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Progress and challenges in photocatalytic ammonia synthesis

The group of Solar Energy and Advanced Materials, led by Prof. Junwang Tang focuses on photocatalytic small molecule activation (including CH₄, N₂, H₂O, C₆H₆ and CO₂) to high-value chemicals/fuels and microwave catalysis (e.g. plastic chemical recycling), together with microwave-intensified chemical processes. In parallel, the group has state-of-the-art spectroscopies to investigate the underlying charge dynamics and kinetics. This review puts forward a few critical issues existing in nitrogen photoreduction, along with a comprehensive discussion of strategies for improvement of NH₃ yield, aiming to make substantial contribution towards advancing this field.
Progress and challenges in photocatalytic ammonia synthesis

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Photocatalytic ammonia (NH₃) synthesis from N₂ and water driven by solar energy is a sustainable and environmentally friendly technology, which has gained considerable attention in recent years. In this review, the recent development in the fundamental understanding of photocatalytic NH₃ synthesis and the methods of precise NH₃ detection are summarized. More importantly the strategy for surface engineering and interface engineering of photocatalysts toward photocatalytic NH₃ production has been thoroughly analyzed with the aim to stimulate critical thinking about the effective methodology for catalyst modification instead of exploring new materials. At the end the challenges and a few concerns are raised from the current reports and future perspectives in this research field are discussed targeting to clarify the reliability and reproducibility of the photochemical process and to direct the future research direction, such as flow reactor design and in-depth understanding of the underlying reaction pathway.

Introduction

Ammonia (NH₃) has been regarded as one of the most important chemical products as feedstock for fertilizers and for various chemicals.¹,² The production of NH₃ plays a prominent role in global economy with an annual yield of more than 200 million tons.³ The industrial synthesis of NH₃ is dominated by the revolutionary Haber–Bosch process, in which pure H₂ and N₂ are reacted under high pressure at high temperature over Fe-based catalysts, which consumes 1–2% of total global fossil fuels and releases around 300 million tons of planet-warming CO₂ into the atmosphere annually.⁴–⁷ The development of environmentally friendly, sustainable strategies with high efficiency of NH₃ production under mild conditions is highly desirable but challenging.

Currently, there are a few green routes reported for ammonia synthesis. One is biological nitrogen fixation which mostly relies on diazotrophs in nature.⁸ Another one is electrochemical reduction of N₂, which uses electricity to produce NH₃.⁹,¹⁰ Compared with the stringent conditions of biological nitrogen fixation and the requirement of highly conductive electrolytes...
Photocatalytic mechanism of NH₃ synthesis

By mimicking the natural process of photosynthesis, researchers intend to convert solar energy, N₂ and water to NH₃ by using photocatalysts. The overall reaction is listed as follows:

\[2\text{N}_2 + 6\text{H}_2\text{O} \rightarrow 4\text{NH}_3 + 3\text{O}_2\]

In this artificial photosynthesis, there are two interlinked conversion processes, water splitting and N₂ reduction reaction (NRR). As shown in Fig. 1a, upon light irradiation, photoinduced electrons generated from photocatalysts populate the conduction band, leaving holes in the valence band. Subsequently, the escaped electrons diffuse across to the reactive sites on the surface of photocatalysts. The adsorbed N₂ and H₂O are catalyzed by these electrons and holes into NH₃ and O₂, respectively, along with the generation of the byproduct H₂. Thus the key point for this process is not only to design a highly active photocatalyst but also to achieve high selectivity to NH₃ instead of H₂ gas as indicated in Fig. 1a.
Up to now, the mechanisms for N₂ reduction to NH₃ on photocatalysts have been divided into two categories based on the surface adsorption characteristics of catalysts: N-hydrogenation (Fig. 1b) and surface-hydrogenation (Fig. 1c).²⁷,²⁸ For the N-hydrogenation mechanism, there are two different pathways for NRR, dissociative and associative pathways. In the dissociative pathway, the N₂ triple bond is broken before hydrogenation, followed by hydrogenation of the N-adatoms into NH₃ on the catalyst surface. The Haber–Bosch process is proved to be the dissociative pathway, which requires prohibitively high energy to cleave the N≡N bond. Compared with the dissociative pathway, the associative pathway without the breaking of the N≡N bond is generally used and accepted far more in the photocatalytic NRR. The nitrogen surface adsorption configuration determines the associative pathways for NRR; as shown in Fig. 1b, the hydrogenation of the adsorbed N₂ preferentially occurs at the terminal-site N owing to its relatively low steric hindrance for both the associative alternative pathway and the associative distal pathway. Having generated the first NH₃, the other N begins a new hydrogenation circulation to produce a second NH₃. It is actually impossible to identify the N₂ configurations due to the fact that they are favored equally by the catalysts.

Recently, a novel surface-hydrogenation mechanism for NRR has been proposed (Fig. 1c),²⁸ in which the adsorbed H⁺ is reduced into *H first, and then N₂ molecules react with surface *H to generate *N₂H₂, and finally the formed *N₂H₂ intermediates are reduced into NH₃. The first step is the trigger step, which enables the NRR to occur at a low potential. Furthermore, the second step is the rate determining step because it needs to conquer a high energy barrier (E_a) to break the N≡N bond. The E_a of hydrogen evolution reaction is always lower than that of NRR, and therefore the NH₃ production rate is relatively low.

As a complete cycle indicated in Fig. 1a is the ideal process but very challenging, in most cases (or in the majority of the reported cases), a half reaction was investigated in which N₂ was reduced to NH₃ in the presence of an efficient hole scavenger which was oxidized other than water oxidation. This half reaction is relatively easy and can be used to preliminarily scan different photocatalysts for N₂ reduction. Bear in mind that this half reaction has advantages but does not indicate whether the photocatalyst selected is good for a complete cycle as indicated in Fig. 1a. The subsequent discussion will underline this in order to remind the audience of the large difference between a half reaction and the complete cycle.

