



Cite this: *RSC Adv.*, 2025, **15**, 48775

Research progress in photochemical synthesis of urea through C–N coupling reactions

Zhidong Yang * and Peixia Li

Urea, as an important organic compound, plays significant role in promoting the development of agriculture, industry and biological sciences. Conventional urea synthesis process requires harsh reaction conditions (high temperature and pressure), making it energy-intensive, emission-intensive, and costly. Photocatalytic C–N coupling is a potentially green and promising alternative strategy for synthesizing valuable urea from CO_2 and inexpensive nitrogen sources (such as N_2 , nitrates, ammonia) as feedstocks under ambient conditions using solar energy. However, the specific details of urea photosynthesis have not been systematically reviewed so far. This article reviews the basic principles of photocatalytic urea synthesis, including the fundamentals, key intermediates, product identification and quantification. Meanwhile, it comprehensively summarizes research advances in photocatalytic urea synthesis, with a focus on the application and design principles of photocatalysts in urea production. Finally, the major challenges and prospective research directions in the field of photocatalytic urea synthesis are thoroughly discussed. We hope that this review will provide useful insights to inspire future research and discoveries.

Received 24th October 2025
 Accepted 3rd December 2025

DOI: 10.1039/d5ra08170j
rsc.li/rsc-advances

1 Introduction

Urea is a compound with critical applications across multiple fields.^{1,2} In agriculture, it serves as an efficient nitrogen fertilizer that improves soil quality, promotes plant growth, and enhances crop yield and quality.^{3,4} In industry, it is an important raw material for fine chemicals such as plastics, resins, coatings, adhesives, pigments, and dyes.^{5,6} In the energy sector, urea can be used as a proton conductor in fuel cells to improve performance, and it can also produce hydrogen through pyrolysis or electrolysis, serving as a potential hydrogen source.⁵ In medicine, urea exhibits functions such as keratin softening, wound healing, antipruritic, and antibacterial effects, making it widely used in dermatological treatments.^{7,8} Therefore, the development of the urea synthesis industry is of great significance for ensuring global food security, advancing chemical technology, promoting economic growth, and improving human living standards.⁹

The industrial urea synthesis process involves the reaction of ammonia and carbon dioxide under harsh conditions to produce ammonium carbamate, which is then dehydrated to produce urea.^{10–13} As one of the reaction feedstocks for urea synthesis, approximately 80% of the ammonia is derived from the Haber–Bosch process.¹⁴ The Haber–Bosch process requires high-temperature and high-pressure conditions (400–650 °C and 200–400 bar), accounting for 1–2% of global energy

consumption.^{14–16} Hydrogen, as another key reactant in the Haber–Bosch process, is typically produced *via* steam methane reforming ($\text{CH}_4 + 2\text{H}_2\text{O} \rightarrow 4\text{H}_2 + \text{CO}_2$) or coal steam reaction ($\text{C} + 2\text{H}_2\text{O} \rightarrow 2\text{H}_2 + \text{CO}_2$), both of which generate significant CO_2 emissions.¹⁷ It is estimated that the Haber–Bosch process contributes to about 1.5% of global anthropogenic CO_2 emissions.¹⁷ As a result, conventional urea synthesis not only consumes substantial energy but also exacerbates environmental burdens. It is urgent to explore more environmentally friendly and energy-saving urea production processes and technologies to reduce environmental pollution and resource consumption.¹⁸

Fortunately, solar-driven photosynthesis techniques are able to initiate chemical reactions under mild conditions, thereby reducing energy consumption and environmental issue and offering prospects for sustainable development and green chemistry.¹⁹ Photocatalytic C–N coupling reactions can convert carbon species such as CO_2 , CO , methanol and nitrogen species such as N_2 , nitrite, nitrate, or NO into valuable urea under ambient conditions and in aqueous solutions.^{20–23} Photosynthesis of urea not only makes use of abundant and free solar energy to alleviate the excessive use of fossil fuels and save energy, but also facilitates efficient and economical CO_2 fixation, which is of great significance to solve the problems of resource scarcity, environmental issues, global warming. In particular, directly coupling N_2 and CO_2 to synthesize urea by photocatalysis have attracted much attention in recent years.^{24–26} In the process of photocatalytic urea synthesis, excited semiconductors first generate electrons and holes that

School of Chemistry and Environment, Ankang University, Ankang 725000, Shanxi, China. E-mail: yangzd1618@163.com



participate in reduction–oxidation reactions to form nitrogen and carbon-containing intermediates. This is followed by the critical step of C–N coupling, and finally, electron-driven hydrogenation through proton-coupled reduction ultimately yields urea. Photocatalytic synthesis of urea is a prospective and innovative green chemical method, which opens up a sustainable pathway for urea synthesis.

