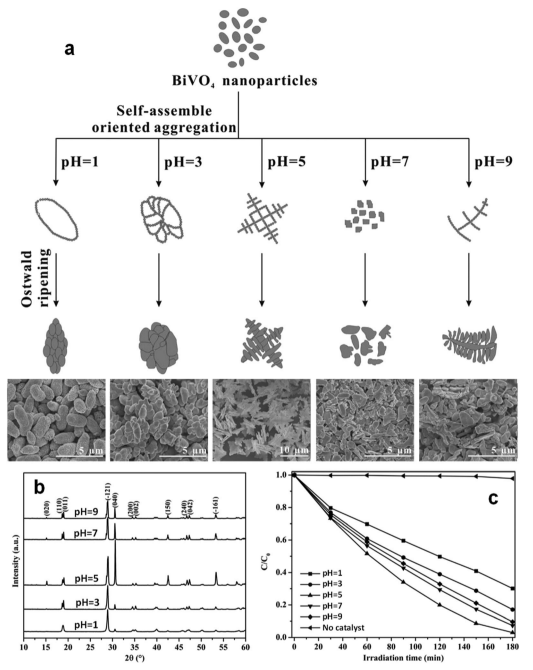


Fig. 17 (a) Schematic diagram of the possible formation mechanism and (b) powder X-ray diffraction patterns (PXRD) patterns of BiVO₄ samples with various morphologies prepared at different pH values; (c) photocatalytic activities of the BiVO₄ samples in the degradation of MB under visible light. Reproduced with permission from ref. 137. Copyright 2016 Elsevier.

on palygorskite. Zhang and colleagues¹⁵⁷ demonstrated that Bi₄Ti₃O₁₂ immobilized on SiO₂ spheres could more effectively decolorize Brilliant Red-X3B because of the good dispersion and smaller size of Bi₄Ti₃O₁₂.

If the substrates are semiconductor materials, a heterojunction will form and facilitate the separation of electrons and holes. Mu and co-workers 158 deposited spindle-like BiVO4 on TiO₂ nanofibers (Fig. 23) using a solvothermal method, and the composite exhibited enhanced visible-light activity than that of a mechanical mixture of BiVO4 and TiO2 in the photocatalytic degradation of RhB. This result could be ascribed not only to the thorough dispersal of the BiVO₄ NPs by immobilization, but also to the improved separation of electrons and holes by the formed heterojunction between BiVO4 and TiO2. Zhang and co-workers¹⁵⁹ reported the enhanced performance of BiOCl nanosheets in the photocatalytic degradation of RhB by immobilizing the BiOCl sheets on TiO₂ arrays over FTO (Fig. 24). This effect was achieved, because the reflection within the ordered array structure endowed the composites with enhanced light absorption. Moreover, the heterojunction between BiOCl and

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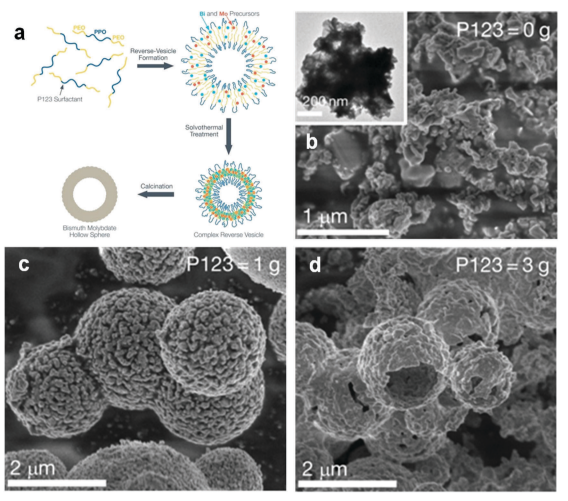


Fig. 18 (a) Schematic diagram of the formation of hollow-structured Bi₂MoO₆ and SEM images of Bi₂MoO₆ fabricated (b) without P123, (c) with 1 g of P123, and (d) with 3 q of P123. Reproduced with permission from ref. 142. Copyright 2016 American Chemical Society.

TiO₂ significantly decreased the charge transfer resistance and facilitated the separation of photo-generated holes and electrons. Kumar et al. 160 prepared Bi₂S₃/TiO₂ composites from Bi₂S₃ nanotubes and TiO₂ NPs. The acquired composites showed improved photocatalytic performance in the degradation of amaranth.

Conductive substrates not only affect the morphologies of the composites, but also act as electron acceptors. For example, when a BiOCl sheet was immobilized on carbon fibers, the performance of the sheet in the photocatalytic degradation of 4-nitrophenol was enhanced due to the good dispersion of BiOCl nanoplates and the electron trapping role of the carbon fibers. 161 Likewise, Weng and co-workers162 loaded Bi4Ti3O12 on carbon fibers using a hydrothermal method with carbon-fiber-supported TiO2 nanosheets as a precursor. The prepared composite exhibited better performance than that of Bi₄Ti₃O₁₂ nanosheets in the photocatalytic degradation of MO. Di et al. 163 deposited small Bi₄O₅Br₂ nanosheets on multi-walled carbon nanotubes (MWCNTs) via an ionic liquidassisted solvothermal method and found that the activity of the composites in the photocatalytic degradation of tetracycline hydrochloride was obviously improved.

Immobilization plays a more important role in gaseous reactions than in other reactions, because well-dispersed

photocatalysts are crucial to the external and internal diffusions of gaseous molecules. Dong's group164 immobilized Bi NPs on g-C₃N₄ and observed that an NO removal efficiency of 60.8% was achieved when Bi NPs of 12 nm were decorated on g- C_3N_4 . This efficiency was higher than that of g- C_3N_4 (38.6%). Meanwhile, Dong's group 165 deposited metal Bi NPs on TiO2 for NO removal and noted that Bi can act as a cocatalyst, similar to a noble metal, due to surface plasmon resonance and can effectively photoactivate TiO2 during NO oxidation under visible-light irradiation. Xia et al. 166 loaded BiOI on porous Al₂O₃ using a hydrothermal method and observed that the obtained BiOI/Al2O3 composite exhibited photocatalytic performance by simultaneously eliminating gaseous NO and SO2.

Heterojunction construction

Heterojunction construction is an effective pathway for enhancing the performance of photocatalysts, because photo-generated electrons and holes can be effectively separated by a heterojunction. 29,42,167-175 For most Bi-based photocatalysts, a narrow band gap ensures that electrons can be excited by visible light.

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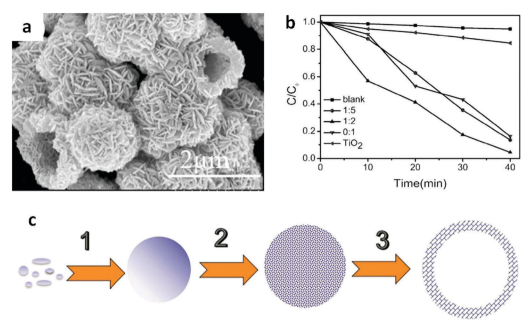


Fig. 19 (a) SEM images of Bi_2WO_6 hollow spheres prepared using a mixed solvent of EG and ethanol (EA) ($V_{EG}: V_{EA} = 1:2$) at 160 °C for 2 h; (b) photocatalytic degradation curves of RhB on Bi_2WO_6 samples prepared using solvents with different V_{EG} : V_{EA} ratios, and (c) schematic diagram of the formation of hollow structures, involving the formation (1), surface roughening (2) and hollowing (3) of amorphous solid microspheres. Reproduced with permission from ref. 143. Open Access 2016 MDPI.

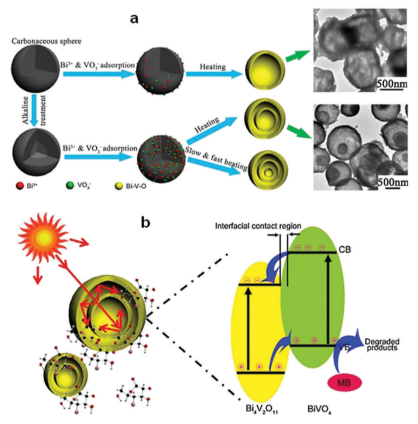


Fig. 20 (a) Schematic diagram of the formation of Bi-V-O multishell hollow spheres and (b) photoabsorption and photocatalytic mechanism of the Bi-V-O multishell hollow spheres. Reproduced with permission from ref. 146. Copyright 2017 Elsevier.

However, the excited electrons recombine with holes soon after. Therefore, heterojunction construction plays a key role in

improving the performance of Bi-based photocatalysts. Bi-Based heterojunctions include conventional and Z-scheme heterojunctions.

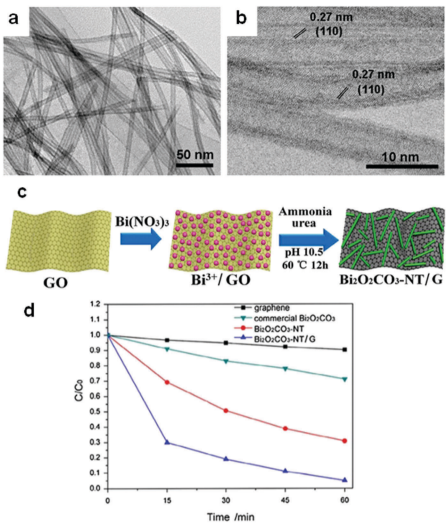


Fig. 21 (a) TEM and (b) HRTEM images and (c) formation mechanism of Bi₂O₂CO₃ nanotubes on a graphene substrate; (d) photocatalytic degradation behavior of samples toward reactive red (X-3B). Reproduced with permission from ref. 147. Copyright 2017 Elsevier.

