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Introduction

A minireview on catalysts for photocatalytic N_2 fixation to synthesize ammonia

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Ammonia (NH₃) is an important feedstock in chemical industry. Nowadays NH₃ is mainly produced via the industrialized Haber-Bosch process, which requires substantial energy input, since it operates at high temperatures (400-650 °C) and high pressures (20-40 Mpa). From the energy conservation point of view, it is of great significance to explore an alternative avenue to synthesize NH₃, which is in line with the concept of sustainable development. Very recently, photocatalytic N₂ fixation (PNF) has been discovered as a safe and green approach to synthesize NH₃, as it utilizes the inexhaustible solar energy and the abundant N₂ in nature to synthesize NH₃ under mild conditions. A highly efficient catalyst is the core of PNF. Up to now, extensive studies have been conducted to design efficient catalysts for PNF. Summarizing the catalysts reported for PNF and unraveling their reaction mechanisms could provide guidance for the design of better catalysts. In this review, we will illustrate the development of catalysts for PNF, including semiconductors, plasmonic metal-based catalysts, iron-based catalysts, rutheniumbased catalysts and several other catalysts, point out the remaining challenges and outline the future opportunities, with the aim to contribute to the development of PNF.

Ammonia (NH₃) is one of the basic raw materials in industrial chemistry and has been widely applied in various fields.¹⁻⁴

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Industrially, NH₃ could be used to produce fertilizers, synthetic fibres, nitrile rubber and so on; medically, NH₃ is often utilized as a drug to treat dizziness and fainting; militarily, NH₃ could serve as a biological disinfectant to disinfect sarin agents. The wide applications of NH₃ make it essential in promoting the development of national economies.

At present, the Haber-Bosch process is the main approach to synthesize NH₃.⁵⁻⁸ However, the Haber-Bosch process is carried out under harsh reaction conditions (the pressure is high up to 20–40 MPa and the temperature is in the range of 400–650 $^{\circ}$ C), which consumes extensive energy.9,10 From the context of global



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Fig. 1 Schematic illustration of catalysts that have been designed for PNF.

energy crisis, it is necessary to search for an alternative avenue to synthesize NH_3 which is in line with the concept of sustainable development.

Photocatalytic N_2 fixation (PNF) is a process which utilizes the inexhaustible solar energy and the abundant N_2 in nature to synthesize NH_3 under mild conditions.^{11–18} It is a safe and green approach and has great potential to solve the problems encountered in the industrialized Haber-Bosch process, provided that efficient photocatalysts are adopted.^{19–31}

In photocatalytic N_2 fixation reaction, the cleavage of the $N \equiv N$ bond is the rate determining step. That is, effective catalysts for PNF should be able to accelerate the $N \equiv N$ bond cleavage process.

Over the past few years, an enormous amount of research effort has been devoted to explore effective catalysts for PNF. A large number of photocatalysts, such as semiconductors, plasmonic metal-based catalysts, iron-based catalysts, rutheniumbased catalysts and several other catalysts, have been designed for PNF (a schematic illustration is shown in Fig. 1).

For each type of these photocatalysts, a specific photocatalyst for PNF could be simply divided into two different functional units. One is the light harvesting unit (which is used to denote the active sites that could adsorb light) and the other is the



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thermal-driven active unit (which is referred as the active center that directly involves in the polarization and activation of N_2). The tailoring of each of the type of functional unit could contribute to the performance of a catalyst in PNF. In this review, we will summarize the photocatalysts that have been reported active for PNF, generalize the principles for the design of efficient catalysts (tailoring the light harvesting unit or the thermal-driven active unit), unravel their reaction mechanisms, point out the remaining challenges and prospect the future development, with the aim to provide guidance for the rational design of more efficient photocatalysts and contribute to the development of PNF.

Photocatalysts for PNF

In this section, the photocatalysts for PNF are classified into five categories, including semiconductors, plasmonic metal-based catalysts, iron-based catalysts, ruthenium-based catalysts and other catalysts. The progress of each category of photocatalysts is summarized and discussed in the following sub-sections.

Semiconductors as photocatalysts for PNF

When semiconductors are adopted to catalyse PNF, suitable energy band levels are required. Generally speaking, semiconductors for PNF are designed according to the following principles, ① the semiconductor could absorb light efficiently and it could be excited by light easily, ② the separation and transfer efficiency of the photogenerated electron hole pairs should be effective, and ③ the energy levels of the semiconductor photocatalyst could meet the standards for the two photo-induced half reactions, referring to the reduction of N₂ and the oxidation of H₂ or H₂O.

Some pristine semiconductors could meet the standards to drive PNF, however, their efficiencies are quite low. Introducing vacancies or foreign elements into the semiconductor, functionalizing the pristine semiconductor, constructing heterojunctions/homojunctions or design a semiconductorbased hydrophilic-hydrophobic catalyst, are approaches to extend the light harvesting spectrum, enhance the light harvesting capacity (which means its capacity to absorb more light), facilitate the separation and transfer of photoinduced electron hole pairs and further accelerate PNF.

Pristine semiconductors. Pristine semiconductors are a type of photocatalyst with only light harvesting unit for PNF. Pristine semiconductors are widely studied in PNF. The catalytic performance of pristine semiconductors could be improved by regulating their optical properties. Theoretical calculations suggest that (110) facet of rutile TiO₂ is capable of activating N₂ into NH₃ *via* both the associative and dissociative mechanism;³² (040) facet of BiVO₄ single crystal could catalyze PNF, with V⁴⁺/V⁵⁺ as the active site, where V⁴⁺ chemisorbs N₂, V⁵⁺ serves as the electron transfer bridge and the photogenerated electrons are the driving force for PNF.³³

The activities of pristine semiconductors have been verified experimentally. For example, ultrathin MoS_2 could convert N_2 into NH_3 *via* a six-electron reduction process (Fig. 2), achieving a NH₃ synthesis rate of 325 µmol g⁻¹ h⁻¹ without any sacrificial reagents or co-catalysts.³⁴ The electron-rich property of ultrathin MoS₂ as well as the high concentration of localized electrons upon light irradiation accounted for its activity.³⁴ Bismuth monoxide (BiO) quantum dots, synthesized *via* a facile hydrothermal method, are reported as a highly efficient catalyst for PNF, recording a NH₃ generation rate of 1226 µmol g⁻¹ h⁻¹ without the assistance of any sacrificial reagents or co-catalysts.³⁵ It is postulated that all the low valence Bi(II) in BiO were potential active sites for N₂ activation.³⁵

The catalytic activity of pristine semiconductor could be improved, in case that a 2-dimensional (2D) material is used as co-catalyst. For instance, P25 itself yielded NH₃ at a rate of 2.11 μ mol g⁻¹ h⁻¹ under full spectrum light irradiation, while the activity was fivefold increased when Ti₃C₂ MXene was used as a co-catalyst.³⁶ Similarly, the catalytic performance of CdS in PNF boosted obviously when black phosphorous nanosheet was adopted as a co-catalyst.³⁷ It is reported that 2D co-catalysts facilitated the separation of electron–hole pairs and promoted N₂ chemisorption and activation.