Although photocatalytic synthesis of urea has been intensively investigated and made some progress in recent years, reported photocatalytic urea synthesis rates remain unsatisfactory. Currently, there are several challenges for the photosynthesis of valuable urea, such as high thermodynamic stability of reaction substrates (e.g., nitrogen and carbon dioxide),^{27–30} undesirable adsorption capacity of substrates/intermediates/products on the surface of the photocatalysts,³¹ high charge-carrier recombination efficiencies,³² slow kinetic processes of multi-electron transfers,^{33,34} and competing reactions in parallel products.^{35,36} These factors collectively result in low solar-to-chemical energy conversion efficiency, severely hindering the development of photocatalytic urea synthesis. In order to improve the photosynthesis performance of urea, a comprehensive understanding is essential to enable rational design of photocatalysts and systematic optimization of the photochemical reaction system. Herein, we present a systematic review of research advances in photocatalytic urea synthesis. The fundamental principles of the photocatalytic urea formation process are first introduced, covering the detection methods of key intermediates, as well as experimental identification and quantification techniques for urea and other parallel products. Subsequently, the performance of both classical and emerging photocatalysts across various reaction systems is summarized, with emphasis on their structure–activity relationships. Finally, current challenges in this field are discussed, and future research directions are outlined (Fig. 1). It is hoped that this review could provide an overview of the current research achievements related to the photocatalytic urea synthesis, thus stimulating the research on the design and synthesis of photocatalysts and the development of more efficient and stable photocatalytic systems.

2 The basic principle details for photocatalytic urea synthesis

2.1. Fundamentals of photocatalytic urea synthesis

Photocatalytic urea synthesis is a green synthetic method that utilizes renewable solar power to convert nitrogen-containing small molecules (e.g., N_2 , NO_3^- , or NH_3) and carbon sources (e.g., CO_2 , CO or CH_3OH) into urea.^{37–40} Its basic principles involve multiple complex steps, mainly including: (1) the absorption of light energy by the photocatalyst, (2) the generation and separation of photogenerated carriers, (3) the adsorption and activation of reactants, and (4) the formation of the urea by C–N coupling reactions.

The main stages in photocatalytic urea synthesis are shown in Fig. 2. When a photocatalyst is exposed to light with an energy greater than its bandgap width, electrons in the valence

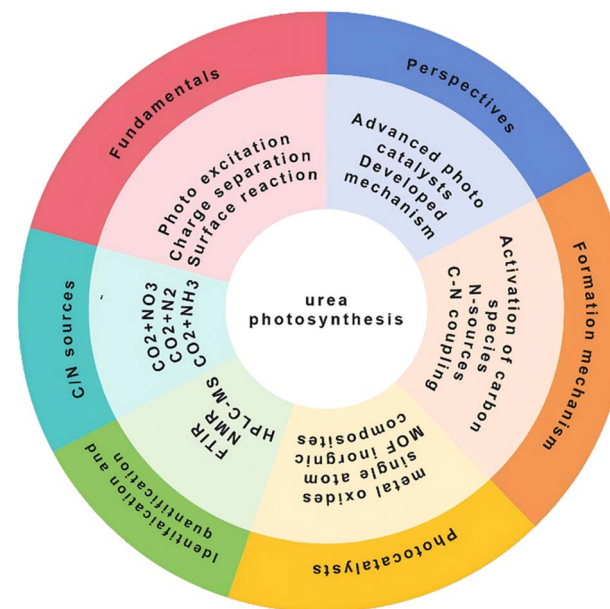


Fig. 1 Core framework of this perspective for urea photosynthesis.

band are excited and transition to the conduction band, forming photogenerated electron (e^-)-hole (h^+) pairs. Then, the separation and transfer of photogenerated electrons and holes. There are several situations regarding the separation and transfer of photogenerated carriers. One is that the photoexcited electrons and holes are transferred to the reduction and oxidation sites on the semiconductor surface, where they undergo reduction and oxidation reactions with the adsorbed substrate molecules respectively, as in process ⑤, ⑥; another is that the excited state carriers undergo recombination during the process of separation and transfer, that is, bulk phase recombination, as in process ③; the third is the electron–hole recombination that occurs on the semiconductor surface, namely surface recombination, such as process ④. It is worth noting that the recombination of carriers severely hinders the solar energy conversion efficiency. Therefore, neither bulk