**Determination of NH₃**

NH₃ detection is a necessary subject for the photocatalytic synthesis of NH₃, which is of great significance in the design of high performance photocatalysts. There are four common methods of NH₃ analysis such as colorimetric assays,¹⁵N
isotope labeling, ion chromatography, and \textit{in situ} infrared spectroscopic measurements.\textsuperscript{12} The colorimetric assays include Nessler’s reagent, phenate, and indophenol blue tests for NH$_3$ detection. Among the colorimetric assays, Nessler’s reagent has been widely used in NH$_3$ detection due to its advantages of simplicity and low cost.\textsuperscript{29} However, it is reported that the sacrificial agents (i.e., alcohol) used in the photocatalytic synthesis of NH$_3$ systems can be oxidized to carbonyl compounds, which will disturb NH$_3$ detection in Nessler’s reagent detection, thus yielding misleading results. Fig. 2a shows photographs of a solution containing Nessler’s reagent and different concentrations of NH$_3$ solutions (0, 1.2, and 4 mg L$^{-1}$).\textsuperscript{29} It can be seen that the solution color is turning from light yellow to brown with the increase in the concentration of NH$_3$. When 4 µg L$^{-1}$ of formaldehyde is added, all the above solutions change into brown, indicating that trace formaldehyde can cause a remarkable color change (Fig. 2b). Therefore, the method of Nessler’s reagent is not able to detect NH$_3$ production in the presence of this or similar sacrificial agents. In contrast, the generated carbonyl compounds have no effect on the testing method of ion chromatography (Fig. 2c and d). It is strongly recommended that the colorimetric assay methods have to be accompanied by other methods such as ion chromatography, $^{15}$N isotope labeling, and \textit{in situ} infrared spectroscopic measurements, to avoid such misleading results. Fig. 2e shows the photocatalytic performance of NH$_3$ synthesis \textit{via} ion chromatography and the indophenol test.\textsuperscript{30} The UV-vis spectra of the reaction solution containing NH$_3$ exhibit a maximum absorption at 630 nm (inset in Fig. 2e), which is ascribed to the indophenols. The production of NH$_3$ measured by ion chromatography increases with the reaction time (Fig. 2e), consistent with the results obtained by the indophenol test, indicating that indophenol is to some extent reliable.

Extensive studies on the determination of NH$_3$ have revealed that the indophenol blue method, Nessler’s reagent method, and the ion chromatography method are accurate when the concentration of NH$_3$ is less than 500 µg L$^{-1}$, but the indophenol blue method tends to be less accurate at higher concentrations or under acidic conditions. Besides, the presence of carbonyl compounds from scavengers will enhance light adsorption in Nessler’s reagent method, thus leading to an interfered detection of NH$_3$. Thus, to guarantee accuracy and reliability of NH$_3$ quantification, at least two detection methods such as the ion chromatography method and Nessler’s reagent method are recommended. Furthermore, because of trace NH$_3$ in the ambient atmosphere, it is necessary to test the isotopically labeled $^{15}$N$_2$ to confirm the source of nitrogen in the product.
Strategies to improve photocatalytic efficiency

As mentioned above there are a few reviews summarizing the activity of diverse photocatalysts.6,13,18,19,22 This critical review will not overlap with these reviews, but will only concentrate on the efficient strategies to improve (i) N2 adsorption and (ii) charge separation. Such strategies include surface engineering and interface engineering for NH3 synthesis. The photocatalytic process of NH3 production generally includes four steps, photogeneration of charge carriers followed by their migration to the active sites, N2 chemisorption, surface photo-oxidation and photo-reduction, and the desorption NH3 from photocatalysts. Following the above key steps, photocatalysts can be engineered to greatly promote their catalytic activities toward NH3 synthesis. As steps (ii), (iii) and (iv) all take place on the surface, through the study of surface regulations and interface modulations on the photocatalysts, their photocatalytic performance can be maneuvered. The surface regulations, including defect engineering and morphology engineering, significantly determine the photocatalytic activity because they can promote the surface adsorption and activation of N2.23,31–40,54–57,64–71 The interface modulations, including modification with cocatalysts and semiconductors, greatly affect the charge transfer and separation efficiency.89–101,109–111 We will discuss these factors one by one below. In order to provide an intuitively systematic comparison of photocatalytic N2 reduction systems, we list photocatalysts, reactants, NH3 production rate, in particular the apparent quantum efficiency and detection method as shown in Table 1.

Surface regulations

Defect engineering. It is widely reported that vacancies in a photocatalyst can improve N2 adsorption, to some extent even charge separation. There are different populations of vacancies, including oxygen vacancies, nitrogen vacancies, carbon vacancies, as well as sulfur vacancies.

Oxygen vacancies. The introduction of oxygen vacancies (OVs) can effectively facilitate N2 adsorption and activation, as well as induce the generation of mid-gap states to promote the separation of photogenerated charge carriers.31–42 For instance, Zhang et al. reported that BiOBr nanosheets with OVs on the exposed [001] facets (BOB-001-OV) could stretch the N≡N bonds from 1.078 Å to 1.133 Å, which gave strong evidence for N2 activation over the OVs.31 Theoretical calculations showed that the OV induced defect states could serve as the electron acceptor (Fig. 3a) to effectively suppress the recombination of electrons/holes and enhance the charge transfer from BOB-001-OV to the N2 molecule. Fluorescence spectroscopy showed that the average lifetime (τ) of BOB-001-OV was 2.15 ns, around two times higher than that of BiOBr without OVs (BOB-001-H, Fig. 3b), proving that the OVs could promote the migration of charge carriers. With the function of OVs, BOB-001-OV showed a NH3 yield rate of 104.3 μmol g-1 h-1 under visible light illumination in the absence of organic scavengers and noble-metal cocatalysts (Fig. 3c), and an apparent quantum efficiency (AQE) of 0.23% at 420 nm. Meanwhile, the amount of generated O2 was stoichiometrically approaching 3/4 of the produced NH3, which proved that water could act as an electron donor to achieve a complete chemical cycle in NH3 photosynthesis.