Among the conventional heterojunctions, the type II junction is the most common one, while in Z-scheme systems, the newlyemerged direct Z-scheme heterojunction appears to be the most effective junction structure used for exploring the capacity of photo-generated carriers.

Conventional type II heterojunctions

Conventional heterojunctions are the most commonly reported in Bi-based photocatalysts, and the two main carrier migrations are illustrated in Fig. 25. In the type I mode, photo-induced holes migrate from the lower VB of semiconductor B (SC-B) to the higher VB of semiconductor A (SC-A), whereas electrons migrate from the higher CB of SC-B to the lower CB of SC-A. During the process, electrons and holes migrate in the same direction, but electrons and holes migrate at different speeds due to their different effective mass or other factors. Consequently, holes and electrons are separated. In the type II mode, holes migrate from the lower VB of SC-B to the higher VB of SC-A, while electrons migrate from the higher CB of SC-A to the lower CB of SC-B. During the process, electrons and holes migrate in opposite directions, resulting in their separation. In any of these two heterojunctions, the electrons and holes can be separated, and the photocatalytic activity of the photocatalysts can be enhanced.

Among the different types of heterojunctions, the type II junction is the most reported one. This kind of heterojunction usually forms between bismuthal compounds and semiconductors with a narrow band gap, when the CB and VB of one semiconductor are lower than that of the coupled one. For example, Zhu and co-workers¹⁷⁶ reported the fabrication of a binary Bi and non-Bi Bi₄Ti₃O₁₂/CeO₂ composite by coupling Bi₄Ti₃O₁₂ with CeO₂ using a molten salt and ion-impregnation method. The obtained Bi₄Ti₃O₁₂/CeO₂ composite exhibited enhanced activity in the photocatalytic degradation of BPA. This phenomenon can be attributed to the important role of the heterojunction between Bi₄Ti₃O₁₂ and CeO₂, which facilitates carrier separation (Fig. 26). In addition, Fan et al. 177 constructed a binary Bi-based Bi₂MoO₆/BiOI composite heterojunction (Fig. 27a and b) using an anion exchange method with BiOI as a precursor. The as-prepared Bi₂MoO₆/BiOI exhibited

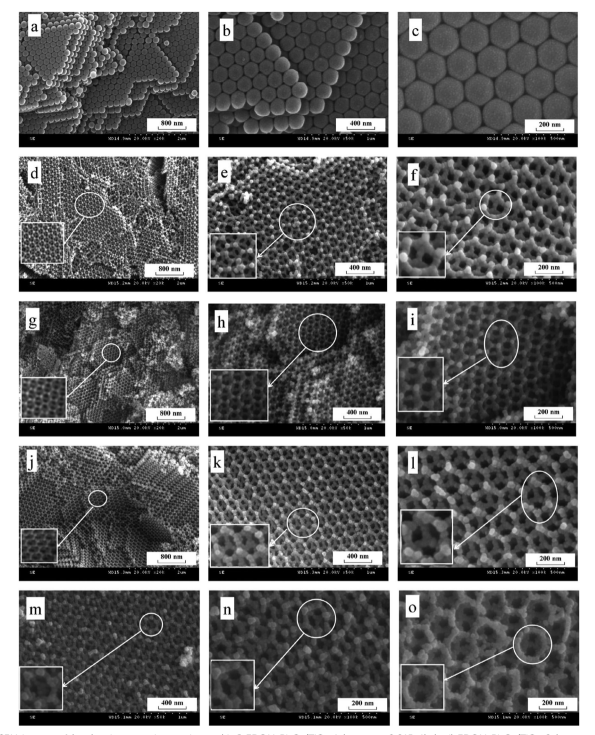


Fig. 22 SEM images of (a-c) polystyrene latex spheres, (d-f) 3DOM Bi_2O_3/TiO_2-1 ($n_B:n_{Ti}=0.015:1$), (g-i) 3DOM Bi_2O_3/TiO_2-2 ($n_{Bi}:n_{Ti}=0.03:1$), (j-l) 3DOM Bi₂O₃/TiO₂-3 (n_{Bi} : n_{Ti} = 0.06:1) and (m-o) 3DOM Bi₂O₃/TiO₂-4 (n_{Bi} : n_{Ti} = 0.09:1). Reproduced with permission from ref. 149. Copyright 2015 Elsevier.

improved photocatalytic activities in the degradation of RhB (Fig. 27c). The sample with a Mo/I molar ratio of 50% exhibited the best activity under visible light excitation due to the formation of a type II heterojunction between Bi₂MoO₆ and BiOI (Fig. 27d). This type of charge migration and separation were also observed for heterojunctions formed between ZnO/BiOI

and BiOCl/Bi₃PO₄ couples. 178,179 If three matched bismuthal semiconductors are coupled together, a ternary heterojunction such as $Bi_2S_3/Bi_2O_3/Bi_2O_2CO_3$ can be constructed that exhibits a better performance. 180 The enhancement in the activity of Bi₂S₃/Bi₂O₃/Bi₂O₂CO₃ is ascribed to the increased light absorption and efficient charge separation by the double type II

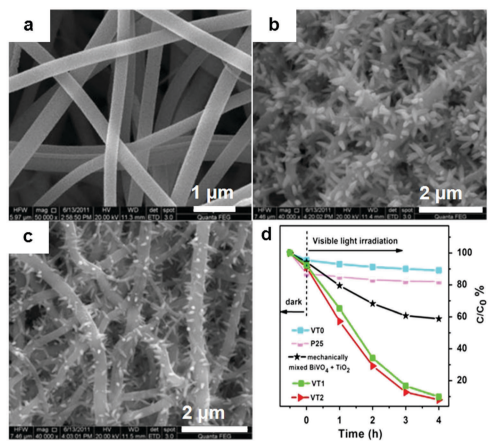


Fig. 23 SEM images of (a) TiO₂ nanofibers (VT0), (b and c) BiVO₄ immobilized on TiO₂ nanofibers using a low (VT1) and high (VT2) concentration of raw material solution, and (d) the degradation curves of RhB over different photocatalysts under visible light. Reproduced with permission from ref. 158. Copyright 2016 Elsevier.

heterojunction (Fig. 28). Type II carrier migrations can also occur in homojunctions. In a homojunction, the interface between two homogenous semiconductors forms due to different phase structures. 181,182

Numerous preparation strategies for conventional heterojunctions (mainly Type II) have been reported, and most of these techniques have revealed enhanced photocatalytic activities (Table 1). Nevertheless, conventional heterojunctions have evident limitations. Photo-generated electrons migrate from a more negative CB to a less negative CB and the holes from a more positive VB to a less positive VB. This type of migration for electrons and holes reduces the redox capacity of the photocatalyst. Consequently, any improvement in the performance of the photocatalysts will be significantly limited.

Given the limited improvement of a simple conventional heterojunction, several strategies combining heterojunction and morphology control, such as the preparation of a MoS₂/ Bi₁₂O₁₇C_{l2} 2D bilayer heterostructure¹⁸³ and a Bi₂S₃ nanorods/ Bi₂O₃ microtubes composite, ¹⁸⁴ have been developed to further enhance the performance of bismuthal photocatalysts. In our previous work, we found that heterojunctions can form between two adjacent (001) and (110) facets of BiOI, whereas their CBs and VBs are located in different positions. Consequently, BiOI with a hierarchical morphology and an optimal ratio of (001)

and (110) facets was found to possess the highest photocatalytic activity.185 However, the combination of heterojunction and morphology control is still unable to overcome the inherent limitations of a conventional heterojunction, although this characteristic results in the better performance of as-prepared photocatalysts.