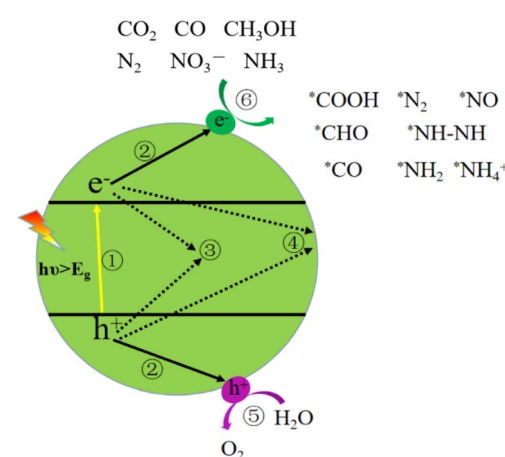


Fig. 2 Schematic diagram of photocatalytic urea production.



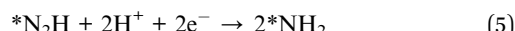
phase nor surface recombination is conducive to the progress of photocatalytic reactions. Furthermore, adsorption and activation of reactants: carbon species and nitrogen sources are adsorbed on the surface of the photocatalyst and activated by photogenerated carriers. Finally, the activated intermediates undergo a C–N coupling reaction to form urea precursors which then proceed a series of hydrogenation and electron transfer processes to ultimately produce urea.

2.1.1. Activation of carbon species. In the photocatalytic urea synthesis reaction, the reduction of CO_2 is a critical step in forming the carbon source of urea. It is generally believed that CO_2 is to form $^*\text{CO}$ intermediate, which then reacts with nitrogen-containing intermediates to generate urea. The specific analysis is as follows: the adsorbed CO_2 molecule couple with one proton and one electron to form the adsorbed $^*\text{HCOO}$ (eqn (1)) which then further are reduced into $^*\text{CO}$ intermediate by the proton-assisted single electron transfer process (eqn (2)). The generated $^*\text{CO}$ intermediates couple with nitrogen-containing intermediates (such as activated N_2 , $^*\text{NH}_2$, $^*\text{NO}$ etc.) to form a C–N bond, ultimately generating urea. For CH_3OH as the carbon source system, more $\cdot\text{CH}_2\text{OH}$ radicals are generated from CH_3OH , which are subsequently oxidized to $^*\text{CHO}$ intermediates on Pt cluster/ TiO_2 .⁴⁰

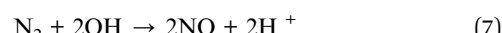
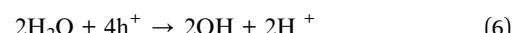


2.1.2. Activation of N-containing sources. The N_2 molecule is the most stable known diatomic molecule, possessing a very short $\text{N}\equiv\text{N}$ bond length of only 109.8 pm with a bond strength of 941 kJ mol⁻¹.⁴¹ A large energy gap (10.82 eV) exists between the highest occupied molecular orbital (HOMO) and lowest unoccupied molecular orbital (LUMO) of N_2 , whilst the molecule has both a high ionization potential (15.85 eV) and a low electron affinity (-1.9 eV) making oxidation or reduction difficult.⁴² The efficient activation of N_2 is a key step in urea synthesis. N_2 activation proceeds *via* two principal pathways: reduction and oxidation-reduction. In reduction systems, such as Cu SA- TiO_2 (ref. 43), Pt cluster/ TiO_2 (ref. 40) and Pd- CeO_2 (ref. 44), N_2 is adsorbed onto metallic sites like Cu, Pt and Pd through “ σ - π^* ” interactions and subsequently hydrogenated to yield $^*\text{N}_2\text{H}$, $^*\text{N}_2\text{H}_2$, $^*\text{NH}_2$ (eqn (3)–(5)). Reductive sites in photocatalysts like $\text{Ti}^{3+}\text{-TiO}_2$, CeO_{2-x} and $\text{Ru-O}_4\text{Ti}_1$ facilitate multi-electron transfer to directly crack the $\text{N}\equiv\text{N}$ bond. For the path of oxidation-reduction, N_2 is first oxidized to nitrogen oxides, which are then coupled with $^*\text{CO}$ and gradually hydrogenated for reduction (eqn (6)–(8)). System like $\text{Ni}_1\text{CdS}/\text{WO}_3$,⁴⁵ the N_2 was converted into NO species by $^*\text{OH}$ radicals generated from photogenerated holes over the WO_3 component, meanwhile, the CO_2 was transformed into $^*\text{CO}$ species over the Ni site by photogenerated electrons. The generated NO and $^*\text{CO}$ species were further coupled to form $^*\text{OCNO}$ intermediate, then gradually transformed into urea. Other systems, such as nitrate and nitrite, mainly underwent hydrogenation reaction (eqn (9)–(15)).