Apart from the research on bismuth oxyhalide materials, Shiraishi’s group utilized a series of commercial TiO2 to study the effect of OVs on the photo-reduction of N2 to NH3.32 A commercial TiO2 of JRC-TIO-6 showed an excellent performance with a steadily increasing NH3 production within 100 h under UV light irradiation in pure water. They proposed that the Ti3+ species formed at the OVs on the surface of TiO2 served as active sites, which could effectively trap the electrons and promote N2 dissociation, resulting in a photocatalytic NH3 production rate of 2.5 μmol h-1.

In order to create the OVs, most studies used oxygen-containing catalysts. In recent years, layered double hydroxide (LDH, Fig. 4a) materials have attracted substantial attention due to their tunable electron structure and low cost.34 Zhang’s group firstly used the CuCr-LDH nanosheets (CuCr-NS) with OVs as efficient photocatalysts for NH3 photosynthesis.34 The OVs over CuCr-NS surface were created by the reduction of thickness and the in-plane compressive strain (Fig. 4b). The optimal CuCr-NS exhibited a NH3 production rate of 78.6 μmol g-1 h-1 under full-spectrum irradiation (Fig. 4c), and an AQE of 2.4% at 400 nm. Subsequently, they further reported another work on ZnAl-LDH nanosheets for optimal OVs by incorporating coordinately unsaturated Cu4+ species.35 The DFT calculations implied that the OVs and Cu4+ species in ZnAl-LDH could effectively promote N2 adsorption and activation, leading to a remarkable NH3 yield of 110 μmol L-1 h-1 in pure water under UV-vis irradiation. Therefore, the creation of OVs on the surface of photocatalysts paves a way for developing other innovative materials toward photocatalytic N2 reduction.

Nitrogen vacancies/carbon vacancies. Inspired by the conclusion of N2 activation and efficient electron transfer by OVs on the catalyst surface, nitrogen vacancies (NVs) were later studied for the NRR process. Because of its excellent optical properties, low cost, and good stability, graphitic carbon nitride (g-C3N4) has attracted much scientific interest in the field of photocatalysis.42–53 Dong et al. have reported the application of NV-incorporated g-C3N4 (V·g-C3N4) for N2 photoreduction.54 They found that the N2 adsorption amount of V·g-C3N4 was 75.1 cm3 g-1, 2.4 times higher than that of g-C3N4 without NVs (30.9 cm3 g-1), which suggested that the NVs on the surface of V·g-C3N4 could provide a number of chemisorption and activation sites for N2 molecules, leading to an NH3 production rate of 1.24 mmol h-1 gcat-1 under visible light irradiation in the presence of methanol as a hole scavenger. Recently, a g-C3N4 catalyst modified with cyano groups (–C≡N) and intercalated K+ (mCNN) was prepared as an efficient photocatalyst for NRR.55 Relative to pristine g-C3N4, the absorption spectrum of mCNN was extended to the whole visible region (Fig. 5a), resulting from the π–π* electronic transitions in the conjugated heterocyclic rings and the n–π* electronic transitions at the
active N defect sites (–C≡N), which indicated that the introduction of the N-defect of –C≡N could significantly change

the optical properties. Fig. 5b displays the band structures of mCNN and pristine g-C₃N₄ through the results of experiments.
Compared with pristine g-C₃N₄, a subgap state was presented for mCNN, which could be ascribed to the N defect, resulting in a broad visible-light response. The enhanced EPR intensity (Fig. 5c) of mCNN compared to pristine g-C₃N₄ was assigned to the strong electron withdrawing groups of –C⁻R⁻ on mCNN, which could delocalize the isolated electrons in mCNN, therefore promoting the separation of photocarriers and increasing the generation of active radical species to boost the activity of NH₃ photosynthesis. The rate of NH₃ generation of the mCNN photocatalyst was measured to be 3.42 mmol g⁻¹ h⁻¹ with ethylene glycol as a scavenger under visible light irradiation (Fig. 5d), which was much higher than that of pristine g-C₃N₄ (1.11 mmol g⁻¹ h⁻¹). The DFT calculations revealed that K⁺ could be linked to the unsaturated C centers by coordination as shown in Fig. 5e, which could adsorb N₂ as the lone-pair electrons of nitrogen would fill the empty state in K⁺ ($D_G = 0.28$ eV), followed by the rearrangement of K⁺ with the C atom and the adsorption of N₂ ($D_G = 0.17$ eV). Because of the large ion hydration free energy of K⁺, it could be extracted back into solution, then forming a C₂N₄ ring ($D_G = 1.64$ eV) intermediate which could undergo a MVK process to regenerate –C≡N, thereby stabilizing the unsaturated C sites.

Apart from the above NVs on the g-C₃N₄ photocatalyst, carbon vacancies (CVs) on a porous sulfur-doped g-C₃N₄ (SCNNSSs) have also been exploited. The introduction of CVs was proved to play the same role as NVs, while the S-doping effect was demonstrated to improve the separation efficiency of photogenerated electron–hole pairs. Taking these advantages, the optimal SCNNSSs showed an NH₃ production rate of 5.99 mmol h⁻¹ g⁻¹.