Despite that some pristine semiconductors are active for PNF, their efficiencies are generally low. To further improve their efficiencies, multiple approaches are adopted to modify the pristine semiconductors.

Semiconductors with vacancies. Introducing vacancies into a semiconductor endows the pristine semiconductor additional properties. On one hand, vacancy introduction could tailor the light harvesting unit of the photocatalyst, such as narrow its band gap, extend its light harvesting range and enhance its light harvesting capacity. On the other hand, the vacancies occasionally serve as thermal-driven active sites for N_2 activation. Defective semiconductors have been widely applied in PNF. The functions of the defective sites vary with the catalytic systems.

(1) The introduction of defective sites engineers the light harvesting unit, such as lowers the conduction band position, engineers the band gap and improves the light harvesting capacity of the semiconductor. Few-layer $g-C_3N_4$ (ref. 38) and one-dimensional $g-C_3N_4$ (ref. 39) are rich in N defects. The existence of N defective sites lowered the conduction band position and increased its light harvesting capacity, which then contribute to their performance in PNF.^{38,39} Band gaps and the light harvesting capacities could be consecutively tuned by dedicatedly controlling the content of surface vacancies.⁴⁰

(2) The defective sites serve as the active sites to enhance the adsorption and activation of N₂. N vacancies on nitrides generally activate N₂ *via* a pathway analogous to Mar-van Krevelen mechanism. That is, N₂ is firstly activated at the N vacancies and then transformed into NH₃ by reacting with H₂ or H₂O. Wang *et al.*'s work is a typical example. In their work, a N deficient g-C₃N₄ catalyst was prepared by the dielectric barrier discharge plasma treatment method, which displayed a NH₃ production rate of 161.8 μ mol g⁻¹ h⁻¹.⁴¹ Mechanism exploration disclosed that N₂ was activated *via* the two-path analogous Mar-van Krevelen mechanism.⁴¹

Oxygen vacancies on oxide semiconductors are active centers for N₂ adsorption and capable of activating N₂.⁴²⁻⁴⁴ Theoretical study revealed that the oxygen vacancies on (001) and (100) facets of MoO_{3-x} nanobelts could chemisorb N₂ via side-on and end-on models, respectively, which accelerates the sluggish rate determining step (N₂ activation) in PNF and boosted its performance.⁴³

Cation defective sites on a semiconductor are electron-rich, which could effectively activate the N \equiv N bond and accelerate catalytic activity in PNF. For instance, Zn deficient Zn₃In₂S₆ exhibited a NH₃ generation rate of 261.2 µmol g⁻¹ h⁻¹ under visible light irradiation, 15 times higher than the one with poor defects.⁴⁵

(3) On some defective semiconductors, the defective sites not only improve light harvesting capacity but also facilitate N₂ activation. For example, Bi₂WO₆ hollow microspheres prepared by a solvothermal template-free method are rich in oxygen vacancies.⁴⁶ The oxygen vacancies induce a sub-band (defect energy level), which not only narrows its band gap and extends its light absorption region (up to 700 nm), but also localizes metastable electrons. These metastable electrons jump to the anti-bonding orbitals of N₂ *via* a manner of non-radiative transfer and activate N₂. Overall, the Bi₂WO₆ hollow microspheres demonstrated a NH₃ yield of ~53 µmol g⁻¹ h⁻¹ under simulated sunlight.⁴⁶

Similarly, in addition to enhance the light harvesting capacity, the oxygen vacancies on BiOCl,⁴⁷ TiO₂ ⁴⁸⁻⁵⁰ and BiOBr⁵¹ also serve as the active sites for N₂ activation and reduction while the oxygen vacancies on Bi₃FeMo₂O₁₂ help adsorb and stabilize the N-H intermediate during N₂ activation,⁵² which cooperatively boost NH₃ production *via* PNF (Fig. 3).⁵³

Intrinsic strain is occasionally induced by vacancies. Thereby the strain and vacancies cooperatively contribute to PNF. Zhang *et al.* synthesized an ultrathin TiO_2 nanosheet catalyst with



Fig. 2 Schematic illustration of trion induced six-electron N_2 reduction. This figure has been adapted from ref. 34 with permission from Elsevier, copyright 2017.



Fig. 3 (a) Strain induced by oxygen vacancies and Jahn-Teller effects in TiO₂ nanosheet. (b) NH₃ yield over different catalysts under ultraviolet (UV)-vis light irradiation for 1 h. This figure has been adapted from ref. 53 with permission from Wiley-VCH, copyright 2019.

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abundant oxygen vacancies *via* a facile copper-doping method, which could absorb visible light high up to 700 nm.⁵³ It was discovered that there was intrinsic compression strain in the asprepared TiO₂ nanosheet. The oxygen vacancies and the strain worked in concert to chemisorb and activate N₂ and H₂O effectively, leading to a high NH₃ production rate (78.9 μ mol g⁻¹ h⁻¹).⁵³

Doped semiconductors. Doping one or more foreign elements into a semiconductor is an important approach to mediate the light harvesting unit (such as introduce new energy levels, engineer its energy structure, tailor its light harvesting capacity) and/or manipulate the active sites for N_2 activation. Doped semiconductor is a category of catalysts for PNF. Here, based on the functions of the dopant, the doped semiconductor catalyst for PNF is review.

(1) Suitable dopant could serve as active sites for promoting N₂ activation. For example, $Mo_{1-x}W_xS_2$ nanosheets, which could be considered as Mo doped WS₂, recorded a NH₃ production rate of 111 µmol g⁻¹ h⁻¹ under visible light irradiation, when the concentration of 1T phase was 33.6% and Mo/W = 0.68 : 0.32.⁵⁴ The doping of Mo into WS₂ resulted in a higher electron density on W 5d orbitals, which polarized the adsorbed N₂ and responded for its PNF activity.⁵⁴

(2) Under most of cases, the introduction of dopant not only engineers the optical properties but also manipulates the active sites for N_2 activation.