N_2 reduction:



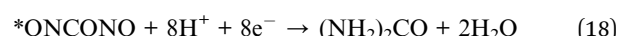
N_2 oxidation-reduction:

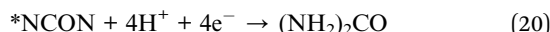


NO_3^- reduction:



2.1.3. C–N coupling. Typically, the urea synthesis from CO_2 and nitrogen-containing species is regarded as a co-reduction process. In general, C–N proceeds *via* three principal pathways. (1) $^*\text{NH}_2$ and $^*\text{CO}$ couple to form the $^*\text{NH}_2\text{CO}$ (eqn (16)), which is followed by coupling with excessive $^*\text{NH}_2$ to produce urea. (2) $^*\text{NO}$ couples with $^*\text{CO}$ forming $^*\text{ONCONO}$ (eqn (17)). Subsequent stepwise hydrogenation of these intermediates yields urea (eqn (18)). (3) Activated $^*\text{N}\equiv\text{N}$ couples with $^*\text{CO}$ to form $^*\text{NCON}$ (eqn (19)), which then undergoes hydrogenation (eqn (20)). Additionally, system like Pt Cluster/ TiO_2 ,⁴⁰ the crucial step of C–N coupling was initiated by the reaction between $^*\text{NH}\text{-NH}$ and $^*\text{CHO}$ intermediate. It was observed that Pt clusters undergo “ σ - π^* ” acceptor-donor interactions with N_2 , resulting in a reduction of the N_2 activation barrier and subsequent activation into $^*\text{NH}\text{-NH}$ intermediates. Furthermore, under the promotion of N_2 molecules and Pt clusters, more $\cdot\text{CH}_2\text{OH}$ radicals were generated from CH_3OH , which were subsequently oxidized to $^*\text{CHO}$ intermediates. Further analysis with DFT calculation demonstrated that $^*\text{NH}\text{-NH}$ and $^*\text{CHO}$ were important precursors in C–N coupling reactions.





2.2. Identification of key intermediates

Various intermediates are formed on the surface of the photocatalyst during activation of CO_2 and nitrogen source, such as *CO_2 , *COOH , *CO , *N_2 , *N_2H_2 , *NH_2 , *NH , *NO_2 , *NO_2OH , *NOOH , *NO , etc. Among them, the determination of C–N coupling intermediates such as *NH_2CO , *OCNO and *NCONO are particularly significant, which ultimately leads to urea formation. Therefore, the determination of key intermediates in the photocatalytic synthesis of urea is of great scientific significance for revealing the reaction mechanism and understanding the catalytic process.