**Sulfur vacancies.** Considering the importance of the sulfur element in nitrogenases, a few have reported the effects of sulfur vacancies (SVs) on N₂ photoreduction. It was reported that SVs could be created on the surface of Mo₀.₁Ni₀.₁Cd₀.₈ by co-doping Mo and Ni into CdS. N₂-temperature-programmed desorption (N₂-TPD) results illustrated that the SVs on the surface of Mo₀.₁Ni₀.₁Cd₀.₈ obviously enhanced N₂ chemisorption (Fig. 6a), which was beneficial for NRR. Importantly, the photocatalytic NH₃ generation rate of Mo₀.₁Ni₀.₁Cd₀.₈ was linearly related to its SV concentration as shown in Fig. 6b. As the concentration of SVs increased, there was an increase in the photocatalytic NH₃ production rate. This result suggested that the SVs played a vital role in the photoreduction of N₂. The Mo₀.₁Ni₀.₁Cd₀.₈ with the highest SV concentration showed the highest NH₃ production rate of 3.2 mg h⁻¹ g⁻¹ (Fig. 6b).

**Doping.** Metal doping is a very promising strategy to change the electronic structure and surface property of photocatalysts to enhance the photocatalytic activity. In one report,
Schrauzer et al. studied the effect of different metal doped TiO₂ on the photocatalytic activity of N₂ reduction, including iron (Fe), chromium (Cr), cobalt (Co), and molybdenum (Mo).¹⁷ They found that Fe doping was the most effective for enhancing the photocatalytic activity of TiO₂ for photosynthesis of NH₃ from N₂ and H₂O. However, O₂ production as the oxidation product was not reported. In this regard, Zhao et al. investigated the Fe-doped TiO₂ with highly exposed (101) facets for N₂ photofixation,⁶⁴ and found that an optimum Fe³⁺ doping content played a key role in inhibiting the recombination of photoinduced electron–hole pairs, which could act as a temporary electron/hole trapping sites, therefore enhancing the concentration of charge carriers and improving the photocatalytic performance. With this in mind, Liu et al. also reported a Fe-doped SrMoO₄ (FSMO) as a potential candidate for N₂ photoreduction.⁶⁵ Further studies revealed that the intrinsic bandgap of SrMoO₄ could be shrunk from 3.98 eV to 2.93 eV with the increase in Fe doping concentration (from 0 to 5.1%), resulting in the extension of light adsorption from the ultraviolet to the visible-light region. Besides that, the Fe doping could induce the formation of surface defects as active sites for N₂ adsorption and significantly retard the recombination of electrons and holes, leading to enhanced N₂ reduction reaction. As a result of these properties, an improved NH₃ production rate of 93.1 μmol g⁻¹ h⁻¹ over the optimal FSMO was achieved compared with that of pristine SrMoO₄ (66.7 μmol g⁻¹ h⁻¹). In another related work incorporating Fe into BiOCl nanosheets, Fe-doped BiOCl nanosheets (BiOCl NsS-Fe) were developed for N₂ photoreduction.⁶⁶ The optimal BiOCl Nss-Fe exhibited a marked enhancement of photocatalytic NH₃ production, and the efficiency was 2.53 times higher than that of pristine BiOCl NSs.

In another study, Mo was successfully doped into W₁₈O₄₉ nanowires to produce Mo-doped W₁₈O₄₉ nanowires (MWO-1) as shown in Fig. 7a.⁶⁷ Compared with OV-rich W₁₈O₄₉ nanowires, Mo-doping had many kinds of effects on photocatalytic N₂ reduction. Fig. 7b displays the electronic band structures of MWO-1 and W₁₈O₄₉; it is observed that the defect-band center could be shifted to the Fermi level by Mo doping, which provided more energetic electrons for NRR. Theoretical simulations revealed that the Mo–W centers could alter electron distribution, leading to a larger adsorption energy (ΔG₂ = 2.48 eV) relative to W₁₈O₄₉ with W–W sites (1.65 eV), which was in favor of N₂ chemisorption and activation (Fig. 7c). Additionally, the enhanced M–O co-valence caused by Mo doping could effectively promote the electron transfer from metal active centers to adsorbed N₂ molecules. Thus, the optimal MWO-1 exhibited an excellent NH₃ yield rate of 195.5 μmol g⁻¹ h⁻¹ under full-spectrum irradiation (Fig. 7d), which was 7 times higher than that of W₁₈O₄₉, and also realized a high AQE of 0.33% at 400 nm.

Furthermore, Cu-doped ultrathin TiO₂ nanosheets (TiO₂-V₁₀ strain) were also studied.⁶⁸ Compared with pristine TiO₂, introducing Cu into TiO₂ could create abundant OVs and cause lattice distortion and strain effects, resulting in a significant increase of N₂ adsorption energy on the surface of TiO₂-V₁₀ strain (ΔG = 0.37 eV) compared with that of pristine TiO₂ (ΔG = 0.17 eV, Fig. 8a). The increased adsorption energy promoted the electron transfer from TiO₂ to N₂, thereby breaking the N≡N bonds. Meanwhile, the required reaction energy for N₂ hydrogenation to