Doping B into g-C₃N₄ led to the formation of a new chemical bond B–N–C, which not only effectively enhanced the light harvesting capacity and suppressed the recombination of photoinduced electron–hole pairs, but also served as an active center for N₂ chemisorption and activation.⁵⁵ In the issue, B-g-C₃N₄ gave a NH₃ yield of 313.9 µmol g⁻¹ h⁻¹ under visible light assistance.⁵⁵ Mn²⁺ could be doped into W₁₈O₄₉ *via* partially replacing the W sites.⁵⁶ The doped Mn²⁺ played multiple roles in PNF. ① Mn²⁺ acted as the active sites for the chemisorption of N₂ and H₂O, ② Mn²⁺ weakened the N≡N bond through proton coupling process and ③ the doped Mn²⁺ facilitated the separation and migration of photoinduced electron–hole pairs. Based on these advantages, the as prepared Mn–W₁₈O₄₉ catalyst exhibited a NH₃ production rate of 97.9 µmol g⁻¹ h⁻¹ under full spectrum irradiation of a 300 W Xe lamp.⁵⁶

When a foreign element is doped into a semiconductor with vacancies sites, the dopants and vacancies sites might synergistically activate N₂ and contribute to PNF. For example, in sulfur doped oxygen deficient TiO_2 ($\text{TiO}_{2-x}S_y$), the oxygen vacancies and sulfur dopant worked in concert to facilitate N₂ adsorption and extend its light absorbing capacity to nearinfrared region.⁵⁷ As a result, $\text{TiO}_{2-x}S_y$ yielded NH₃ at a rate of 114.1 µmol g⁻¹ h⁻¹ under full spectrum light irradiation.⁵⁷ Similar phenomena were also observed over Br doped BiOCl with oxygen vacancies,⁵⁸ Ni doped vacancy-rich TiO₂,⁵⁹ as well as S doped g-C₃N₄ with carbon vacancies.⁶⁰

Doping a foreign element into a semiconductor sometimes leads to the generation of vacancies. Tang *et al.* fabricated a carbon doped TiO₂ nanosheet catalyst (C-TiOx) from Ti₃SiC₂ *via* a bottom-up approach (Fig. 4a).²⁹ It was discovered that the doping of carbon led to the generation of oxygen vacancies in TiO₂. As charge compensation, controllable Ti³⁺ sites were produced. The oxygen vacancies broadened its light harvesting region and the electron-rich Ti^{3^+} were active for N₂ activation. With Ru/RuO₂ as co-catalyst to promote the separation and migration of photoinduced electron–hole pairs, the optimal C-TiO_x recorded a 109.3 µmol g⁻¹ h⁻¹ NH₃ synthesis rate under visible light irradiation and an apparent quantum yield of 1.1% at 400 nm (Fig. 4b).²⁹ Similarly, oxygen vacancies could also be generated by doping carbon into BiOI.⁶¹ Carbon dopant decreased the band gap, extended the light harvesting region, facilitated the separation and migration of electron–hole pairs and consequently hastened PNF, leading to a NH₃ generation rate of 311 µmol g⁻¹ h⁻¹ under the illumination of simulated sunlight.⁶¹

As for doping, it is clear that most of the doped semiconductors exhibited a NH₃ production rate of ~100 μ mol g⁻¹ h⁻¹, with only two exceptions on B-g-C₃N₄ and C– BiOI, who gave NH₃ yields over 300 μ mol g⁻¹ h⁻¹. Even though limited by the number of studies, it might provide a clue that nonmetal element doping could endow the doped semiconductors better performance in PNF.

Functionalized semiconductors. Grafting functional groups onto a semiconductor or modifying a pristine semiconductor is an important approach to mediate the two functional units and consequently enhance their performance for PNF.

(1) Functionalized semiconductors with engineered light harvesting unit.

Acid treated semiconductors. Tian *et al.* reported that salicylic acid (SA) modification could enlarge the Brunauer–Emmett–Teller surface area, improve the optical absorption capacity as well as promote the separation of photoinduced electron–hole pairs in $g-C_3N_4$, which resulted in a much enhanced PNF activity over $g-C_3N_4$ -SA.⁶²

Base treated semiconductors. It has been reported that base treated semiconductors exhibited improved electron-hole separation efficiency. Typical examples are given by Yangjeh *et al.*⁶³ and Wang *et al.*,⁶⁴ who reported that MgO decorated g- C_3N_4 and KOH treated g- C_3N_4 gave superior performance in PNF than pristine g- C_3N_4 , and the reduced recombination rate of electron-hole pairs was one of the reasons for their premier activities.

Metal decorated semiconductors. Cu, Fe, Ni and Pd decoration could promote the charge transfer efficiency. Cu, Fe or Ni decorated TiO₂, synthesized *via* microwave assisted hydrothermal method, not only exhibited promoted charge transfer efficiency, but also had larger specific surface area and stronger capacity in harvesting visible light.⁶⁵ As a result, they exhibited 1.9–6.0 times higher PNF activities than the pristine TiO₂ under simulated light irradiation.⁶⁵ Pd decorated TiO₂, prepared by one-pot microwave synthesis techniques, also demonstrated reduced recombination of photoinduced electron–hole pairs and led to a 4 fold higher NH₃ production rate than the unmodified one.⁶⁶

Quantum dots modified semiconductors. Graphene quantum dots modified Bi_2WO_6 , with graphene quantum dots uniformly dispersed on the surface of Bi_2WO_6 , exhibited remarkably enhanced PNF activity than the two single component counterparts. Characterization results implied that the recombination of photoinduced electron-hole pairs was



Fig. 4 (a) Schematic illustration for the synthesis of C-TiO_x. (b) Photocatalytic performance of C_n -TiO_x (n denotes the treatment time of Ti₃SiC₂) and Ti₃SiC₂ in PNF under visible light irradiation. Ru/RuO₂ was used as co-catalyst. This figure has been adapted from ref. 29 with permission from Wiley-VCH, copyright 2021.

significantly reduced and the junction between graphene quantum dots and ${\rm Bi}_2{\rm WO}_6$ helped to boost the photocatalytic activity.⁶⁷

(2) Functionalized semiconductors with tailored active sites. **Hydrogenated semiconductors.** Unsaturated Mo atoms in Bi₂MoO₆ are the active sites for N₂ chemisorption, activation and reduction, on the contrary, the Mo atoms within the crystal are always inert. Focusing on this standpoint, Zhang *et al.* hydrogenated Bi₂MoO₆ with the aim to expose more unsaturated Mo atoms by inducing the escape of oxygen atoms in saturated Mo–O bond and enhance its N₂ activation capacity. As expected, the hydrogenated Bi₂MoO₆ exhibited a NH₃ production rate 9.5 times higher than that of untreated one, up to 1.3 mmol g⁻¹ h⁻¹.⁶⁸

(3) Functionalized semiconductors with two functional units modified.

Semiconductors with functional groups. Grafting amine groups onto g- C_3N_4 could enhance the charge separation efficiency and consequently nearly double its activity in PNF.⁶⁹ Introducing cyano groups onto g- C_3N_4 not only improves the separation and migration of photoinduced electron-hole pairs but also enhances N_2 activation owing to its electronwithdrawing characteristics. Thus, cyano group functionalized g- C_3N_4 resulted in a 12.8 times promoted activity in PNF than pristine g- C_3N_4 .⁷⁰ With the assistance of K, the cyano group on modified g- C_3N_4 could be regenerated readily *via* the analogous Mars van Krevelen mechanism, which ensured the stability of the functionalized semiconductors.¹²

Hydrogenated semiconductors. Dong *et al.* reported that hydrogen treatment could withdraw the oxygen atoms and leave oxygen vacancies in BiOBr, which then broadened the photoelectricity absorption window, triggered the electron transfer from BiOBr to the adsorbed N_2 , and responded for the 2.6 times improved PNF efficiency, compared to the untreated counterpart.⁷¹