Combining *in situ* characterization techniques with theoretical calculations can capture key intermediates, reveal active sites and reaction pathways, thereby deeply understanding the reaction mechanism and optimizing the catalytic system. *In situ* FTIR (DRIFTS) is an important technique for studying key intermediates in the photocatalytic synthesis of urea. It can monitor the dynamic evolution of adsorbed species on the catalyst surface in real time and reveal the reaction path. For example, Sheng *et al.* used *in situ* diffuse reflection infrared Fourier transform spectroscopy (*in situ* DRIFT) to monitor the photosynthesis process of urea. Upon illumination, the emergence and growth of several intermediate bands were observed in both cases (Fig. 3a). The two vibrational bands located at 1332 and 1173 cm^{-1} were attributed to the wagging and deformation modes of NH_2 species, respectively, suggesting the one electron oxidation of NH_3 . For C–N intermediates, the 1439 cm^{-1} band corresponded to the C–N stretching vibration, closely resembling the standard band for urea, while the vibrational band located at 2205 cm^{-1} was assigned to the stretching mode of $O=C-NH_2$, derived from the coupling of NH_2 with CO .⁴⁶ Zheng *et al.* utilized *in situ* DRIFT spectroscopy to track the evolution of the critical intermediates in reactions.⁴⁵ As illustrated in Fig. 3b, an obvious broadening band at about 1594 cm^{-1} was observed, which could be attributed to the co-existence of the stretching mode of adsorbed NO species at 1590 cm^{-1} and the adsorbed CO species at 1610 cm^{-1} . More importantly, they captured the *OCNO intermediates at 2120 cm^{-1} , which was generated by C–N coupling from *CO and *NO . Theoretical calculations can predict the stability of intermediates, energy barriers and the reaction mechanism, complementing experimental data. By calculating the adsorption configurations and energies of possible intermediates (such as *CO , *NH_2 , *CONH_2 , etc), thermodynamically feasible paths were screened out. For instance, Pang *et al.* performed DFT calculations to explore the photocatalytic C–N coupling mechanism for urea production (Fig. 3c).³⁷ Comparing the Gibbs free-energy (ΔG) profiles for CO_2 reduction, revealed that pure Fe_2O_3 faced challenges in generating *CO because of its inherently high CO_2 activation barrier (with a rate-determining step of 1.28

eV). In contrast, $CuL-Fe_2O_3$ exhibited a more moderate rate-determining step (0.73 eV), thereby facilitating the generation of *CO intermediates. In addition, the desorption barrier stabilized the *CO intermediates for subsequent C–N coupling reactions. Considering the molecular orbital matching between *CO and *N_2 , the formation of *N_2CO intermediates was a spontaneous exothermic reaction (Fig. 3d). Coupled with mass spectrometry (MS) and nuclear magnetic resonance spectroscopy (NMR), isotope labeling enables precise tracking of reaction pathways. Zhang *et al.* detected *NCONO intermediate (2100 cm^{-1}) on $Ru-O_4Ti_1$ sites using *In situ* ATR-FTIR spectroscopy coupled with ^{15}N -NMR (Fig. 3e–g).⁴⁷ DFT calculations revealed unique Ru nanostructure ($Ru-O_4Ti_1$), which effectively triggered the activation of inert N_2 molecules, facilitated the formation of crucial $^*NN(OH)$ intermediates (Fig. 3h), lowered the energy barrier of the potential determining step (Fig. 3i), and served as an “electronic pump” for electron migration from nitrogen to TiO_2 support during urea photosynthesis.

2.3. Product identification and quantification

Photocatalytic urea synthesis is accompanied with the generation of various by-products, including CO , CH_4 , C_2H_4 , $HCOOH$, CH_3OH , C_2H_5OH , H_2 , NH_3 , CH_3NH_2 , $C_2H_5NH_2$, CH_3CONH_2 , etc. Gas chromatography can be used to determine gas-phase products. For liquid-phase or soluble products, the identification and quantification could be achieved by nuclear magnetic resonance (NMR) spectroscopy, the diacetyl monoxime method or the urease decomposition method, high-performance liquid chromatography (HPLC) and high-performance liquid chromatography-mass spectrometry (LC-MS) spectroscopy.

For the typical identification and quantification of urea by 1H NMR spectroscopy, dimethyl sulfoxide- d_6 (DMSO- d_6) (99.8 atom % D with 0.03% [v/v] trimethylsilane) and disodium maleate aqueous solution as internal standards are added into reaction solution without any preliminary treatment. Subsequently, the mixed solution is transferred to a NMR tube and analyzed. However, the signal intensity of active hydrogen will be influenced by pH value and solution concentration.

Regarding the diacetyl monoxime method, typically, 5 g of diacetylmonoxime (DAMO) and 100 mg of thiosemicarbazide (TSC) are dissolved in distilled water to prepare the DAMO-TSC solution with 1000 mL in total volume. Subsequently, the preparation of acid-ferric solution is mainly to add 100 mL concentrated phosphoric acid and 300 mL concentrated sulfuric acid to 600 mL deionized water, and then dissolve 100 mg $FeCl_3$ in the above solution. For the quantification of urea, 1 mL of DAMO-TSC solution and 2 mL of acid-ferric solution were added to 1 mL of sample solution. After vigorous mixing, the mixed solution was heated to 100 °C and maintained at this temperature for 15 minutes. When the solution is cooled to 25 °C, UV-vis absorption spectra were collected at a wavelength of 525 nm. The concentration-absorbance curve is calibrated using a standard urea solution with different concentrations. This method requires strict control of reaction conditions (temperature, time), and its accuracy can be affected by nitrates, ammonia, etc.