Fig. 4 (a) Schematic representation of the layered LDH structure with defective MO₆ octahedra. (b) Schematic of the in-plane biaxial compressive strain in CuCr-NS. (c) The yield of NH₃ over different LDH photocatalysts under visible-light illumination (λ > 400 nm). Reproduced from ref. 34 with permission from Wiley-VCH.
N–NH* was only 0.365 eV on TiO$_2$-Vo-strain, which was lower than that on pristine TiO$_2$ (2.115 eV, Fig. 8b). A result, the optimal TiO$_2$-Vo-strain (6% TiO$_2$) achieved an enhanced NH$_3$ yield rate of 78.9 mmol g$^{-1}$ h$^{-1}$ under full solar irradiation, about 5.2 times higher than that of the TiO$_2$ nanosheets with OVs (0% TiO$_2$, Fig. 8c). In addition, the yield rate of O$_2$ (59.1 mmol g$^{-1}$ h$^{-1}$) and NH$_3$ mentioned above over TiO$_2$-Vo-strain was close to the stoichiometric ratio of 3:4, revealing that the protons in NH$_3$ were from the H$_2$O molecules. Most recently, a novel bimetallic system with Fe–Pt loaded g-C$_3$N$_4$ was used for efficient ammonia synthesis under mild conditions. Further investigations proposed that the doping of Pt onto the Fe nanocluster over the surface of g-C$_3$N$_4$ could cause an uplift of the energy band of semiconductors and form a large Schottky barrier, thus leading to an improved separation of photogenerated carriers and enhanced N$_2$ reduction. Accordingly, 0.3 wt% Pt doped on 3 wt% Fe@g-C$_3$N$_4$ exhibited a high NH$_3$ production rate of 63 μg h$^{-1}$ g$^{-1}$ with gaseous H$_2$ and N$_2$ as reactants under visible light irradiation and the AQE was tested to be 0.15% between 450 and 500 nm.

Apart from metal doping, a metal-free B-doped g-C$_3$N$_4$ nanosheet (BCN) with exposed active N atoms was recently synthesized for highly efficient ammonia synthesis. Theoretical studies revealed that the exposed N atoms could be stabilized by B–N–C coordination on BCN, which was different from that in the pristine g-C$_3$N$_4$. The B dopants were proved as active sites for N$_2$ adsorption and activation and could effectively retard charge recombination and improve light utilization. The optimal BCN with 13.8 wt% B-dopants exhibited a remarkable NH$_3$ yield rate of 313.9 mmol g$^{-1}$ h$^{-1}$ under visible light irradiation in the presence of Na$_2$SO$_3$ as a hole scavenger, which was much higher than that (32.8 mmol g$^{-1}$ h$^{-1}$) over pristine g-C$_3$N$_4$ and also achieved a good QE of around 0.64% at 420 nm. In addition, carbon-doped TiO$_2$ was also reported for N$_2$ photoreduction, which exhibited an NH$_3$ yield rate of 109.3 mmol g$^{-1}$ h$^{-1}$.

**Morphology engineering**

Facets. Since the surface atomic distribution has a huge effect on the active sites and electronic structures of photocatalysts, many researchers attempted to control the crystal facet of semiconductors to enhance the photocatalytic N$_2$ reduction. For instance, two Bi$_2$O$_3$-I nanosheets with different {100} and {001} facets were synthesized successfully via hydrolysis and calcination methods, respectively. Further investigations...
revealed that Bi$_5$O$_7$I-001 had more negative conduction band position (−1.45 eV) compared with that in Bi$_5$O$_7$I-100 (−0.85 eV) and exhibited higher efficiency for separation of photoinduced carriers, thus leading to a high photocatalytic activity for N$_2$ photoreduction. Accordingly, a remarkable NH$_3$ generation rate of 111.5 μmol g$^{-1}$ h$^{-1}$ over Bi$_5$O$_7$I-001 was achieved using methanol as a hole scavenger and the AQE of that was increased to 5.1%.

In another report, Zhang et al. also systematically investigated the influence of {001} and {010} facets of OV-rich BiOCl nanosheets on N$_2$ adsorption and activation. Experimental results showed that {010} facet BiOCl nanosheets (BOC-010) exhibited a superior performance with a NH$_3$ yield of 4.62 μmol g$^{-1}$ h$^{-1}$, which was around 2.5 times higher than that on {001} facet BiOCl nanosheets (BOC-001, Fig. 9a). Meanwhile, it was observed that N$_2$H$_4$ which served as a main intermediate was accumulated on BOC-010 during N$_2$ photoreduction reaction within 30 min and was consumed gradually to be converted into NH$_3$ (Fig. 9b). Importantly, when N$_2$H$_4$ was used as the reactant, both BOC-010 and BOC-001 showed a similar photocatalytic conversion efficiency of N$_2$H$_4$ under simulated solar irradiation (Fig. 9c), which in turn meant that BOC-010 had a stronger ability to generate N$_2$H$_4$ intermediates. Further DFT calculations revealed that the N$_2$ fixation via proton-coupled electron transfer compared with that via a distal pathway occurring on BOC-001 facets, thus resulting in enhanced NH$_3$ production (Fig. 9d and e).

**Nanostructure engineering.** Nanomaterials have preternatural interface structures and unique functions, such as the small size effect, the surface and boundary effect, quantum size and so on. In order to increase the catalytic active sites of the photocatalytic reduction of N$_2$, Sun et al. synthesized BiO quantum dots with an average size of 2–5 nm (Fig. 10a), which showed an NH$_3$ production rate of 1226 μmol g$^{-1}$ h$^{-1}$ without a cocatalyst and a sacrificial agent. Meanwhile, the amount of produced O$_2$ increased continuously with increasing irradiation time, consistent with the stoichiometric chemistry. Kinetic analysis and quantum chemical calculations suggested that the highly efficient photocatalytic activity of BiO quantum dots could be attributed to the fact that the smaller size endowed the surface and edge of BiO rich in Bi$^{2+}$ species, which had the potential to increase the electron donation to the anti-bonding π$^*$ orbitals of N$_2$, thereby could act as active sites to adsorb and activate N$_2$ (Fig. 10b). More recently, a tubular OV-rich Bi$_2$O$_3$-Br (Bi$_2$O$_3$-Br-NT) with a diameter of 5 nm (Fig. 10c) was prepared and it exhibited a maximum NH$_3$ yield rate of 1.38 mmol g$^{-1}$ h$^{-1}$ (Fig. 10d) upon changing the OV concentration.