Acid treated semiconductors. Wang *et al.* reported that phosphate acid treated LaFeO₃ could catalyze PNF effectively, producing NH₃ at a rate of \sim 250 µmol g⁻¹ h⁻¹ under simulated

light irradiation. Mechanism exploration suggested that phosphate acid served as the Lewis acid center, it worked synergistically with Fe in LaFeO₃ to activate N₂ *via* the "push-pull" hypothesis. That is, the electron density is pulled from Fe and pushed into N₂ by the adjacent hydrogen bonding sites. In addition, phosphate modification facilitated H₂O dissociation.⁷²

Metal decorated semiconductors. In case that a metal is decorated onto a semiconductor with vacancy sites, the doped metal and the vacancies work in concert to promote N₂ activation. For instance, Dong *et al.* constructed a Bi/BiOBr heterostructure with abundant oxygen vacancies *via* a one-step solvothermal strategy. In BiOBr, the oxygen defective sites were the active centers for N₂ adsorption. Bi and oxygen vacancies promoted the interfacial charge transfer from Bi/ BiOBr to the adsorbed N₂, facilitated charge separation efficiency and accounted for a remarkably high NH₃ production rate (1350 µmol g⁻¹ h⁻¹).⁷³

Heterojunctions and homojunctions. Heterojunctions and homojunctions are fabricated by integrating two or more semiconductors. Heterojunctions and homojunctions generally inherit the merits of each single semiconductor counterpart, exhibit stronger light harvesting capacities and deliver high photocatalytic activities. Further introducing other dopants or vacancies into heterojunctions or homojunctions might tailor the active centers for N_2 activation. The advantages of heterojunctions and homojunctions make them applicable in PNF.

(1) Heterojunctions/homojunctions with modified light harvesting unit. Bi₂Te₃/BiOCl,⁷⁴ g-C₃N₄/ZrO₂,⁷⁵ MoO₂/BiOCl⁷⁶ perovskite/attapulgite⁷⁷ and CdS/LDH (LDH: layered double hydroxide)⁷⁸ are typical heterojunctions that have been reported active in PNF. Here CdS/LDH is taken as an example for elaboration. In the case that CdS/LDH heterojunction was constructed between (003) or (012) facet of LDH and (002) facet of CdS, a build-in electric field would be induced under light irradiation, which promoted charge transfer from the heterostructure to the reaction media for N₂ activation *via* a favorable configuration and resulted in a better catalytic activity.⁷⁸

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Z scheme is a special case of heterojunctions. Z schemes AgBr/ Bi₄O₅Br₂,⁷⁹ Bi₂O₃@CoAl-LDHs,⁸⁰ 3,4-dihydroxybenzaldehydefunctionalized Ga₂O₃/g-C₃N₄,⁸¹ g-C₃N₄/Mg_{1.1}Al_{0.3}Fe_{0.2}O_{1.7},⁸² nano-MOF@defected g-C₃N₄ (MOF: metal organic frameworks)⁸³ and SiW₉Co₃/PDA/BWO (PDA: poly-dopamine; BWO: Bi₂WO₆)⁸⁴ have been reported active in PNF. In these Z schemes, the separation of photoinduced electron-hole pairs was significantly improved, which played a dispensable role in boosting their catalytic activities.

(2) Heterojunctions/homojunctions with modified light harvesting unit and active sites.

The catalytic activities of heterojunctions in PNF could be further improved by doping foreign elements, introducing defective sites or loading another component to one of the semiconductors of heterojunctions. N deficient g- $C_3N_4/Cu_2(-OH)_2CO_3$,⁸⁵ N deficient g- C_3N_4/Ag_2CO_3 ,⁸⁶ Bi₂MoO₆/oxygenvacancy-rich BiOBr,⁸⁷ MoS₂/C–ZnO (C was loaded onto ZnO),⁸⁸ TiO₂@C g- C_3N_4 ,⁸⁹ B doped g- C_3N_4/Ni_2P^{90} and In₂O₃/In₂S₃ (oxygen vacancies are generated *in situ*)⁹¹ are representatives. In this type of heterojunctions, the junctions between the two semiconductors as well as the doped/loaded component or defective sites synergistically contributed to their activities in PNF.

Homojunction catalyst such as ordered/disordered TiO_2 exhibited a superior activity in PNF, affording a NH₃ formation rate of 432 µmol g⁻¹ h⁻¹ under solar illumination.⁹² In the homojunction catalyst, ordered TiO₂ exhibited a stronger N₂ adsorption capacity with a reduced activation barrier while the disordered TiO₂ was rich in oxygen vacancies which selectively chemisorbed N₂ and enhanced visible light harvesting. The synergistic effect between ordered TiO₂ and disordered TiO₂, together with the rapid interfacial charge separation, ensured its superior activity (Fig. 5).⁹²

Semiconductor-based hydrophilic-hydrophobic catalyst. In case that H_2O is the proton donor and solid catalyst is designed, PNF occurs at the gas (N_2) -liquid (H_2O) -solid (catalyst) triphase

interface. The poor solubility and the poor diffusion rate of N₂ in H₂O severely limit the N₂ fixation efficiency. In order to overcome this obstacle, Fan *et al.* designed a Bi₄O₅Br₂/ZIF-8 catalyst (ZIF-8 is a kind of MOFs), where Bi₄O₅Br₂ is hydrophilic whereas ZIF-8 is hydrophobic.⁹³ Such a hydrophilic–hydrophobic catalyst allowed the direct delivery of N₂ and H₂O into the reaction interface, without the diffusion of N₂ in H₂O. The rapid supply of N₂ ensured the efficient utilization of photoinduced electrons and led to a superior activity (NH₃ production rate was high up to 327 µmol g⁻¹ h⁻¹).⁹³

Plasmonic metal-based catalysts for PNF

Plasmonic metals, such as Au and Ag, exhibit localized surface plasmon resonance (LSPR) effect upon light irradiation. The LSPR effects empower the plasmonic metal-based catalysts applicability in PNF. Over most of the plasmonic metal-based catalysts, the light harvesting unit and the active center work synergistically for improved performance. As the particle sizes, morphologies as well as the particle-particle distances of plasmonic metals are crucial for their light harvesting capacities, the tailoring of these properties are generally adopted with the aim to accelerate the reaction rate of plasmonic metal-based catalysts in PNF.