Direct Z-scheme heterojunctions

In 2013, Yu et al. proposed a direct Z-scheme heterojunction concept to explain the enhancement in the photocatalytic activity of a ${
m TiO_2/g\text{-}C_3N_4}$ composite photocatalyst. 285 This direct Z-scheme heterojunction is different from conventional liquidphase Z-scheme heterojunctions and all-solid-state Z-scheme heterojunctions. The direct Z-scheme heterojunction does not require an electron medium.²⁸⁶⁻²⁹⁰ The transfer of charge carriers in this heterojunction is through the built-in electric field between the interface of SC-A and SC-B (Fig. 29a). However, the transfer of charge carriers in the liquid-phase (Fig. 29b) and all-solid-state (Fig. 29c) Z-scheme heterojunction is through the use of ions in the solution and noble metal NPs as electron conductors, respectively. 291-295

The structure of a direct Z-scheme heterojunction is similar to that of a type-II heterojunction (Fig. 25b) but with different charge carrier transport mechanisms. In a typical type-II

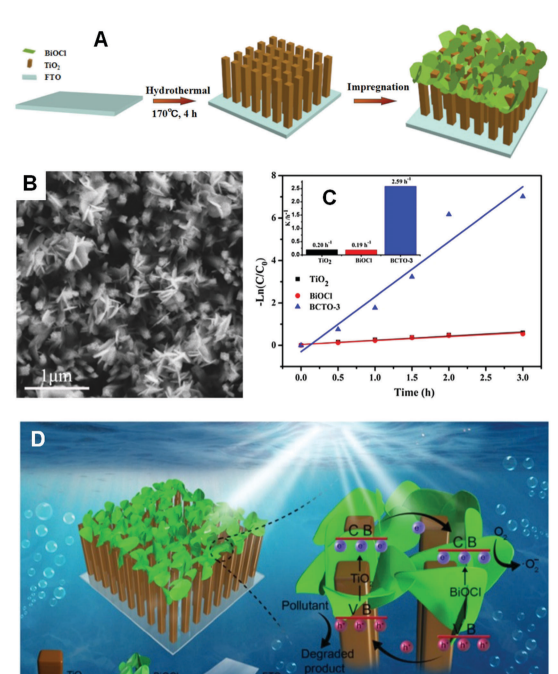


Fig. 24 (A) Schematic diagram of BiOCl/TiO₂ composite formation; (B) FESEM images of a BiOCl/TiO₂ composite (BCTO-3); (C) kinetic curves of RhB photocatalytic degradation on TiO2, BiOCl, and BCTO-3; (D) schematic diagram of the reaction mechanism over the immobilized BiOCl/TiO2 composite under visible light. Reproduced with permission from ref. 159. Copyright 2017 Elsevier.

heterojunction photocatalyst, SC-A has higher CB and VB positions than SC-B. Under light irradiation, the electrons in SC-A and holes in SC-B will transfer to the CB of SC-B and VB of SC-A, respectively. This process results in the spatial separation of the electron-hole pairs. However, the reduction ability of photo-generated electrons in the CB of SC-B and the oxidation ability of holes in the VB of SC-A are greatly reduced.

In contrast, the direct Z-scheme heterojunction exhibits a completely different mechanism of charge carrier migration. A photo-generated electron with a low reduction potential in SC-B will recombine with a photo-generated hole with a low oxidation potential in SC-A (Fig. 29a). Finally, the photo-generated electron and hole will remain in the CB of SC-A and VB of SC-B, respectively. The direct Z-scheme heterojunction will result in

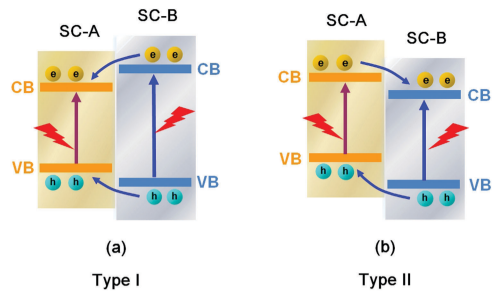


Fig. 25 Schematic diagram of the carrier migrations in conventional (a) type I and (b) type II heterojunctions between semiconductor A (SC-A) and semiconductor B (SC-B).

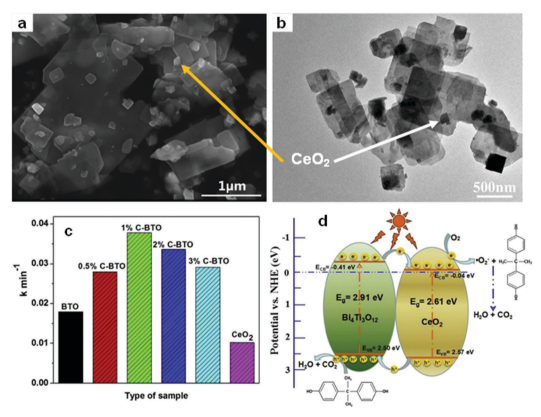


Fig. 26 (a) SEM and (b) TEM images of CeO₂/Bi₄Ti₃O₁₂ composites (1%C-BTO); (c) comparison of pseudo-first-order kinetic constants of different photocatalysts for BPA; and (d) a schematic illustration of the charge migration in the 1%C-BTO composites under light irradiation. Reproduced with permission from ref. 176. Copyright 2016 Elsevier.

the spatial separation of useful charge carriers, remove useless charge carriers, and optimize the redox ability of the SC-A and SC-B composition system. Therefore, an enhanced photocatalytic performance can be observed in a direct Z-scheme heterojunction photocatalyst. In addition, the transfer mechanism of the charge carriers in a direct Z-scheme heterojunction is also different from that of conventional liquid-phase and all solid-state Z-scheme heterojunctions. No electron conductor is Review Nanoscale Horizons

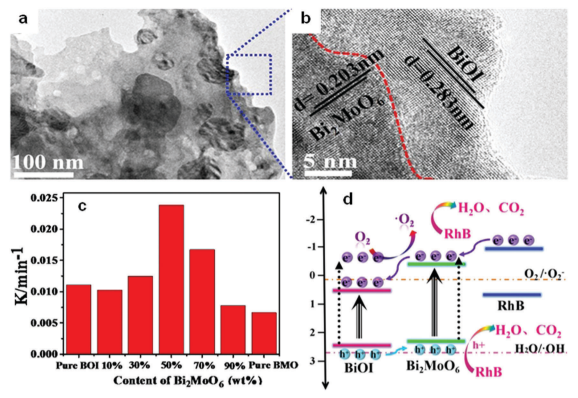


Fig. 27 (a) TEM images and (b) the corresponding HRTEM images of Bi₂MoO₆/BiOI composites (BMO/BOI = 50%); (c) reaction rate constants for RhB degradation on BiOI (BOI), Bi₂MoO₆ (BMO) and BMO/BOI; and (d) a schematic diagram showing the charge transfer of a BMO/BOI heterostructure. Reproduced with permission from ref. 177. Open Access 2016 World Scientific.

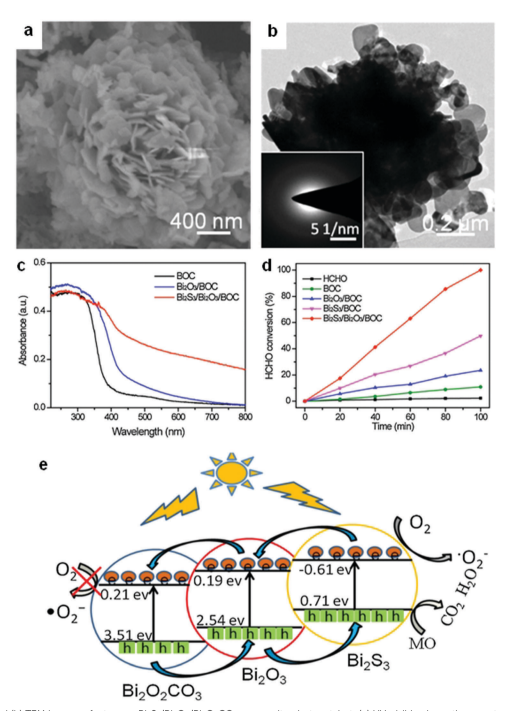
needed in a direct Z-scheme heterojunction, and the charge carrier can directly migrate using the built-in field at the SC-A and AC-B interfaces. Moreover, the fabrication cost of the direct Z-scheme photocatalyst is greatly reduced. Significantly, the direct Z-scheme heterojunction offers at least two obvious merits, namely, a low fabrication cost and high redox ability.

Many Bi-based direct Z-scheme heterojunction photocatalysts have been prepared and reported. For example, Liu et al. 296 prepared a CdTe/Bi₂S₃ direct Z-scheme heterojunction photocatalyst using an electrostatic adsorption and self-assembly method. The obtained CdTe/Bi₂S₃ exhibited higher photocurrent responses and microcystin-LR sensitivity than the individual CdTe and Bi2S3 nanorod components. Meanwhile, Dai and colleagues²⁹⁷ formed a g-C₃N₄/Bi₂MoO₆ direct Z-scheme heterojunction between g-C₃N₄ and Bi₂MoO₆ using a hydrothermal method. This g-C₃N₄/Bi₂MoO₆ photocatalyst exhibited a higher activity than the individual g-C₃N₄ and Bi₂MoO₆ components in the photocatalytic degradation of MB (Fig. 30). Ullah and coworkers²⁹⁸ reported the synthesis of a Se/BiVO₄ direct Z-scheme heterojunction photocatalyst and its higher photocurrent density than that of its individual Se and BiVO4 components. Huang and colleagues²⁹⁹ demonstrated a direct Z-scheme electron migration mechanism between g-C₃N₄ and BiOIO₃ in their prepared g-C₃N₄/BiOIO₃ composite under UV-visible light irradiation. The photocatalytic activity of the g-C₃N₄/BiOIO₃ composite was shown to be markedly higher than those of g-C₃N₄ and mechanically mixed g-C₃N₄/BiOIO₃ for the removal of RhB. We recently fabricated a Bi₂O₃/g-C₃N₄ direct Z-scheme heterojunction photocatalyst using an *in situ* room-temperature approach, including photoreduction deposition of Bi3+ and subsequent air-oxidation of the resultant metallic Bi. 300 In the prepared composite, Bi₂O₃ quantum dots were uniformly distributed on the surface of g-C₃N₄ sheets (Fig. 31a), and the photocatalytic activity of Bi₂O₃/g-C₃N₄ was found to be higher than those of pure Bi₂O₃ and g-C₃N₄ in the photocatalytic degradation of phenol under visible light (Fig. 31b). This phenomenon can be attributed to the formed direct Z-scheme heterojunction (Fig. 31c). More reports on direct Z-scheme heterojunctions are listed in Table 2 for reference.