A high specific surface area of up to 96.56 m$^2$ g$^{-1}$ was observed for Bi$_2$O$_3$-Br-NT, which indicated the exposure of abundant surface
OVs for the chemisorption and activation of N₂. It was found that the OVs on the Bi₅O₇Br-NT surface could not only effectively promote N₂ adsorption and activation, but also could be regenerated by capturing O atoms from H₂O after the reaction to maintain the good stability of Bi₅O₇Br nanostructures. As a result, the calculated AQE for the Bi₅O₇Br-NT photocatalyst was 2.3% under visible light irradiation at 420 nm (Fig. 10e). Furthermore, it was reported that metal-free black phosphorus nanoflakes with abundant edges (eBP NFs) synthesized via a facile chemical etching exfoliation method exhibited a remarkable NH₃ yield rate of 2.37 mmol g⁻¹ h⁻¹ under visible light irradiation in the presence of Na₂SO₃ and Na₂S·9H₂O as scavengers.88 The photoelectrochemical characteristics (PEC) and transient absorption (TA) studies revealed that the efficient ammonia synthesis was attributed to rich edges on the surface, which could provide abundant active sites for enhanced N₂ adsorption and activation. Similarly, a nanocomposite (SiO₂/C-RP) prepared by loading red phosphorus on SiO₂ nanoparticles showed superior charge separation.89 Besides, this hybrid nanostructure exhibited a large surface area, good water dispersibility and large light adsorption. As a result of these benefits, SiO₂/C-RP exhibited an NH₃ production yield of 0.73 μmol h⁻¹ under full-spectrum irradiation.

Interfacial modulation

Cocatalyst loading. With the similar idea of using metal cocatalysts to inhibit the recombination of electrons and holes and enhance the photostability of catalysts in other photocatalytic processes, e.g. water splitting,90–102 Ranjit et al. studied different metal loaded TiO₂ toward the photoreduction of N₂, including ruthenium (Ru), rhodium (Rh), palladium (Pd), and platinum (Pt).93 It was found that the order of the photocatalytic activity of these metals on TiO₂ was Ru > Rh > Pd > Pt. The excellent performance of Ru-loaded TiO₂ could be assigned to the high metal–hydrogen bond strength which well correlated with the high production of NH₃. In recent years, single-atom catalysts (SACs) have been well studied as excellent candidates for boosting the catalytic activity and improving the utilization efficiency of metal atoms during the photocatalytic process.94 To further improve the performance of the photocatalytic NH₃ synthesis, a single atomic Ru decorated TiO₂ nanosheet with abundant OVs was prepared.95 The DFT calculations indicated that the single atomic Ru species were favorable to be formed and dispersed on the OVs over the TiO₂ nanosheet surface. Further investigations revealed that the isolated Ru atom could serve as the active site for N₂ adsorption and activation, promote the separation of photogenerated charge carriers efficiently, as well as inhibit the competitive H₂ evolution reaction, thus leading to an improved NH₃ photosynthesis. Thus, a 1 wt% single Ru atom decorated TiO₂ nanosheet exhibited a superior performance with an NH₃ yield of 56.3 μg h⁻¹ g⁻¹ cat⁻¹, which was two times as high as that of pure TiO₂. Recently, Au-anchored TiO₂ nanosheets with OV (Au/TiO₂-OV) photocatalysts were developed for N₂ photofixation.96 As shown in Fig. 11a, Au nanospheres...
with a diameter of 20 nm ± 1.4 nm were uniformly distributed on the surface of TiO$_2$-OV nanosheets. The UV-vis spectrum of Au/TiO$_2$-OV showed absorption characteristics of Au loaded TiO$_2$ (Au/TiO$_2$) and TiO$_2$-OV, displaying a broad localized
surface plasmon resonance (LSPR) peak at 550 nm and an obvious absorption tail (Fig. 11b). Therefore, in the Au/TiO\textsubscript{2}-OV hybrid, OVs on the surface of TiO\textsubscript{2} nanosheets chemisorbed and activated N\textsubscript{2} molecules, which were reduced to NH\textsubscript{3} by plasmon-induced hot electrons from Au nanoparticles in the presence of methanol as a hole scavenger (Fig. 11c). As a result, the optimal Au/TiO\textsubscript{2}-OV exhibited an NH\textsubscript{3} photoproduction rate of 78.6 \textmu mol g\textsuperscript{-1} h\textsuperscript{-1}, which was 98 and 35 times higher than that of Au/TiO\textsubscript{2} and TiO\textsubscript{2}-OV respectively (Fig. 11d). Recently, we reported a hybrid of 2D layered Ti\textsubscript{3}C\textsubscript{2} and P25 TiO\textsubscript{2} (Ti\textsubscript{3}C\textsubscript{2}-P25), in which Ti\textsubscript{3}C\textsubscript{2} served as the cocatalyst to facilitate charge separation due to its excellent electrical conductivity. The DFT calculation (Fig. 11e) revealed that the N\textsubscript{2} adsorption energies were ranked as Ti\textsubscript{3}C\textsubscript{2} (2.731 eV) > VOs-TiO\textsubscript{2} (0.342 eV) > TiO\textsubscript{2} (0.170 eV), implying that Ti\textsubscript{3}C\textsubscript{2} had a much stronger N\textsubscript{2} chemisorption effect compared with VOs-TiO\textsubscript{2} and TiO\textsubscript{2}. Therefore the introduction of Ti\textsubscript{3}C\textsubscript{2} on P25 could accelerate N\textsubscript{2} chemisorption, leading to five times higher photocatalytic yield of NH\textsubscript{3} than that on pure P25. More importantly, water acted as the electron donor in this study and stoichiometric O\textsubscript{2} production was also observed, which was one of a few examples that could observe the stoichiometric reduction and oxidation products.\textsuperscript{103}