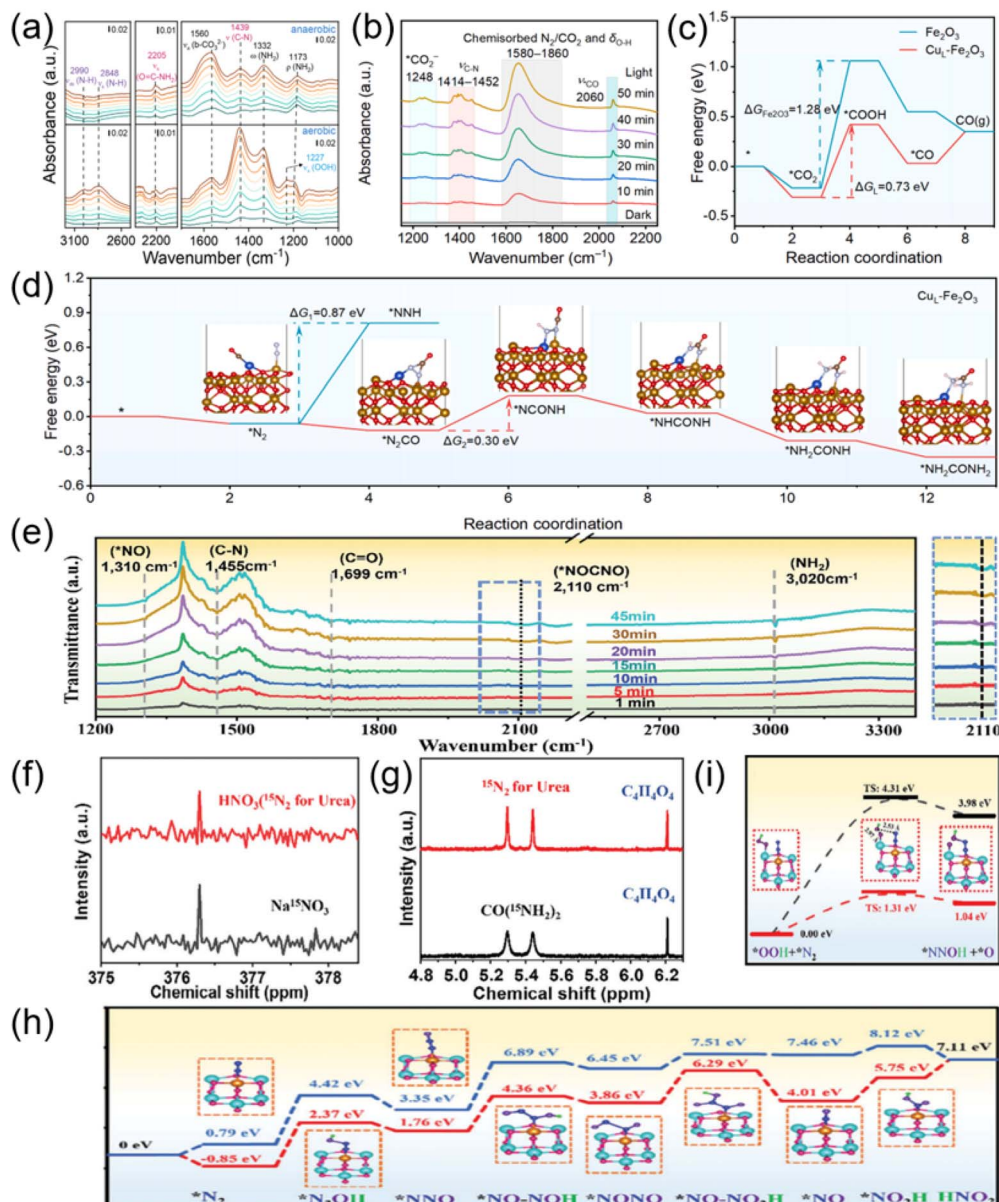


Fig. 3 (a) *In situ* FT-IR spectra of the photocatalytic urea synthesis under anaerobic (80% CO and 20% Ar) or aerobic (80% CO and 20% CO) condition on P25-4 h. Reproduced with permission from ref. 46. Copyright 2025, Wiley-VCH. (b) *In situ* DRIFT of $\text{Ni}_1\text{-CdS}/\text{WO}_3$ during photocatalytic urea synthesis. Reproduced with permission from ref. 45. Copyright 2024, Wiley-VCH. (c) Gibbs free-energy profiles of the CO_2 reduction process for Fe_2O_3 and $\text{CuL-Fe}_2\text{O}_3$. (d) Gibbs free-energy profiles and the geometric structure diagram of intermediates evolution during the urea synthesis process for $\text{CuL-Fe}_2\text{O}_3$ by alternating hydrogenation pathway. Reproduced with permission from ref. 37. Copyright 2025, Elsevier. (e) *In situ* ATR-FTIR spectroscopy measurements of Ru-TiO_2 over time during the coupling of N_2 and CO_2 . The enlargement at 2100 cm^{-1} is shown in blue on the right. (f) ^{15}N -NMR spectra of $^{15}\text{NO}_3^-$ (standard) and test product from NOR using $^{15}\text{N}_2$ as feeding gas. (g) ^1H -NMR spectra of $\text{CO} (^{15}\text{NH}_2)_2$ (standard) and test product from urea using $^{15}\text{N}_2$ as feeding gas. (h) Calculated free energy diagram for possible NOR mechanism on TiO_2 and Ru-TiO_2 (blue line: TiO_2 ; red line: Ru-TiO_2). (i) Reaction barrier diagrams from $^* \text{N}_2$ to $^* \text{NNOH}$ on TiO_2 and Ru-TiO_2 (black line: TiO_2 ; red line: Ru-TiO_2). Reproduced with permission from ref. 48. Copyright 2024, Wiley-VCH.

Additionally, the quantification of urea is achieved by HPLC (e.g. Agilent 1220 Infinity II) spectroscopy. HPLC was performed on a Luna 5 μm NH_2 column ($250\text{ mm} \times 4.6\text{ mm}$). The corresponding mobile phase, flow rate, and detected wavelength is needed. Measurement of urea in the reaction product: firstly, the product was rinsed repeatedly with ultrapure water to ensure that it was adequately collected, then pumped and

filtered and transferred to a volumetric flask and diluted to a volume of 50 mL. 5 mL of it was concentrated to 1 mL for the assay. The advantages of the HPLC method are high resolution, good repeatability, and less impurity interference. Compared with HPLC, LC-MS has higher sensitivity and is suitable for trace analysis, and reaction mechanism research in

photocatalytic synthesis of urea, but the instrument and maintenance costs are relatively high.