Surface modification

Surface modification is an effective and widely used method for enhancing the activity of a photocatalyst.314 It also plays an important role in improving the performance of Bi-based photocatalysts. An extensive range of materials can be adopted to modify surfaces. Carbon-based materials, metals, ions, polymers, and semiconductors can be used to modify bismuthal photocatalysts. If a semiconductor is used as a modifier, a nanosized heterojunction will be formed, and the carriers will transfer faster in the tiny heterojunction than in bulk junctions because of the larger interface ratio. For example, the dandelion-like MoS₂-decorated Bi₂S₃ prepared by Li et al.³¹⁵ exhibited improved

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 $\textbf{Fig. 28} \hspace{0.2cm} \textbf{(a) SEM and (b) TEM images of a ternary } Bi_2S_3/Bi_2O_3/Bi_2O_2CO_3 \hspace{0.2cm} \textbf{composite photocatalyst; (c) UV-visible absorption spectra of } Bi_2O_2CO_3(BOC), \\ \textbf{(b) TEM images of a ternary } Bi_2S_3/Bi_2O_3/Bi_2O_2CO_3 \hspace{0.2cm} \textbf{(c) UV-visible absorption spectra of } Bi_2O_2CO_3(BOC), \\ \textbf{(c) UV-visible absorption } Bi_2O_2CO_3(BOC), \\ \textbf{(d) SEM and (d) TEM images of a ternary } Bi_2S_3/Bi_2O_3/Bi_2O_3/Bi_2O_3(BOC), \\ \textbf{(d) SEM and (d) TEM images } Bi_2O_3/Bi$ $Bi_2O_3/BOC, \text{ and } Bi_2O_3/BOC; \text{ (d) comparison of the photocatalytic HCHO removal of } Bi_2O_2CO_3(BOC), \\ Bi_2O_3/BOC, Bi_2O_3/BOC, \text{ and } Bi_2S_3/Bi_2O_3/BOC, \\ Bi_2O_3/BOC, \text{ and } Bi_2S_3/Bi_2O_3/BOC, \\ Bi_2O_3/BOC, \text{ and } Bi_2O_3/BOC, \\ Bi_2O_3/BOC, \text{ and } Bi_2O_3/BOC, \\ Bi_2O_3/BOC, \text{ and } Bi_2O_3/BOC, \\ Bi_2O_3$ BOC; and (e) a schematic diagram showing the charge-transfer process in ternary Bi₂S₃/Bi₂O₂CO₃ double heterojunctions. Reproduced with permission from ref. 180. Copyright 2016 Elsevier.

photocatalytic performance in the degradation of RhB due to the tiny heterojunctions formed between MoS₂ and Bi₂S₃ (Fig. 32). Further to this, SnO₂ modified BiVO₄ was found to be able to reduce CO₂ to CH₄, with the introduced SnO₂ acting as a redox reaction platform capable of accepting the photogenerated electrons from BiVO₄ nanoplates during the conversion of CO₂. 316 AgBr, graphene-like BN (g-BN), and some bismuthal

compounds with a narrow band gap have also been reported as semiconductor modifiers for Bi-based photocatalysts (Table 3).

Coupling bismuthal photocatalysts with conductive materials is another effective surface modification strategy. The conductive materials usually act as electron acceptors to promote the migration of photo-generated electrons and consequently facilitate the separation of carriers. Metals, carbon-based materials,

 Table 1
 Conventional heterojunction strategies of nanoscale Bi-based photocatalysts