In addition to TiO\textsubscript{2}-based photocatalysts, Pt-loaded ZnO was reported as an efficient photocatalyst for N\textsubscript{2} photoreduction which exhibited an NH\textsubscript{3} yield rate of 860 mmol g\textsuperscript{-1} h\textsuperscript{-1}.\textsuperscript{104} More recently, O-doped 1T-MoS\textsubscript{2} nanosheets with a large amount of SVs (SV-1T-MoS\textsubscript{2}) were used as cocatalysts over CdS nanorods for...
photocatalytic N₂ reduction. The DFT calculations revealed that the SVs and the metal properties of 1T-MoS₂ could effectively promote the separation of h⁺/e⁻ and the presence of SV-1T-MoS₂ could also provide abundant active sites, resulting in enhanced N₂ adsorption and immobilization. Accordingly, a remarkable performance over the optimal SV-1T-MoS₂/CdS was realized with a superior NH₃ yield rate of 8220.83 μmol L⁻¹ h⁻¹ g⁻¹ under simulated solar light irradiation. Besides, NiS was also studied as a cocatalyst over CdS nanorods for ammonia photosynthesis. The DFT calculation indicated that the NiS cocatalyst could reduce the adsorption energy of N₂ from 1.16 eV on the CdS surface to 0.55 eV on the Ni-doped CdS surface. Further characterization revealed that charge separation could also be enhanced by loading NiS on the CdS surface. Because of these properties, a favorable NH₃ yield of 2.8 mg L⁻¹ was reported within the first hour in pure water under full-spectrum irradiation. Recently, single-atom Pt modified triazine framework (CTF) nanosheets (Pt-SACs/CTF) were developed for photocatalytic NH₃ production. It was found that the Pt–N₃ sites were formed in Pt-SACs/CTF, leading to more negative conduction band position with accelerated thermodynamics and fast interfacial charge migration ability, and therefore a high NH₃ production rate of 171.4 μmol g⁻¹ h⁻¹ under visible light irradiation. Most recently, we synthesized a ternary Ru/RuO₂/g-C₃N₄ system for N₂ photoreduction, which exhibited a photocatalytic NH₃ yield of 13.3 μmol g⁻¹ h⁻¹ whilst no NH₃ could be measured for pure g-C₃N₄ under the same condition. Such significant improvement in photocatalytic NH₃ synthesis for Ru/RuO₂/g-C₃N₄ was not only because of Ru and RuO₂ acting as cocatalysts to promote electron and hole transfer respectively, but also due to the advantage of Ru for N₂ chemisorption and activation. Besides that, carbon-based materials especially graphene are also often used as cocatalysts to prevent the photoinduced electron–hole recombination and promote the electron transfer due to their prominent electrical conductivity. For instance, a Fe modified three-dimensional graphene (Fe@graphene) photocatalyst was reported for NH₃ production. In the Fe@graphene system (Fig. 11f), graphene generated hot electrons under light irradiation and transferred those electrons to Fe, while Fe acted as an electron sink and provided the catalytic active sites for the adsorption and activation of N₂.

**Junction structure.** The combination of two semiconductors as a photocatalyst would overcome the shortages of each component, and construct the heterostructure or Z-scheme system with improved photoelectric properties for broad applications. In one work, Bi₂MoO₆ was implanted on g-C₃N₄ nanosheets to enhance the photocatalytic N₂ reduction. In this case, the obtained g-C₃N₄/Bi₂MoO₆ heterojunction showed effective separation of charge carriers and enhanced photocatalytic performance for NH₃ production. In another report, MXene-derived TiO₂@C incorporated with g-C₃N₄ (TiO₂@C/g-C₃N₄, Fig. 12a) was prepared for efficient NH₃ photocatalysis. Compared with the pristine g-C₃N₄, TiO₂@C/g-C₃N₄ showed a significantly decreased PL intensity (Fig. 12b), which indicated that the heterostructure promoted the charge transfer, which in turn inhibited the recombination of photoinduced electron–hole pairs. Furthermore, the transient photocurrent density of TiO₂@C/g-C₃N₄ was about 1.7 times higher than that of g-C₃N₄ (Fig. 12c), suggesting the improved separation efficiency of the charge carriers. The N₂-TDP result revealed that the Ti⁺³ species on TiO₂ and carbonaceous components were responsible...
for the adsorption and activation of N₂ in TiO₂@g-C₃N₄ (Fig. 12d). As a result, a superior performance over the optimal TiO₂@g-C₃N₄:10 (10 represents the molar ratio of starting materials of melamine/Ti₃C₂) was realized with an NH₃ production yield of 250.6 μmol g⁻¹ h⁻¹ using methanol as a scavenger under visible light irradiation (Fig. 12e). Recently, a p-n junction of Bi₂MoO₆/OV-BiOBr combining n-type Bi₂MoO₆ nanorods with OV-rich p-type BiOBr nanosheets displayed efficient charge separation and a broad range of light absorption. Moreover, OVs on the surface of Bi₂MoO₆/OV-BiOBr were beneficial for the chemisorption and activation of N₂ and the nanosheet morphology could provide a large specific surface area with more active sites for photoreduction of N₂. Benefiting from these advantages, the optimal Bi₂MoO₆/OV-BiOBr exhibited a photocatalytic yield of NH₃ of 81.0 μmol g⁻¹ h⁻¹ under visible light illumination in pure water, and no O₂ production was reported here.