Overall, it is necessary to balance the detection target (urea concentration, matrix complexity), equipment availability and data requirements when choosing specific method. For mechanism research, NMR (isotope tracking) or LC-MS (product identification) are preferred. The HPLC and DAMO-TSC methods are recommended for routine testing in clinical/environmental applications. For trace detection, the HPLC and LC-MS methods offer ultra-high sensitivity for detecting urea.

3 The progress of urea photosynthesis

3.1. Urea synthesis from CO_2 and NO_3^-

In recent years, significant progress has been made in urea synthesis from inexpensive carbon and nitrogen sources. In particular, urea synthesis originates from CO_2 and NO_3^- from industrial emission or agricultural fertilizer.^{4,15,48}

Photochemical synthesis of urea was first reported by Yoneyama in 1998.⁴⁹ Inspired by the high activities for reduction of CO_2 to methanol and the favourable reaction rate for the reduction of NO_3^- to NH_4^+ of size-quantized TiO_2 semiconductor nanocrystals immobilized in polyvinylpyrrolidinone gel film (Q- TiO_2 /PVPD), the group explored the simultaneous reduction of CO_2 and NO_3^- (from LiNO_3) to form urea using Q- TiO_2 /PVPD in the propylene carbonate solution containing isopropanol as a hole scavenger. Apart from the target urea, the by-products formed included methanol, ammonia, hydrogen, Ti^{3+} and acetone. Except for acetone, which was the only observed oxidation product derived from the oxidation of isopropyl alcohol, the other five products are reduction products. The sum of the quantum efficiencies of the reduction products (15.4) was very close to that of the oxidation products (15.9). Besides, NH_2OH and NO were also used as nitrogen sources instead of NO_3^- , and HCOOH and CO gas were used instead of CO_2 as carbon sources. All cases tested in the study gave urea, but the reaction rate and distribution of products were largely influenced by the combination of carbon and nitrogen sources ($\text{CO}_2\text{-NO}_3^-$, $\text{CO}\text{-NO}_3^-$, $\text{HCOOH}\text{-NO}_3^-$, $\text{CO}_2\text{-NH}_2\text{OH}$, and $\text{CO}_2\text{-NO}$) used. It should be noted that the reduction of NO_3^- was hypothesized as the rate-determining step in the urea photosynthesis reaction, as evidenced by the significant increase in urea yield when the nitrogen source is switched to NO or NH_2OH , both of which are intermediates in the reduction reaction of NO_3^- .