For example, Wang *et al.* encapsulated Au nanoparticles into a MOF membrane (Uio-66) and realized a NH₃ production rate of 359.1 µmol g⁻¹ h⁻¹ under visible light irradiation ($\lambda >$ 400 nm, 100 mW cm⁻²).⁹⁴ It was discovered that N₂ adsorbed on Au nanoparticles. Upon light irradiation, hot electrons generated on Au nanoparticles. The hot electrons on Au activated N₂ *via* two pathways, ① induced an electromagnetic field to polarize N₂ and ② directly injected into the anti-bonding orbitals of N₂ molecules (Fig. 6a). Moreover, the gas permeable nature of MOF membrane facilitated the mass transfer of the reactants, which further boosted its photocatalytic activity at



Fig. 5 Energy band structure of disordered TiO_2 (short for Na-A_d) and ordered TiO_2 (represented by R_o) as well as the reaction mechanism. This figure has been adapted from ref. 92 with permission from Royal Society of Chemistry, copyright 2020.

the gas (N₂)-membrane (Uio-66 encapsulated Au nanoparticles)solution (H₂O) interface (Fig. 6b).⁹⁴

Yang et al. reported that Au nanoparticles supported on nitrogen deficient g-C₃N₄ could catalyze PNF, achieving a NH₃ production rate of \sim 783 µmol g⁻¹ h⁻¹ under visible light irradiation.95 In this catalytic system, the nitrogen vacancy sites adsorbed N2; Au and g-C3N4 harvested visible light and induced electrons upon light irradiation, which then injected into N2 for its activation.95 Liao et al. synthesized small Ag doped g-C₃N₄ catalyst and unraveled that Ag enhanced N2 adsorption, generated more electrons, facilitated the separation and migration of photogenerated electron-hole pairs and consequently resulted in high activity in PNF.96 Wang et al. supported Au nanocrystals onto Mo doped W18O49 and realized a NH3 synthesis rate of \sim 399 µmol g⁻¹ h⁻¹.⁹⁷ Au not only harvested visible light but also decreased the desorption energy of the product NH₃, which accelerated the regeneration of the active sites for next catalytic cycle.97 Introducing a moderate amount of alkali metal cations could further promote N2 activation and then enhance the performance of these plasmonic metal-based catalysts in PNF.98

Iron-based catalysts for PNF

Iron is the most active metal for N_2 activation. The strong N_2 activation capacity of iron makes it applicable as active center for PNF. However, the light harvesting capacity of iron is very weak. Integrating iron with a material that could harvest light efficiently is an approach to design the photocatalyst for PNF. Based on this principle, doping iron into semiconductors, loading iron onto semiconductors, constructing a Fe-based heterostructure catalyst and loading iron onto other light harvesting unit and the active center are regulated. Up to now, various iron-based catalysts have been designed. In this section, we will discuss the types of iron-based catalysts for PNF and emphasize their progress.

(1) Fe doped into semiconductors. Doping Fe into a semiconductor could tune the local electronic structure of the

catalysts and thereby facilitate N2 activation. For example, when Fe is doped into TiO_2 , Fe will substitute Ti atoms in TiO_2 owing to their similar radii, which creates oxygen vacancies at the neighbor of Fe atoms to meet the local charge balance.⁹⁹ The Fe atoms and oxygen vacancies work in concert to facilitate N₂ adsorption and polarization, which enables N₂ hydrogenation via the favorable associative distal pathway and contributes to PNF. As shown in Fig. 7a and b, 5wt% Fe/TiO₂ (5-FTNFs) afforded a stable NH₃ production rate of \sim 64 µmol g⁻¹ h⁻¹ under the full spectrum illumination of a 300 W Xe lamp.99 Isotope labeling experiment as well as ¹H-NMR (NMR: Nuclear magnetic resonance, Fig. 7c and d) indicated that the generated NH₃ was originated from N₂ instead of other contaminants.⁹⁹ Similarly, Fe-doped BiOBr with oxygen vacancies afforded a NH3 yield of 46.1 $\mu mol~g^{-1}~h^{-1}$ without any sacrificial reagent under 400 mW cm $^{-2}$ visible light irradiation.¹⁰⁰ The doped Fe and the oxygen vacancies synergistically modulated the band structure, improved charge transfer and thereby enhanced the photocatalytic activities.100

Doping exerts functions beyond creating oxygen vacancies. For instance, the Fe atoms on the surface of Fe doped SrTiO₃ $(Fe_xSr_{1-x}TiO_3)$ could not only chemisorb and activate N₂, but also promote the electron transfer from $Fe_xSr_{1-x}TiO_3$ to N₂, which resulted in high N2 fixation capacity and a NH3 production rate of 30.1 μ mol g⁻¹ h⁻¹ over Fe_xSr_{1-x}TiO₃ (x = 0.1) under Xe lamp illumination.101 Doping Fe into Mo-based semiconductors could narrow their band gaps, extend their light absorption capacities as well as generate new Fe-Mo active centers, which enables Fe doped Mo-based semiconductor to harvest more solar light and facilitate the electron-hole separation and migration efficiency.¹⁰² Chang et al. doped Fe into 2D MoTe₂ nanosheets to construct Fe-Mo active centers.103 It is reported that Fe doped into MoTe₂ facilitated the separation and transfer of photoinduced electron-hole pairs, prolonged their lifetime and accounted for an obviously boosted NH3 production rate.103 Similarly, Fe doped $SrMoO_4$ (Fe/Sr = 1.6) achieved a NH_3 production rate of 93.1 µmol g⁻¹ h⁻¹ under Xe lamp illumination.¹⁰²

(2) Fe loaded onto semiconductors. Zhang *et al.* reported that $Fe/TiO_{2-x}H_y$ could serve as a dual temperature zone catalyst for



Fig. 6 (a) Schematic illustration of PNF over Uio-66 encapsulated Au nanoparticles. (b) Schematic illustration of the interface design for PNF. This figure has been adapted from ref. 94 with permission from Royal Society of Chemistry, copyright 2021.

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Fig. 7 (a) Performance of Fe doped TiO_2 (x-FTNFs, where x is the loading of Fe) in PNF. (b) Stability test of 5-FTNFs in PNF. (c) Timedependent NH₃ production over x-FTNFs. (d) Isotope labeling experiment (left panel) and 1H-NMR (right panel) of the products. This figure has been adapted from ref. 99 with permission from Royal Society of Chemistry, copyright 2021.

PNF.¹⁰⁴ Under solar light irradiation, catalyst surface temperature reached 495 °C, with a temperature difference between Fe and TiO_{2-x}H_y of 137 °C, owing to the plasmonic local heating effect of Fe. Then Fe acted as the hot zone catalyst to dissociate N₂ *via* its photogenerated hot electrons, while TiO_{2-x}H_y accommodated the N atoms from Fe and hydrogenated them into NH₃.¹⁰⁴ Fe/TiO_{2-x}H_y delivered a NH₃ concentration of 109.5 µmol g⁻¹ h⁻¹, an order of magnitude higher than the commercial Haber-Bosch Fe catalyst.¹⁰⁴

(3) Fe based heterostructure catalysts. Fe₂O₃/g-C₃N₄ prepared by thermal treatment method exhibited a NH₃ production rate of 7044 μ mol g⁻¹ h⁻¹ under the illumination of a 300 W Xe lamp, where the heterostructure facilitated light harvesting and Fe₂O₃ played key roles in N2 adsorption.105 Oxygen deficient Fe2O3/ZnO could stably produce NH₃ at a rate of 80 μ mol g⁻¹ h⁻¹ without any sacrificial agent under visible light irradiation.¹⁰⁶ Over Fe₂O₃/ ZnO, the synergistic effect between oxygen vacancies and Fe_2O_3 activated N2, while the heterostructure of Fe2O3/ZnO prohibited the recombination of electron-hole pairs, which accounted for its superior photocatalytic performance.¹⁰⁶ Fe-modified palygorskite supported FeS2, synthesized by microwave hydrothermal method, could serve as a Z scheme type photocatalysts for PNF and recorded a NH₃ production rate of 147 μ mol g⁻¹ h⁻¹ under solar light irradiation.107 Its activity was ascribed to the narrowed band gap, widened light harvesting region of Fe-modified palygorskite as well as the facilitated charge transfer between Femodified palygorskite and FeS2.107