Bi, O ₁	Bismuthal compound	Coupled material	Method	Application	Ref.
Bi_O_0	· ·		~	1-11	
Bi-O ₂		-		2	186
Bi.Co.					187
Bis.5					188
Bis.S.					189
Bis Policy Poli					190
Bio.Co.co					191
Bi_TiO_0 manshees					192
Bi_17i0_0_ g-C_3N,					193
Bil Tilo 2		_			194
Bi_17_0_1 CeO_					195
Bi,O,CO,					196
Bi,O,CO ₂ Åg,PO ₄ Hydrothermal method and precipitation RhB 1 BiFeO ₂ CuO Hydrothermal synthesis and impregnation RhB 2 BiFeO ₂ Ling Hydrothermal synthesis and impregnation RhB 20 BiFeO ₂ Ling Impregnation RhB 20 BiVO ₄ TiO ₅ fibers Hydrothermal method Brilliant red X-3B 20 BiVO ₄ 3DOM TiO ₂ Hydrothermal method Brilliant red X-3B 20 BiVO ₄ 2CO ₂ Coprecipitation and evaporation RhB 20 BiVO ₄ 2CO ₂ Coprecipitation and evaporation MO 22 BiVO ₄ 2CO ₂ Coprecipitation and evaporation MO 22 BiVO ₄ 2CO ₂ Coprecipitation and evaporation MO 22 BiVO ₄ 2CO ₂ Copol Impregnation and evaporation MO 22 BiVO ₄ 2CO ₂ Ag-0 Impregnation and evaporation MO 22 BiVO ₄ 2CO ₂ Agloa					176
BiFeO ₂					197
BiFeO, CuO Hydrothermal synthesis and impregnation RhB 20					198
Bishago, Ing.O, Ing.O, Impregnation RhB and p-thlorophenol 20 BIOCOOPH Ags.O Solvothermal deposition RhB and p-thlorophenol 20 BIOVO, TiOp fibers Hydrothermal method RhB 20 BIOVO, G.Ci.N, Ultrasonic assembly RhB 20 BIOVO, G.Co.N, Ultrasonic assembly RhB 20 BIOVO, Ags.O Impregnation and subsequent annealing RhB 20 BIOVO, Ags.O Impregnation and evaporation RhB 20 BIOVO, Ags.O Impregnation and evaporation RhB 20 BIOVO, Co.O.O, Hydrothermal method MB 20 BIOVO, SinO_2 Hydrothermal method Water oxidation 21 BIOVO, CO.O.O, Drop casting method Water oxidation 21 BISOO_C TiO_1 (Hydrothermal method RhB and MrO 21 BISOO_C TiO_2 (Hydrothermal method RhB and phenol 21 BISOO_C TiO_3 (Hydrothermal method RhB 20 BISOO_C					199
BiOCOOFH		CuO			200
BiVO ₄ TÖ ₂ fibers Hydrothermal method Brilliant red X-3B 2c BiVO ₄ g-G,N, Ultrasonic assembly CO ₂ reduction 2c BiVO ₄ g-G,N, Ultrasonic assembly CO ₂ reduction 2c BiVO ₄ Ag ₂ O Impregnation and subsequent annealing MO 2c BiVO ₄ SnO ₂ Impregnation and evaporation MO 2c BiVO ₄ SnO ₂ Hydrothermal method MB 2c BiVO ₄ SnO ₂ Hydrothermal method Water oxidation 21 BiVO ₄ PEDOT-PSS Electrodeposition Photo-photocurrent 21 Bi ₂ WO ₆ TO ₂ fibers Hydrothermal method Rh B and MO 21 Bi ₂ WO ₆ TO ₂ fibers Hydrothermal method Rh B and phenol 21 Bi ₂ WO ₆ WO ₃ Hydrothermal method Rh B 21 Bi ₂ WO ₆ CeO ₂ Hydrothermal method Rh B 21 Bi ₂ WO ₆ Ag ₂ Cl Hydrothermal method Rh B 22 <td></td> <td></td> <td></td> <td></td> <td>201</td>					201
BNO ₀ 3DOM TiO ₂ Hydrothermal method RhB 2c BNO ₀ C _C O ₂ Coprecipitation and subsequent annealing RhB 2c BNO ₀ C _C O ₂ Coprecipitation and subsequent annealing RhB 2c BNO ₀ ZnO Annealing of mixture RhB 2c BNO ₀ SnO ₀ Hydrothermal method MB 2c BNO ₀ C _O O ₀ Drop-casting method Water oxidation 2c BNO ₀ TiO ₂ Hydrothermal method RhB and MO 2c BNO ₀ TiO ₃ Hydrothermal method RhB and MO 2c BNO ₀ TiO ₄ Hydrothermal method RhB and MO 2c BNO ₀ TiO ₂ Hydrothermal method RhB and MO 2c BNO ₀ TiO ₂ Hydrothermal method RhB and MO 2c BNO ₀ Hydrothermal method RhB and MO 2c BNO ₀ Hydrothermal method RhB 2c BNO ₀ AgCl	BiOCOOH				202
BiVO ₄ $g_{-}C_3N_s$ Ultrasonic assembly C_2 reduction C_2 Coprecipitation and subsequent annealing C_2 BiVO ₄ Ag_2O Impregnation and evaporation C_2 MO C_2 C_3 C_3 C_4 C_4 C_4 C_5	$BiVO_4$		Hydrothermal method	Brilliant red X-3B	203
BiNO ₂ Ceo. Coprecipitation and subsequent annealing RhB 22 BiNO ₄ ZnO Annealing of mixture RhB 22 BiNO ₄ ZnO Annealing of mixture RhB 22 BiNO ₄ SnO ₂ Hydrothermal method MB 22 BiNO ₄ TiO. Drop-casting method Water oxidation 21 Bi, WO ₈ TiO. Hydrothermal method RhB and MO 22 Bi, WO ₈ TiO. Hydrothermal method RhB and phenol 22 Bi, WO ₈ WO ₁ Hydrothermal method MB 21 Bi, WO ₈ VP ₆ O ₂ Electrospinning with sintering RhB 22 Bi, WO ₈ AgCl Hydrothermal method RhB 22 Bi, WO ₉ GeC ₈ and ZnS Surface	$\mathrm{BiVO_4}$	3DOM TiO ₂		RhB	204
BINO, Ag,O Impregnation and evaporation MO 22 BINO, SnO, Hydrothermal method MB 22 BINO, SnO, Hydrothermal method MB 22 BINO, TO Annealing of misture RhB 22 BINO, SnO, Hydrothermal method MB MS 22 BINO, TO, Drop-asting method MB MS 22 BINO, MTO PEDOT-PSS Electrodeposition Photo-photocurrent 22 BINO, TO, Hydrothermal method RhB and MNO 21 BINNO, TO, Fibers Hydrothermal method RhB and Phonol 22 BINO, MNO, TO, Fibers Hydrothermal method RhB and Phonol 22 BINNO, MNO, Hydrothermal method RhB MS 22 BINNO, MNO, Hydrothermal method RhB 22 BINNO, MNO, AgCl Hydrothermal method RhB 22 BINNO, AgCl Solvothermal method RhB 22 BINNO, AgCl Solvothermal method RhB 22 BINNO, AgCl Solvothermal method and precipitation RhB 22 BINNO, AgCl Hydrothermal method and precipitation RhB 22 BINNO, AgCl Precipitation-deposition RhB BINNO, AgCl Prec	$\mathrm{BiVO_4}$	$g-C_3N_4$	Ultrasonic assembly	CO ₂ reduction	205
BiVO ₄	BiVO_4	CeO ₂	Coprecipitation and subsequent annealing	RhB	206
BiVO ₄	BiVO ₄	Ag_2O	Impregnation and evaporation	MO	207
BiVO ₄ SnO ₂ Hydrothermal method MB 22 BiVO ₄ FTO PEDOT:PSS Electrodeposition Photo-photocurrent 21 BiVO ₄ TiO ₅ PEDOT:PSS Electrodeposition Photo-photocurrent 22 Bi ₂ WO ₄ TiO ₅ Hydrothermal method RhB and phenol 22 Bi ₃ WO ₄ TiO ₅ fibers Hydrothermal method RhB and phenol 22 Bi ₃ WO ₄ WO ₆ TiO ₅ fibers Hydrothermal method RhB and phenol 22 Bi ₃ WO ₄ WO ₆ Hydrothermal method RhB and phenol 22 Bi ₃ WO ₆ WO ₇ Hydrothermal method RhB RhB 22 Bi ₃ WO ₆ CoO ₅ Hydrothermal method RhB RhB 22 Bi ₃ WO ₆ AgC Chemical precipitation RhB RhB 22 Bi ₃ WO ₆ AgC Chemical precipitation RhB RhB 22 Bi ₃ WO ₆ AgC Chemical precipitation RhB RhB 22 Bi ₃ WO ₆ AgC Chemical precipitation RhB 22 Bi ₃ WO ₆ CdS and ZnS Surface functionalization RhB 22 Bi ₃ MO ₆ TiO ₂ Solvothermal method RhB RhB 22 Bi ₃ MO ₆ TiO ₂ Solvothermal method RhB RhB 22 Bi ₃ MO ₆ TiO ₂ Solvothermal method RhB RhB 22 Bi ₃ MO ₆ Ag ₃ R Precipitation and thermal RhB 22 Bi ₃ MO ₆ Ag ₃ R Precipitation Agostition RhB 22 Bi ₃ MO ₆ Ag ₃ R Precipitation-deposition RhB Ag ₃ WO ₄ Hydrothermal method and precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation-deposition RhB and BPA 22 Bi ₃ MO ₆ Ag ₄ R Precipitation-deposition RhB and BPA 22 Bi ₃ MO ₆ Ag ₄ R Precipitation-deposition RhB and BPA 22 Bi ₃ MO ₆ Ag ₄ R Precipitation-deposition RhB and BPA 22 Bi ₃ MO ₆ Ag ₄ R Precipitation-deposition RhB and BPA 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ Ag ₄ R Precipitation RhB 22 Bi ₃ MO ₆ R	BiVO ₄			RhB	208
BiVO, Co,O, Depotasting method Water oxidation 21 BiVO/FTO PEDOT:PSS Electrodeposition Photo-photocurent 22 Bi,WOc, TiO, Hydrothermal method RhB and MO 21 Bi,WOc, TiO, fibers Hydrothermal method RhB and MO 22 Bi,WOc, WO, Hydrothermal method RhB and phenol 22 Bi,WOc, WO, Hydrothermal method RhB and phenol 22 Bi,WOc, WO, Hydrothermal method RhB 22 Bi,WOc, WO, Hydrothermal method RhB 22 Bi,WOc, WO, Hydrothermal method RhB 23 Bi,WOc, WO, AgCI Hydrothermal method RhB 23 Bi,WOc, AgCI Hydrothermal method RhB 21 Bi,WOc, AgCI Hydrothermal method RhB 22 Bi,WOc, AgCI Hydrothermal method RhB 22 Bi,WOc, AgCI Hydrothermal method RhB 23 Bi,WOc, CdS and ZnS Surface functionalization RhB 23 Bi,WOc, TiO, Solvothermal method Phenol and nitrobenzene 22 Bi,MOCo, TiO, Solvothermal method Phenol and nitrobenzene 22 Bi,MOCo, AgR Precipitation-deposition RhB 22 Bi,MOCo, AgI Precipitation-deposition RhB 24 Bi,MOCo, AgI Precipitation-deposition RhB 24 Bi,MOCo, AgI Precipitation-deposition RhB 25 Bi,MOCo, AgI Precipitation-deposition RhB 26 Bi,MOCo, AgI Precipitation-deposition RhB 26 Bi,MOCo, AgI Precipitation-deposition RhB 26 Bi,MOCo, AgI Precipitation-deposition RhB 27 Bi,MOCo, AgI Precipitation-deposition RhB 28 Bi,MOCo, AgI Precipitation-deposition RhB 28 Bi,MOCo, AgI Precipitation-deposition RhB 28 Bi,MOCo, AgI Precipitation-deposition RhB 29 Bi,MOCo, AgI Precipitation-deposition RhB 29 Bi,MOCo, AgI Precipitation-deposition RhB 22 Bi,MOCo, AgI Precipitation-deposition RhB 22 Bi,MOCo, AgI Precipitation-deposition RhB 22 Bi,MOCo, AgI Precipitation-deposition RhB 23 Bi,MOCo, AgI Precipitation-deposition RhB 24 Bi,MOCo, AgI Precipitation		SnO_2		MB	209
$ \begin{array}{c ccccccccccccccccccccccccccccccccccc$				Water oxidation	210
Bi ₃ WO ₆	BiVO ₄ /FTO			Photo-photocurrent	211
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Bis WOok (001) TiO (001) Hydrothermal method MB 21 Bis WOok WO3 Hydrothermal method RhB 21 Bis WOok C≠Fe_O_3 Electrospinning with sintering RhB 21 Bis WOok AgCl Hydrothermal method RhB 21 Bis WOok AgCl Hydrothermal method RhB 21 Bis WOok Ag_CO Chemical precipitation RhB 22 Bis WOok Ag_SO CDC and ZnS Surface functionalization RhB 22 Bis MOOk TiO2 Solvothermal method Phenol and nitrobenzene 22 Bis MOOk TiO2 Solvothermal method and precipitation RhB 22 Bis MOOk AgBr Precipitation-deposition RhB 22 Bis MOOk AgBr Precipitation-deposition RhB Band BPA 22 Bis MOOk MOO3 Hydrothermal method Photoanode 22 Bis MOOk MOO3 Hydrothermal method RhB 23 Bis OCl </td <td></td> <td></td> <td></td> <td></td> <td>213</td>					213
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	Bi_2O_3 - Bi_2S_3		Hydrothermal method	RhB	184

Table 1 (continued)