**Conclusion and perspectives**

The photosynthesis of NH₃ is an energy-saving process and has high potential for substantial contribution to economic and social sustainability. Compared with the traditional Haber–Bosch process and the emerging electrochemical process, photocatalytic NH₃ synthesis exhibits very low efficiency, mostly at micromoles g⁻¹ h⁻¹ if using water as the electron donor, and thus is far from practical application. However as photocatalysis can be driven by abundant solar energy, it is a carbon-free process. Besides, it requires a solar energy input of 208.3 MJ kg⁻¹-NH₃ compared with that of 339.1 MJ kg⁻¹-NH₃ in the electrochemical process. Thus, it has strong potential to achieve NH₃ synthesis with affordable costs in particular taking into account the sustainability and the trend of decarbonization in the industry. However there is a long way to go in order to realize this potential.

This review does not detail different photocatalysts as there are a few reviews on them, but analyses surface engineering and interface engineering in photocatalysts developed recently for the photocatalytic NH₃ synthesis, involving oxygen vacancies, nitrogen vacancies, carbon vacancies, sulfur vacancies, metal and non-metal doping, facet modulation, nanostructure engineering, and heterostructure construction such as loading with cocatalysts and modification with other semiconductors, together with the mechanism of N₂ reduction reaction and the reliable means for the detection of NH₃.

Great progress has been made in the study of photoreduction of N₂ to NH₃ as briefed in Fig. 13, but there are still many problems, some of which are very critical. First, the majority of the studies so far reported use an efficient and expensive hole scavenger to get long lived photoelectrons for N₂ reduction. However, a practical pathway should use water as the only electron donor and to provide protons for the photosynthesis of NH₃, as such NH₃ synthesis by photocatalysis is scientifically meaningful and economically sustainable. The slow progress linked to this profile is mainly due to the very limited understanding in this field. Second, there are few reports specifying the stoichiometric chemistry in the photochemical process. Without the stoichiometric O₂ gas produced, such N₂ reduction to NH₃ is questionable. Furthermore, as O₂ could be leaked from air, the isotopic measurement of O₂ production is necessary. In addition, since H₂ production would compete with NH₃ synthesis, a comprehensive study should detail the selectivity between the two products and discuss the strategy to favor NH₃ synthesis. In addition an experimental error bar is very crucial for such NH₃ synthesis in order to improve the confidence as almost all studies were carried out in a batch reactor with quite a small amount of NH₃ produced, which was quite easy to be interfered by occasional factors.

Apart from the issues mentioned above, photocatalytic NH₃ synthesis also faces a few tough challenges. The majority of the studies show that the NH₃ amount produced is less than 10 μmol h⁻¹ (instead of the unit of μmol g⁻¹ h⁻¹) when using water as the electron donor and 10 times enhancement is achieved when using an organic hole scavenger. This amount is lower than the NH₃ amount in the river water which is about 0.01 mM. Besides, since the NH₃ yield is extremely easy to be interfered by environmental impurities and different detection agents, it is difficult to obtain reliable and reproducible results by just one analysis method. At least two methods have to be applied to prove the consistent results and to improve the reproducibility of the process.

The surface states of a catalyst can dominate its performance, such as surface defects. However, to quantify the correlation of the surface defects with the catalytic activity is very challenging at present. This should be undertaken in the future in order to distinctly guide catalyst surface state control. Furthermore the cocatalyst is another key factor to control both charge separation and catalytic performance. The majority of the cocatalysts reported are either large particles or nanosized particles. Single atomic catalysis is an emerging area, which...
presents unprecedented activity in some chemical processes. Such unique reaction sites should be introduced to ammonia synthesis with in-depth exploration of their intrinsic function. In addition, a junction structure is widely accepted to help charge separation, which was also reviewed in the previous session. The new design strategy should be applied to construct new junctions, such as polymer and oxide junctions as the former has likely better N₂ adsorption due to the potential hydrogen bond between the organic polymer and N₂ while the latter has a very positive VB for water oxidation.

The modified catalyst discovery is very informative. Equally, the mechanism and reaction pathways for photocatalytic N₂ fixation are scarce and much less investigated, forming an obstacle for the rational design of highly efficient photocatalysts. Currently, most of the given reaction mechanisms and reaction pathways were based on assumptions and lacked solid evidence. Thus, the current catalyst development is more or less a try-and-error process. Some advanced operando technologies in functional characterization should be applied to comprehensively study the reaction pathway such as by time resolved spectroscopy to investigate charge carrier dynamics, and by transient and stationary spectroscopy to observe the reaction intermediates.

In addition, it is of paramount importance to design a flow reaction system for N₂ photoreduction. The current reaction system is composed of a batch reactor, which is far from the requirement of the industry. The typical batch reactor is easy to be constructed; however, its efficiency is severely restricted by the low mass transfer between N₂ gas and aqueous solution due to the low solubility of N₂, thereby resulting in a low photo-catalytic efficiency. Besides, NH₃ accumulated in the batch reactor during the reaction process will be oxidized to other byproducts, thereby resulting in a low conversion rate and even bad selectivity. Moreover, based on the industrial viewpoint, it is quite costly to separate the photocatalysts from reactants and products in a batch system. Thus, an intensified flow reactor such as using N₂ and water vapor as reactants should be explored to overcome these drawbacks and to meet the industrial requirement in the future, underlining the crucial collaborations between chemical scientists and experts in reaction engineering.

Last, modelling will be expected to combine with experimental results to figure out the reaction mechanism and speed up the rational design of highly efficient photocatalysts. Meanwhile, the big data, together with artificial intelligence (AI) and machine learning (ML), will be a future key research area to improve the reproducibility of the catalyst synthesis, to facilitate catalyst screening, to enhance the experimental safety and to save the experimental costs and time.

Overall, this research field is at an infant state and all these areas should be enhanced in order to improve the NH₃ synthesis rate with high reproducibility and reliability of the procedure.

Conflicts of interest

There are no conflicts to declare.

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