(4) Other light responsive materials, such as graphdiyne and MOFs, supported Fe catalysts. For instance, Li *et al.* reported that the morphology, coordination environment and the valence state of iron oxide could be manipulated by encapsulating versatile shaped Fe_3O_4 by graphdiyne.¹⁰⁸ The encapsulation of graphdiyne endowed Fe_3O_4 @graphdiyne

heterojunctions strong light harvesting capacity, a structural evolution during PNF, as well as a NH₃ yield of an unprecedented level of ~1762 μ mol g⁻¹ h⁻¹.¹⁰⁸ The Fe atoms in Fe-based MOFs (*e.g.*, MIL-101(Fe), MIL-100(Fe), MIL-88(Fe)) have high electron densities, low reaction barriers for the activation of N₂ and H₂ to N–H bond, and could serve as the catalytic active center for PNF.¹⁰⁹ Taking MIL-101(Fe) as an example, it gave a NH₃ production rate of 100.7 μ mol g⁻¹ h⁻¹ under the illumination of a 300 W Xe lamp.¹⁰⁹

Ruthenium-based catalysts for PNF

Ruthenium is another metal capable of activating N_2 under thermally-driven conditions. Ruthenium-based catalysts that could harvest solar light effectively have been successfully applied in PNF. Since the light harvesting capacity of ruthenium is weak, the light harvesting unit of ruthenium-based photocatalysts are generally semiconductors or other light responsive materials. Similar as iron-based catalysts, over most of the ruthenium-based catalysts for PNF, both the light harvesting unit and the active center are regulated. Based on the nature of the light responsive components, these ruthenium-based catalysts could be divided into the following categories.

(1) Semiconductor, such as TiO₂, GaN, C₃N₄ and CeO₂, supported ruthenium catalysts. For example, Ru/TiO₂, in which singly dispersed Ru atoms were decorated onto TiO2 nanosheets rich in oxygen vacancies, is active for PNF.110 Over Ru/TiO2, the single Ru atoms were possibly located at the oxygen vacancy sites and stabilized by the vacancies.110 The isolated Ru atoms promoted the chemisorption of N₂, boosted the electron-hole separation and overall recorded a NH₃ generation rate of 3.3 μ mol g⁻¹ h⁻¹ upon irradiation by a 300 W high pressure Xe lamp.110 Ru/P25, prepared by the facile synthetic method, exhibited Ru particle sizes of 2-3 nm. In PNF, Ru/P25 dissociated H₂O to hydrogen atoms continuously and then hydrogenated N₂ molecules via a distal reaction pathway at the gas (N_2) -liquid (H2O) interface.¹¹¹ The NH3 yield over Ru/P25 was high up to 5.2 μ mol g⁻¹ h⁻¹ under the irradiation of a Xe lamp.¹¹¹ GaN supported Ru catalysts, Ru/GaN, behaved tailorable electronic and morphological properties.112 The interfacial Schottky junction between Ru and GaN facilitated the electron transfer from GaN to Ru, then the electron tank in Ru promoted N=N bond dissociation and achieved NH3 synthesis at low temperatures. Notably, 5 wt% Ru/GaN afforded an average NH₃ production rate of 120 μ mol g⁻¹ h⁻¹ after 2 h UV irradiation at 10 °C (Fig. 8).112

Modifying the semiconductor supported Ru catalysts could speed up NH₃ production rate. For instance, modifying Ru/g-C₃N₄ or Ru/TiO₂ by K could enrich the electrons in Ru, enhance the catalyst capacities in activating N₂ and consequently improve NH₃ generation rate.^{113,114} Doping Zr⁴⁺ into CeO₂ could increase the electron densities on Ce and create oxygen vacancies, which strengthened the interactions between Ru nanoparticles and supports.¹¹⁵ The strong interaction upshifted the Ru d-band center relative to Fermi level and enhanced N₂ cleavage.¹¹⁵ Ternary heterostructure Ru/RuO₂/g-C₃N₄ catalyst gave an average NH₃ production rate of 13.3 µmol g⁻¹ h⁻¹, 6 times higher than Ru/g-C₃N₄.¹¹⁶ Characterization results indicated that under light





irradiation, the electrons transferred to Ru whereas holes migrated to RuO₂ to facilitate the reduction and oxidation reactions, respectively, meanwhile, the electron-rich Ru activated N₂ effectively.¹¹⁶ Decorating Ru/g-C₃N₄ catalyst by S-deficient CoS_x could construct a bimetallic center at the interface of Ru/CoS_x, which facilitated N₂ polarization and activation *via* electron transfer from Ru and Co to N₂ upon light irradiation and ultimately gave a NH₃ production rate high up to 440 µmol g⁻¹ h⁻¹.¹¹⁷

(2) Other light responsive materials supported Ru catalysts. For example, coal-based carbon nanosheet supported Ru catalyst yielded 55.3 µmol g⁻¹ h⁻¹ NH₃ under a 300 W Xe lamp irradiation.¹¹⁸ Graphene oxide/silica could sufficiently disperse Ru and enhance the LSPR effect of Ru species, which excited more electron-hole pairs upon light irradiation and accelerated NH₃ generation.¹¹⁹ TiO₂-Mxene hybrid nanostructure supported Ru catalysts afforded an ammonia production rate of ~5.7 µmol g⁻¹ h⁻¹, which was principally stemmed from the synergetic effects among TiO₂, Mxene and Ru.¹²⁰

Other catalysts for PNF

Numerous other catalysts have also been designed for PNF. Phosphorus is one of them. Yu et al. disclosed that the edges of black phosphorus (BP) could absorb and reduce N2. They synthesized an edge-rich BP nanostructure with a flake-like shape via chemical etching exfoliation method. The edge-rich BP is of good dispersibility in H₂O, which allows its full contact with the reactants. Owing to the abundant active sites for N₂ chemisorption and reduction as well as the efficient contact between the reactants and catalyst, the edge-rich BP delivered a NH₃ production rate of 2370 µmol g⁻¹ h⁻¹.¹²¹ Lin et al. loaded red phosphorus (RP) onto photoinactive SiO₂ via a facile sublimation-deposition method, in which RP was modified by in situ formed carbon.122 The hybrid SiO2/C-RP catalyst was of large specific surface area, strong light harvesting capacity and high charge separation efficiency, which accounted for a NH₃ production rate of 36.5 μ mol g⁻¹ h⁻¹.¹²²