Bismuthal compound	Coupled material	Method	Application	Ref.
Bi ₂ O ₃ /BiVO ₄		Solvothermal synthesis followed by annealing	Phenol	246
Bi ₂ O ₃ /BiVO ₄		Hydrothermal method	RhB	247
Bi ₂ O ₃ QDs/BiVO ₄ fibers		Direct heat treatment	RhB	248
Bi_2WO_6/Bi_2O_3		Solid-state reaction	RhB	249
Bi ₂ O ₃ /BiOCl		Alkaline treatment	MO	250
β-Bi ₂ O ₃ /BiOI		In situ reaction	MO	251
Bi ₅ O ₇ I/Bi ₂ O ₃		Chemical etching	Malachite green	252
Bi ₂ S ₃ /BiOCl		Solvothermal synthesis	Salicylic acid, RhB	253
Bi_2S_3/Bi_2WO_6		Anion exchange approach	RhB	254
Bi_2S_3/Bi_2WO_6		Hydrothermal method	Reduction of Cr(VI)	255
Bi_2S_3/Bi_2VVO_6 $Bi_2S_3/Bi_4Ti_3O_{12}$ nanofibers		In situ ion exchange	RhB	256
Bi ₂ S ₃ /Bi ₂ O ₂ CO ₃		Anion exchange	Gaseous NO	257
Bi ₂ S ₃ /Bi ₂ O ₂ CO ₃ hollow		One-pot room temperature route	RhB	258
microspheres		one por room temperature route	MID	230
Bi ₂ Ti ₂ O ₇ /Bi ₄ Ti ₃ O ₁₂		One-step molten salt method	MO, RhB	259
Bi ₁₂ Ti ₂ O ₇ /Bi ₄ Ti ₃ O ₁₂ Bi ₁₂ TiO ₂₀ /Bi ₂ WO ₆		Hydrothermal method	RhB	260
		Hydrothermal method	RhB	261
BiVO ₄ /Bi ₂ O ₂ CO ₃			Gaseous <i>o</i> -dichlorobenzene	262
BiPO ₄ /BiOBr		Mixing in solvent		
Bi ₂ MoO ₆ /BiVO ₄		Spin-coating	Photoelectrode	263
Bi ₂ MoO ₆ /BiPO ₄		Hydrothermal method	RhB	264
$Bi_2W_xMo_{1-x}O_6/BiOCl$		Solvothermal method	RhB	265
Bi _{3.64} Mo _{0.36} O _{6.55} /Bi ₂ MoO ₆		Hydrothermal method	RhB	266
BiOI/BiVO ₄		Coprecipitation	Pseudomonas aeruginosa	267
BiOI/BiVO ₄		Precipitation-deposition	MO	268
BiOCl/BiVO ₄		Coprecipitation method	RhB	269
BiOCl/Bi ₁₂ O ₁₇ C _{l2}		Hydrothermal method	MO	270
BiOI/Bi ₂ MoO ₆		Precipitation-deposition	BPA	271
Bi ₂ MoO ₆ /BiOI		hydrothermal method	RhB	177
Bi ₂₄ O ₃₁ Cl ₁₀ /BiOCl		Phase transformation by annealing	Conversion of benzyl alcohol	272
$\mathrm{Bi_4O_5I_2/Bi_5O_7I}$		In situ phase transformation	BPA and RhB	273
$\mathrm{Bi_4O_5I_2/Bi_5O_7I}$		Hydrothermal method	Propylparaben	274
Bi ₂ O ₂ CO ₃ /BiOI		Pore impregnation	RhB	275
Bi ₂ O ₄ -Decorated BiOBr		Alkali posttreatment assisted light	MO	276
		irradiation		
Monoclinic BiVO ₄ /tetragonal		Hydrothermal method	RhB	277
BiVO ₄ BiVO ₄ /Bi ₄ V ₂ O ₁₁		Precursor conversion	Photoelectrodes	278
Ternary heterojunctions		** 1 .1 .1 .1 .1	0 1 1 1 1 1 1	
$Bi_2O_3/Bi_2S_3/MoS_2$		Hydrothermal method	O ₂ evolution and MB degradation	279
AgI/BiOI-Bi ₂ O ₃		Etching-deposition	Cr(v1) reduction	280
$BiOCl_x/BiOBr_y/BiOI_z$		Electrospinning and the sol-gel methods	Trichloroethylene	281
Bi ₂ S ₃ /Bi ₂ O ₃ /Bi ₂ O ₂ CO ₃		Heat treatment and ion exchange	HCHO, MO, and phenol	180
Bi ₇ O ₉ I ₃ /AgI/AgIO ₃		Chemical deposition	MO and gaseous NO	282
Bi ₇ O ₉ I ₃ /Bi ₅ O ₇ I/g-C ₃ N ₄		Hydrothermal method	Crystal violet	283
BiOBr/Co(OH) ₂ /PVP		Solvothermal synthesis	MO	284

Rhodamine B, RhB; methylene blue, MB; methyl orange, MO; bisphenol A, BPA; ciprofloxacin, CIP; PEDOT:PSS = Poly(3,4-ethylenedioxythiophene):poly-(styrenesulfonate).

conductive polymers, and metallic Bi have been adopted as electron scavengers to enhance the photocatalytic activities of bismuthal compounds (Table 3). Commonly employed noble metals include Au, Ag, Pt, and Pd. Hirakawa et al.317 obtained a composite by loading Au NPs on BiVO₄. The prepared Au/BiVO₄ exhibited an enhanced production of H2O2 from water than that of pure BiVO₄ under visible-light irradiation due to the selective two-electron reduction of O₂ by Au. In addition to noble metals, other metals can also serve as electron trapping agents. Park and co-workers318 decorated Ni NPs on W-doped BiVO4 nanofibers and obtained a fibrous composite (Ni@NiO/W:BiVO4 NFs). The process was followed by a calcination method, during which Ni NPs were partly oxidized to NiO. The as-prepared composite

exhibited a higher photocurrent and O2 evolution than W-doped BiVO₄ nanofibers and Pt-decorated fibers in the photocatalytic oxidation of water. The enhancement was attributed to the unique structure of the Ni@NiO decoration, in which the Ni metal traps the photo-generated electrons and NiO accepted holes. Moreover, the metal Bi-decorated Bi₂WO₆ exhibited a photocatalytic efficiency three times higher than that of pure Bi₂WO₆ in the degradation of phenol under visible light, because metal Bi not only acts as an electron acceptor, but also promotes the separation of carriers.319 Finally, the surface plasmon resonance of Bi was also embodied in the Bi/BiOBr composite, which possessed a higher activity for NO oxidation than that of pure BiOBr. 320

Z-scheme heterojunction

Review

Internal Solid Redox electric electron ion pairs field mediator Reduction 1 Reduction 4 Reduction 1 Electron cceptor Electron Oxidation Oxidation Oxidation SC-A SC-A SC-B SC-B SC-B (b) Liquid-phase (c) All-solid-phase (a) Direct

Fig. 29 Schematic illustration of (a) direct Z-scheme, (b) liquid-phase Z-scheme and (c) all-solid-state Z-scheme heterojunctions (SC represents semiconductor).

Z-scheme heterojunction

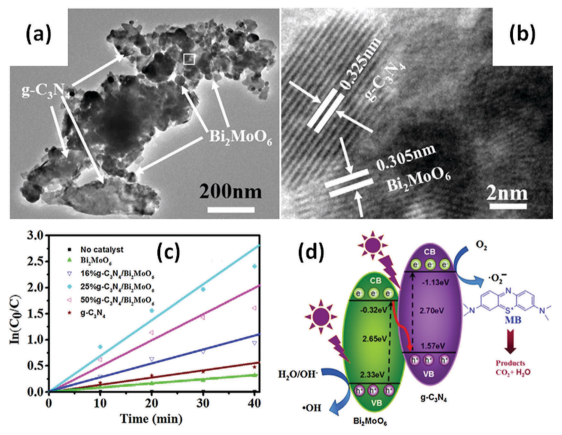


Fig. 30 (a) TEM and (b) HRTEM images of a $g-C_3N_4/Bi_2MoO_6$ composite; (c) pseudo first-order fitted kinetic curves of MB degradation on $g-C_3N_4/Bi_2MoO_6$ composites $(25\%g-C_3N_4/Bi_2MoO_6)$; and (d) a schematic illustration of electron and hole migration in a $g-C_3N_4/Bi_2MoO_6$ direct Z-scheme heterojunction. Reproduced with permission from ref. 297. Copyright 2015 Elsevier.

Carbon materials, such as carbon dots (CDs), CNTs, graphene, graphene oxide (GO), reduced graphene oxide (RGO), carbon quantum dots (CQDs), and graphene quantum dots (GQDs),

can be used as modifiers. Among these materials, graphenebased materials are more favorable because of their special graphitic structure and ideal conductivity.321 Priya et al.322

Z-scheme heterojunction

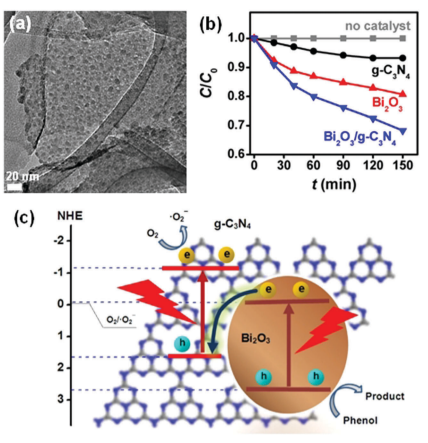


Fig. 31 (a) TEM images of a $Bi_2O_3/g-C_3N_4$ composite; (b) comparison of the photocatalytic performance of Bi_2O_3 , $g-C_3N_4$, and $Bi_2O_3/g-C_3N_4$ in the $degradation \ of \ phenol; \ and \ (c) \ the \ photocatalytic \ mechanism \ of \ a \ Bi_2O_3/g-C_3N_4 \ direct \ Z-scheme \ heterojunction. \ Reproduced \ with \ permission \ from \ phenol; \ and \ (c) \ the \ photocatalytic \ mechanism \ of \ a \ Bi_2O_3/g-C_3N_4 \ direct \ Z-scheme \ heterojunction.$ ref. 300. Copyright 2018 Elsevier.