MOFs have also been successfully applied in PNF. Chen *et al.* unraveled that the Ce species in MOF-76(Ce) was an electron tank, which accepted photoinduced electrons to its 4f orbitals

and then donated the electrons to the anti-bonding orbitals of N_2 .¹²³ As a result, MOF-76(Ce) gave an average NH₃ yield of 34 µmol g⁻¹ h⁻¹ under ambient conditions.¹²³ Ye *et al.* reported that functionalized MIL-125(Ti) could act as photocatalysts for PNF under visible light irradiation without any sacrificial reagent.¹²⁴ Notably, amine-functionalized NH₂-MIL-125(Ti) afforded a NH₃ production rate of 12.3 µmol g⁻¹ h⁻¹. The electron transfer from the ligand to metal in MIL-125(Ti) induced Ti³⁺, which was the active sites for N₂ activation. Functionalization extended the light harvesting capacity of MIL-125(Ti) and further enhanced catalyst activity in PNF.¹²⁴

LDH with oxygen defects and electron-rich metals have been discovered active in PNF, in which vacancies and metal centers synergistically promote N₂ adsorption, facilitate the separation of photoinduced electron-hole pairs, and thereby boost activity in PNF.^{125,126} Zhang *et al.* uncovered that 0.5% mol Cu modification could impart ZnAl-LDH oxygen vacancies and electronrich unsaturated Cu^{δ +} (δ < 2);¹²⁵ NaOH treatment could also induce vacancies and low-coordinated metal centers in ZnCr-LDH, ZnAl-LDH and NiAl-LDH.¹²⁶ Taking Cu^{δ +}-ZnAl-LDH as an example, it realized a NH₃ production rate of 110 µmol g⁻¹ h⁻¹ under UV-vis light irradiation.¹²⁵

In addition, Mo-based catalysts (*e.g.*, $Mo_1/g-C_3N_4$),¹²⁷ carbontungstic-acid hybrids,¹²⁸ Pr³⁺:LiNbO₃,¹²⁹ Pt GO/SiO₂ (GO: graphene oxide)¹³⁰ have also been utilized in PNF. These catalysts produce NH₃ at the magnitude of µmol g⁻¹ h⁻¹. Meanwhile, the studies on these catalysts are quite limited and the reaction mechanism is not fully understood.

Summary and outlooks

Ideally, PNF is a safe and green approach to synthesize NH_3 under ambient conditions, using the inexhaustible solar light as the sole energy input and the abundant N_2 as a reactant. Catalyst is the key for PNF. Over the past few years, extensive studies have been conducted to search for efficient catalysts for PNF. Semiconductor, plasmonic metal-based catalysts, ironbased catalysts, ruthenium-based catalysts and several other catalysts, have been reported active in PNF. Table 1 lists the

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Table 1 The performance of some typical catalysts in PNF reaction

Catalysts	Catalytic types	Reaction conditions	$h_{\rm H3}$ yield ($\mu mol \ g^{-1} \ h^{-1}$
Ultrathin MoS ₂	Pristine semiconductors	Water (200 mL), catalyst (15 mg), reaction temperature (25 °C), N ₂ bubble, under light irradiation (500 W $2 > 420$ nm Ye lamn)	325
BiO quantum dots	Pristine semiconductors	With the section of the section temperature (25 °C), N ₂ bubble, under light irradiation (500 W. Xe lamb	1226
PFL-g-C ₃ N ₄ (PFL: porous few- laver)	Defective semiconductors	20% CH ₃ OH (100 mL), catalyst (20 mg), N ₂ (30 min), under light irradiation (500 W, $AM 1.5G_{100} \text{ mW cm}^{-2}$, $Re Jann)$	8200
D-CN (1D defective g-C ₃ N ₄)	Defective semiconductors	0.1 mol L^{-1} K ₂ SO ₄ solution (95 mL) and methanol (5 mL), catalyst (200 mg), reaction temperature (25 °C), N ₂ bubble (2 h), under light irradiation (600 mW cm ⁻²), reaction time (6.0 m cm ⁻²).	17.4
BOC/OV (surface oxygen vacancies modified micro-nanosheet structure Bi,O,CO ₃)	Defective semiconductors	utilie (8 II) 0.1 mmol L ⁻¹ Na ₂ SO ₃ solution (50 mL), catalyst (30 mg), N ₂ (60 mL min ⁻¹ , 30 min), under light irradiation (300 W, $\lambda > 420$ nm, Xe lamp)	14.7
A-SmOCl (amorphous SmOCl	Defective semiconductors	Water (20 mL), catalyst (10 mg), N_2 (30 mL min ⁻¹ , 30 min), under light irradiation (250 mW cm ⁻² 320–780 nm. Ye land)	426
ΠO_2 (B) nanotubes	Defective semiconductors	Mixture of water (90 mL) and methanol (10 mL), catalyst (25 mg), ultrasonic oscillation (10 min), reaction time (10 min), N_2 (30 min) under simulated sunlight irradiation (300 W. 60 min, AM 1.5G, xe lamb)	106
TiO ₂ -OVs (reduced TiO ₂)	Defective semiconductors	Mixture of water (90 mL) and methanol (10 mL), catalyst (50 mg), ultrasonic oscillation (15 min), N_2 (30 mL min ⁻¹), reaction temperature (25 °C), under light irradiation (300 W, Xe lamp)	324.86
BCN (B-doped g-C ₃ N ₄ nanosheets)	Doped semiconductors	Aqueous solution of Na ₂ SO ₃ (40 mL, 1.0 × 10 ⁻³ mol L ⁻¹), catalyst (20 mg), N ₂ (30 mL min ⁻¹ , 30 min), reaction time (1 h), under light irradiation (250 W, λ > 400 nm, 500 mW cm ⁻² . Xe lamp)	313.9
C-BiOI (carbon-doped BiOI)	Doped semiconductors	Mixture of water (90 mL) and ethanol (10 mL), catalyst (50 mg), under light irradiation (300 W. Xe lami)	311
NCN/MgO (g-C ₃ N ₄ nanosheets decorated with MgO nanovarticles)	Functionalized semiconductors	Water (4.0 mL) and 40 µL absolute ethanol (0.789 g L ⁻¹), catalyst (40 mg), ultrasonic oscillation (6 min), N_2 (1 h), reaction temperature (25 °C), under light irradiation (500 W. 100 mW cm ⁻² , $\lambda > 420$ nm. Xe lamp)	4554
KOH treated g-C ₃ N ₄	Functionalized semiconductors	CH ₃ OH (150 mL), catalyst (20 mg), reaction temperature (25 °C), under light irradiation (300 W, 100 mW cm ⁻² , Xe lamp)	3632
Cu/TiO ₂ (transition metal modified TiO ₂)	Functionalized semiconductors	5.0 vol% glycerol aqueous solution (200 mL), catalyst (100 mg), under simulating solar light irradiation (300 W, AM 1.5 filter, Xe lamp)	6780
P-LFO (phosphate modified LaFeO ₃)	Functionalized semiconductors	Water (40 mL), catalyst (20 mg), N_2 (1 h), under light irradiation (500 W, λ > 420 nm, Xe lamp)	250
Bi@BiOBr	Functionalized semiconductors	Water (100 mL), catalyst (10 mg), N_2 (80 mL min ⁻¹ , 30 min), reaction temperature (15 °C), under simulating solar light irradiation (300 W, 1.63 W cm ⁻² , Xe lamp)	1350
MOF@DF-C ₃ N ₄ (nano- MOF@defected thin film C ₃ N ₄)	Heterojunctions and homojunctions	Mixture of water (48 mL) and methanol (2 mL), catalyst (10 mg), ultrasonic oscillation (15 min), reaction temperature (25 °C) under light irradiation (300 W, $\lambda > 400$ nm, Xe lamp)	2320
MoS ₂ /C-ZnO	Heterojunctions and homojunctions	Mixture of water (190 mL) and ethanol (10 mL), catalyst (100 mg), reaction time (5 h), air, under light irradiation (300 W, $\lambda > 420$ nm, Xe lamp)	49.1
rio ₂ @C/g-C ₃ N ₄	Heterojunctions and homojunctions Heteroiunctions and	20 vol% CH ₃ OH (100 mL), catalyst (50 mg), N ₂ (60 mL min ⁻¹), under light irradiation (300 W, $\lambda < 420$ nm, Xe lamp)	250.6 432
	homojunctions		404