Table 2 Nanoscale Bi-based direct Z-scheme heterojunction photocatalysts

Bismuthal compound	Coupled material	Method	Application	Ref.
Bi ₂ O ₃	g-C ₃ N ₄	Ball milling and heat treatment	MB	301
$Bi_2Sn_2O_7$	g-C ₃ N ₄	High-temperature solid-state reaction	MB and acid red 18	302
Bi_2MoO_6	g-C ₃ N ₄	Hydrothermal method	MB	297
BiOBr	g-C ₃ N ₄	Reflux process	RhB and BPA	303
BiOIO ₃	g-C ₃ N ₄	Hydrothermal method	MO, RhB, and dichlorophenol	299
BiOI	CdS	Hydrothermal method	RhB	304
Bi ₂ WO ₆	MoS_2	Hydrothermal method	RhB	305
Bi ₂ MoO ₆	CuO, Co ₃ O ₄ , NiO	Precipitation	RhB	306
Bi ₂ O ₃	NaNbO ₃	Ball milling method	RhB	307
BiVO ₄	Se film	Chemical vapor deposition	Photocurrent enhancement	298
BiVO ₄	SiC	Precipitation followed by calcination	O ₂ evolution	308
BiOBr	Bi ₂ MoO ₆	Two-step coprecipitation method	RhB and CIP	309
$BiO_{1-r}Br$	Bi ₂ O ₂ CO ₃	Solvothermal method	CIP	310
BiPO ₄	$Bi_2O_2(OH)(NO_3)$	Hydrothermal method	Dichlorophenol	311
MoS ₂ /BiOI/AgI	2 - 2(-)(3)	Precipitation	RhB	312
BiVO ₄ /ZnIn ₂ S ₄ /g-C ₃ N ₄		Impregnation and calcination	Congo red and metronidazole	313

Rhodamine B, RhB; methylene blue, MB; methyl orange, MO; bisphenol A, BPA; ciprofloxacin, CIP.

loaded a Bi₂O₃/BiOCl heterojunction on a prepared graphene sand composite using a wet impregnation method. The obtained composite showed an improved performance over that of Bi₂O₃/ BiOCl for the mineralization of ampicillin and oxytetracycline. The improvement was attributed to the electron trapping role of the graphitic carbon on the graphene sand composite.

When Bi₂WO₆ microspheres were wrapped with GO using a freeze-drying dehydration method, they exhibited a higher activity than that of pure Bi₂WO₆ on the photocatalytic degradation of RhB due to the improved separation of the electrons and holes by GO.323 GQDs possess a small particle size and high specific surface area. Therefore, GQDs are more favorable for the surface modification of

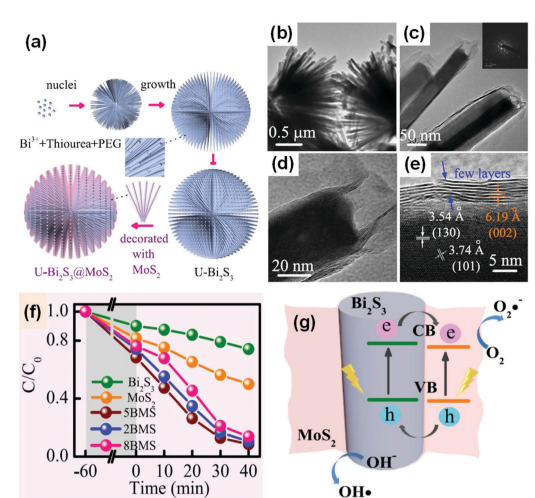


Fig. 32 (a) Schematic illustration of the formation of MoS2-decorated Bi₂S₃ (BMS); (b-d) TEM and (e) HRTRM images of MoS2-decorated Bi₂S₃; (f) comparison of the photocatalytic performance of RhB on MoS_2 , Bi_2S_3 and BMS (2MBS, 5MBS, and 8MBS present in molar ratios of Mo^{4+} -to- Bi^{3+} of 20%, 50%, and 80%) under visible-light irradiation; and (g) a schematic diagram of the charge transfer at a MoS₂/Bi₂S₃ heterojunction. Reproduced with permission from ref. 315. Open Access 2017 Springer Nature.

photocatalysts. For example, Bi₂MoO₆ decorated with GQDs via a self-assembly approach, showed a remarkable increase in photocatalytic degradation efficiency of different target pollutants, including BPA, MB, TC, CIP, and phenol.324 Nevertheless, GO or RGO materials may suffer from instability in long-term reactions. Subramanian and colleagues³²⁵ found that RGO coupled with Bi₂Ti₂O₇ might suffer from decomposition and result in a reduced performance of the composite after long-term illumination in an oxidation environment (Fig. 33a and b). The degradation of RGO was attributed to the oxidation by the formed OH (Fig. 33c). Several conductive polymers, such as polyaniline, polypyrrole, and polythiophene, can play the role of electron scavenger and promote the electron migration in bismuthal photocatalysts. 326-328 However, the conductivities of these polymers are much weaker than those of noble metal and carbon materials. Thus, the related research is less reported. Usually, amorphous carbon is not suitable for electron trapping because of its poor conductivity. Nevertheless, it also plays a positive role in the enhancement of photocatalytic activity due to it enhancing the visible light absorption and carrier separation of the photocatalyst.329

Introducing vacancies or elements with different chemical valence states may also improve the performance of photocatalysts. Wang et al. 330 introduced vacancies into the surface of Bi₆S₂O₁₅ nanowires by increasing the molar ratio of Na₂SO₄ and $Bi(NO_3)_3$ from 1:3 to 3:1 during hydrothermal synthesis. The light absorption of vacancy Bi₆S₂O₁₅ nanowires at 370-500 nm was stronger than that of pure Bi₆S₂O₁₅ nanowires. Vacancy Bi₆S₂O₁₅ nanowires also showed improved photocatalytic activity for the degradation of phenol and MB. The enhancement was attributed to the improved visible-light response induced by the surface vacancies (Fig. 34). Zhang and colleagues³³¹ treated BiVO₄ at different temperatures under different atmospheres. The sample treated at 973 K in N2 exhibited the best activity for the formation of EG from aldehyde. The enhancement was attributed to the formation of V⁴⁺. Bi₂WO₆ with a large fraction of {100} high-energy facets showed relatively high efficiency in the photocatalytic degradation of diclofenac, because "Bi-O" dimer vacancy pairs formed on the {100} facets could bring about a narrower band gap and less photoexcited charge recombination.332 Moderate oxygen-deficient defects in BiOBr

 Table 3
 Surface modification strategies for nanoscale Bi-based photocatalysts

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g-BNBiPO $_4$ Solvothermal methodEnrofloxacinBi $_2O_3$ QDsBiVO $_4$ CalcinationRhBBi $_2O_4$ BiOBrAlkali posttreatment under light irradiationMOBiOIBi $_2$ MoO $_6$ Deposition-precipitationMOBiOBrBiPO $_4$ Hydrothermal methodMB	341 248 276 342
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BiOBr BiPO $_4$ Hydrothermal method MB	
Carbon $Bi_4Ti_3O_{12}$ Coprecipitation MO	344
Carbon Bi ₂ MoO ₆ Two-step hydrothermal method RhB	345
Carbon microspheres BiFeO ₃ Hydrothermal method RhB	346
Carbon microspheres Bi _{0.5} Dy _{0.5} VO ₄ Self-assembly H ₂ generation	347
Carbon nanotubes $Bi_4O_5Br_2$ Solvothermal method Tetracycline hydrocl	
Carbon dots BiOI Immersion MO	348
CQDs Bi ₂₀ TiO ₃₂ Oil bath Isoproturon	349
CQDs $Bi_4O_5I_2$ Solvothermal method RhB	350
	351
CQDs BiOI Hydrothermal method RhB GO BiVO ₄ Hydrothermal method RhB	352
GO BiOCl Two-step synthesis RhB	353
	354
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3	355
GO Bi ₂ O ₂ CO ₃ Precipitation Gaseous NO	356
GO Bi ₂ WO ₆ Liquid nitrogen freezing and freeze-drying dehydration RhB	323
GO BiOI Self-assembly Phenol	357
RGO BiOI Hydrothermal method MO	358
RGO BiOBr Two-step hydrothermal method Higher activity for M	
RGO Bi ₂₅ FeO ₄₀ Hydrothermal method MO	360
RGO Bi ₂ WO ₆ Wrapped RhB	361
RGO Bi ₂ S ₃ Hydrothermal method 2,4-Dichlorophenol	362
RGO BiPO ₄ Solvothermal method Chlorpyrifos detecti	
Graphene Ag/BiOBr _{0.2} I _{0.8} Combined solvothermal method and photodeposition RhB	364
Graphene BiOCl _{0.7} Br _{0.3} Solvothermal method RhB	365
Graphene Bi ₂ O ₂ CO ₃ Hydrothermal method RhB	366
Ag and graphene Bi ₂ WO ₆ Hydrothermal method followed by photodeposition RhB	367
GQDs Bi ₂ MoO ₆ Self-assembly BPA, MB, TC, CIP, a	•
Au and RGO Bi ₂ MoO ₆ Solvothermal synthesis and photochemical reduction RhB	368
Ag and graphene $\mathrm{Bi}_2\mathrm{Fe}_4\mathrm{O}_9$ Multistep synthesis MB	369
Metal and ion Pd Bi ₂ MO ₆ Photoreduction Phenol	370
2 0	371
	372
T 1	2
Pt BiOBr Photodeposition p-Nitrophenol	374
Pt BiFeO ₃ Impregnation and thermal reduction MO	375
Pt Bi ₂ WO ₆ Chemical reduction Rhodamine 6G	376
Pt Bi ₂ WO ₆ /TiO ₂ Photodeposition CO ₂ reduction	377
Au BiOCl Photodeposition Formaldehyde	378
Au BiVO ₄ Pulsed laser deposition Congo red and water	
Au BiPO ₄ Solvothermal method CO oxidation	380
Au Bi ₂ O ₂ CO ₃ Hydrothermal method NO	381
Ag $BiFeO_3$ film Sol spin coating and annealing O_2 evolution	382
Ag BiOBr Solvothermal method Tetracycline hydrocl	
Ag BiVO ₄ Hydrothermal method Reduced electron-tra	
Ag Bi ₄ Ti ₃ O ₁₂ Sonochemical method RhB	385
Ag Bi ₂ WO ₆ Irradiation RhB	386
Bi Bi ₂ WO ₆ In situ reduction Phenol	319
Bi Bi ₂ WO ₆ nanorod Hydrothermal reaction Rh6G	387
Bi Bi ₂ MoO ₆ Solvothermal reduction Rh6G	388
Bi BiPO ₄ Solvothermal treatment MB	389
Bi BiOBr Reduction reaction at room temperature MO	390
Bi BiOI/CNFs Solvothermal method MO	391
Cr $Bi_4Ti_3O_{12}$ Sol-gel hydrothermal process H_2 generation	69

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Table 3 (continued)

Modifier	Bi-Based material	Method	Target material of enhanced perfo	rmance Ref.
Ni	BiOI	Photo-assisted deposition	Reducing Cr(vI) to Cr(III)	392
Fe(III)	$Bi_2O_2CO_3$	Impregnation	MO	393
Fe	$Bi_2Ti_2O_7$	Precipitation	MO	394
Fe_2O_3	Macroporous BiVO ₄	Impregnation	4-Nitrophenol	395
CoO_x	BiFeO ₃	Photodeposition	O ₂ evolution	396
Rh(III)	BiOCl	Impregnation	Gaseous acetaldehyde (Vis)	397
Organic material				
Polythiophene	Bi ₂ MoO ₆	<i>In situ</i> chemical oxidative polymerization	RhB	326
Polyaniline	Bi ₂ O ₂ CO ₃	Low-temperature chemical method	RhB	327
PVP	BiOBr	Solvothermal method	RhB	328
MIL-101 (MOF)	$BiVO_4$	Hydrothermal method	RhB	398
Ionic liquid	${ m BiPO}_4$	Hydrothermal method	RhB	399
Surface defect				
Bi defects	$Bi_6S_2O_{15}$	Hydrothermal method	Phenol	330
Dimer vacancy	Bi ₂ WO ₆	Solvothermal method	Diclofenac	332
Defect	BiPO ₄	Ball milling	MB	400

GO: graphene oxide; RGO: reduced graphene oxide; MOF: metal-organic framework; CQDs: carbon quantum dots; GQDs: graphene quantum dots; Rh6G: Rhodamine 6G; MB: methylene blue; PEDOT:PSS: poly(3,4-ethylenedioxythiophene):poly(styrenesulfonate).

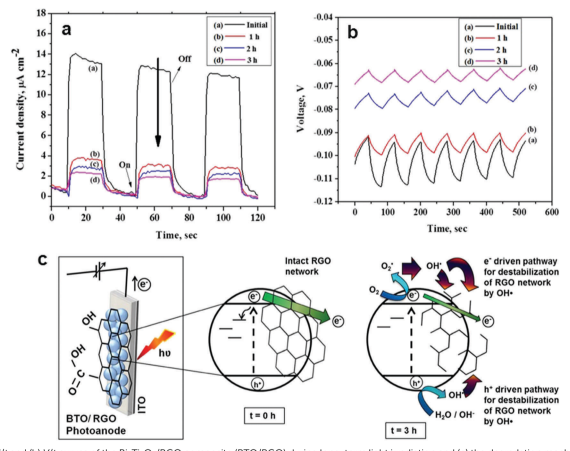


Fig. 33 (a) //t and (b) V/t curves of the Bi₂Ti₂O₇/RGO composite (BTO/RGO) during long-term light irradiation and (c) the degradation mechanism of the RGO on a BTO/RGO film under light illumination for 3 h in an air-equilibrated solution. Reproduced with permission from ref. 325. Copyright 2016 Electrochemical Society.

also play an indispensable role for superior photocatalytic CO₂ reduction, by acting as the active sites for CO2 adsorption and activation, trapping photogenerated electrons and thus impact upon the recombination of the electron-hole pairs. 333 Oxygen

vacancies were also found to have a positive influence on the performance of BiOCl on photocatalytic CO2 reduction because oxygen vacancies can induce exciton dissociation and provide more photo-induced electrons for CO₂ reduction.³³⁴

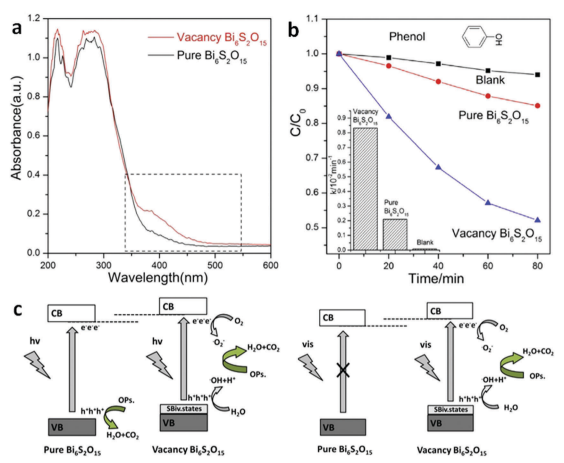


Fig. 34 (a) UV-visible diffuse reflectance spectra of pure $Bi_6S_2O_{15}$ and vacancy $Bi_6S_2O_{15}$ nanowires; (b) degradation rate constant k of pure $Bi_6S_2O_{15}$ and vacancy Bi₆S₂O₁₅ nanowire photocatalysts; (c) schematic diagram of the carrier separation and photocatalytic reaction of pure Bi₆S₂O₁₅ and vacancy Bi₆S₂O₁₅ nanowires under UV-visible irradiation. Reproduced with permission from ref. 330. Copyright 2015 Elsevier.

Conclusions and prospects

A layered structure and excellent visible-light response endow Bi-based photocatalysts with excellent photocatalytic activity and promising prospects for application in the fields of environmental science and energy. Studies centered on Bi-based photocatalysts will be beneficial for the future development of photocatalysts. Considerable work has been performed to control the composition, morphology, and structure of these catalysts to improve the photocatalytic performance of Bi-based photocatalysts. These photocatalysts are used for the visible-light photocatalytic degradation of organic pollutants, H2 generation, water splitting, and CO₂ reduction. Although significant progress has been achieved, great efforts are still required to further explore the advantages of Bi-based photocatalysts as visible-light photocatalysts. The following five aspects deserve special attention:

(1) Previous studies on Bi-based photocatalysts have mainly focused on the photocatalytic degradation of organic pollutants. Few studies have reported on the application of these photocatalysts in photocatalytic H₂ generation, water splitting, and CO₂ reduction. Several investigations have focused on BiVO₄. The main obstacle in the study of Bi-based photocatalysts is the less negative CB and the poor reducing abilities of bismuthal compounds. More

bismuthal materials should be extensively explored to overcome these obstacles. The construction of nanosized direct Z-scheme heterojunctions by coupling bismuthal semiconductors with semiconductors that have a more negative CB is a promising method for exploring Bi-based photocatalysts for reduction applications.

- (2) Bi-based photocatalyst photodegradation has been primarily evaluated in aqueous solutions instead of under gaseous conditions. The latter condition is favorable for the use of Bi-based photocatalysts because visible light can be more fully utilized in the gas phase. The removal of formaldehyde and benzene from air is important for environmental remediation and indoor air purification. Therefore, extending the application of Bi-based photocatalysts to the degradation of gaseous contaminants is meaningful.
- (3) The aim of photocatalysis is to solve serious environmental and energy problems in the world. We hope that this review can further stimulate the research into and application of Bi-based photocatalytic materials in the near future. We believe that the understanding and knowledge on Bi-based photocatalysts can be greatly enhanced and the experimental and characterization methods improved.
- (4) Theoretical simulations can provide new insight into understanding the photocatalytic mechanism and relationship

between the microstructure and performance. The theoretical investigations on Bi-based photocatalytic materials should be further strengthened, which will contribute to a deep understanding on the surfaces, interfaces, nanostructures and activity

at the atomic and molecular level of the material properties.

(5) Although Bi-based photocatalysts have been continuously investigated over the past ten years, their photocatalytic performance is still far from the requirements for practical application. More research work is still needed to reach their real application.

Conflicts of interest

Review

There are no conflicts to declare.

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