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Table 1 (Contd.)

Catalysts	Catalytic types	Reaction conditions	NH ₃ yield ($\mu mol \ g^{-1} \ h^{-1}$)	Ref.
Na-A _d /R _o (Na treatment of P25- TiO ₂ , A _d : Disordered anatase, R _O : disordered rutile)		Water (50 mL), catalyst (50 mg), isopropyl alcohol (7 mL), N ₂ (0.3 L min ⁻¹) under simulated AM 1.5G sunlight irradiation (1000 W, Xe lamp) in a double-layered jacket with cooling water circulating line		
${\rm Bi}_4{\rm O}_5{\rm Br}_2/{ m ZIF-8}$	Semiconductor-based hvdrophilic-hvdrophobic catalyst	Water (50 mL), catalyst (50 mg), N_2 (80 mL min ⁻¹) under simulated sunlight irradiation (300 W, 200–800 nm. Xe lamp)	16.4	93
Au@UiO-66	Plasmonic metal-based catalysts	Aqueeus solution of K_2SO_4 (50 mL, 0.5 mol L^{-1}), catalyst (15 mg), N ₂ (80 mL min ⁻¹ , 30 min), reaction temperature (25 °C) under light irradiation (300 W, λ > 400 nm, 100mW cm ⁻² . Xe lamp) in a convertional solution-based (<i>i.e.</i> , PiS) protocol	359.1	94
Au/HCNS-NV (HCNS: Hollow mesoporous carbon nitride sphere, NV: nitrogen vacancies)	Plasmonic metal-based catalysts	Mixture of water (80 mL) and methanol (20 mL), catalyst (50 mg), ultrasonic oscillation (10 min), reaction temperature (room temperature), N_2 (100 mL min ⁻¹ , 30 min), under light irradiation (300 W, Xe lamp)	783.4	95
Au/P25-K ⁺	Plasmonic metal-based catalysts	Water (50 mL), catalyst (5 mg), N_2 (60 mL min ⁻¹) under light irradiation (300 W, $\lambda > 400$ nm, 100mW cm ⁻² , Xe lamp)	430	98
Fe-MoTe ₂	Iron-based catalysts	Milli-Q (80 mL), catalyst (10 mg), reaction temperature (25 °C), N ₂ (50 sccm, 60 min), under light irradiation (300 W, $\lambda < 420$ nm, 400 mW cm ⁻² , h, Xe lamp)	129.08	103
MIL-101(Fe)	Iron-based catalysts	Water (100 mL), catalyst (50 mg), ultrasonic oscillation (10 min), reaction temperature (room temperature), N_2 (80 mL min ⁻¹ , 30 min), under simulated light irradiation (300 W, 1 h, Xe lamp)	100.7	109
Ru-TiO ₂	Ruthenium-based catalysts	20% ethanol solution (100 mL), catalyst (40 mg), reaction temperature (25 $^{\circ}$ C), N ₂ (1 h), light irradiation (300 W, Xe lamp), reaction time (4 h)	3.3	110
Ru-Vs-Cos/CN (Ru/Cos _x with S- vacancy on graphitic carbon nitride nanosheets)	Ruthenium-based catalysts	10% methanol solution (50 mL), catalyst (25 mg), \dot{N}_2 (30 min), light irradiation (300 W, 200 mW cm ⁻² , Xe lamp)	438	117
SiO ₂ /C-RP (RP: red phosphorous)	Other catalysts	Water (40 mL), catalyst (20 mg), ultrasonic oscillation, N_2 , reaction temperature (25 °C) under light irradiation (300 W, 320 mW cm ⁻² , Xe lamp)	36.5	122
Mo-PCN SACs (PNC: polymeric carbon nitride, SACs: single-atom catalysts)	Other catalysts	Water (6 mL), catalyst (3 mg), ultrasonic oscillation (60 min), pH = 5, N_2 , reaction temperature (room temperature), reaction time (12 h), light irradiation (300 W, Xe lamp)	830	127
Pr ³⁺ : LiNbO ₃	Other catalysts	Aqueous solution (100 mL), catalyst (40 mg), N ₂ (30 min), reaction temperature (30 °C), under light irradiation (300 W, $\lambda > 420$ nm, Xe lamp)	38.4	129

performances of some typical catalysts. This review summarizes the progress of each category of the catalysts designed for PNF, with a special attention on semiconductor-based catalysts. Generally speaking, the catalyst development is still in the infant stage and huge challenges need to be overcome.

(1) NH₃ production rates are still low. Over most of the catalysts, NH₃ production rates are in the magnitude of mmol g⁻¹ h⁻¹ or even µmol g⁻¹ h⁻¹. It is far away from the industrial applications. Meanwhile, other chemicals (*e.g.*, N₂H₄) are occasionally generated as byproducts. Therefore, persistent efforts should be devoted to design catalysts that could drive PNF efficiently and selectively to the desired product NH₃. Adopting novel materials as catalysts for PNF might benefit this research area.

(2) There is a long way to make clear the reaction mechanism. In spite that some studies carried out mechanism explorations, little progress have been made in understanding the fundamental mechanism. The physicochemical properties of the photocatalysts under working states remain unclear; N_2 chemisorption, activation and reduction pathway on the active sites are not clarified; the electron-hole transfer and migration routes need to be understood; in-depth understanding on the structure/property-performance correlations in PNF needs to be unraveled. Theoretical studies together with *in situ* characterization techniques might offer potential approaches to make clear the reaction mechanism.

In summary, despite that the development of catalysts for PNF is still in the primary stage, progress has been made. PNF has been proved as a promising avenue to replace the industrialized Haber-Bosch process to produce NH₃. Numerous opportunities exist to move the research field forwards.

Conflicts of interest

The authors declare no competing interests.

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