Subsequently, they studied that photocatalytic reduction of CO_2 in the presence of nitrate ions using TiO_2 nanocrystals embedded in SiO_2 matrices.⁵⁰ It was found that the urea production was influenced by the kind of solvents, including ethylene glycol monoethyl ether, acetonitrile, sulfolane, and water. From the results, the highest polarity solvent, which was water, yielded the highest amounts of urea. It was explained that these results may be due to the different degrees of dissociation of LiNO_3 in solvents. With the increase of solvent polarity, the dissociation degree of LiNO_3 increased, and the amount of NO_3^- available for the reduction reaction on TiO_2 also increased.

Aligned with their previous report, the primary conclusion of this study was that the reduction of NO_3^- was the rate-determining step in the photocatalytic production of urea.

In 2005, the photosynthesis of urea from CO_2 and NO_3^- was reported by Shchukin *et al.* on TiO_2/Cu particles encapsulated inside poly(styrene sulfonate)/poly(allylamine hydrochloride) capsules of different size (2.2, 4.2, and 8.1 μm) in aqueous solution.⁵¹ Poly(vinylalcohol) was employed as electron donor to facilitate the photosynthetic process. Fig. 4a showed an assembly scheme for spherical microreactor. The highest yield of urea photosynthesis (1.7 mM) was achieved for Cu-modified TiO_2 nanoparticles encapsulated inside 2.2 μm poly(styrene sulfonate)/poly(allylamine hydrochloride) capsules (Fig. 4b). The decreasing the size of the confined microvolume of poly-electrolyte capsules accelerated the NO_3^- photoreduction, which was considered to be a limiting stage of the urea photosynthesis, and correspondingly improving the efficiency of urea production.

The same year, Ustinovich *et al.* reported that urea was synthesized by the photoinduced reduction of CO_2 in the presence of NO_3^- , using titania-stabilized perfluorodecalin-in-water (PFD: TiO_2) emulsions and isopropanol as the hole scavenging reagent.⁵² The production of urea was obtained by irradiation PFD: TiO_2 emulsion and TiO_2 suspension, evidenced that in the case of emulsion the efficiency of urea photoproduction dramatically increased (Fig. 4c). Cu-modified nano-dispersed TiO_2 emulsion also studied the simultaneous photoreduction of CO_2 and NO_3^- , the urea photoproduction rate reached its maximum at copper loading of *ca.* 3wt% and was higher than that of PFD: TiO_2 emulsion (Fig. 4c). The increase of efficiency and selectivity of urea photoproduction observed for titania-stabilized perfluorocarbon-in-water emulsion can be attributed to high concentration of CO_2 in the oleic phase contacting with photocatalyst and favourable conditions for stabilizing of the reaction intermediates to form C-N bonds in the case of two-phase reaction medium.

Despite the inspiring findings, early studies lacked discussion of the reaction mechanism. It was not until 2011 that Srinivas *et al.* first proposed a reaction pathway in his study on the photocatalytic synthesis of urea (Fig. 5a). Firstly, they investigated influence of hole scavengers (isopropanol and oxalic acid) on nitrate reduction for urea formation over TiO_2 catalyst.¹² The carbonaceous hole scavengers not only served as electron donors but also underwent further oxidation to produce CO_2 , thus serving as a carbon source, whereas the NO_3^- ion was used as the nitrogen source in their work (Fig. 5b). The results showed that the urea yield slightly increases under the isopropanol as the hole scavenger because the formation of ammonium oxalate, which was not desired under the oxalic acid as the hole scavenger, was avoided. To improve the adsorption of nitrate and to prevent the oxidation of ammonia, which was formed during nitrate reduction, the immobilization of TiO_2 on zeolite was envisaged. The high activity of the TiO_2 (10 wt%) supported over zeolite sample can be ascribed to the high dispersion of TiO_2 on the surface, strong adsorption of substrates and also because of lower recombination of electron-hole pairs generated.



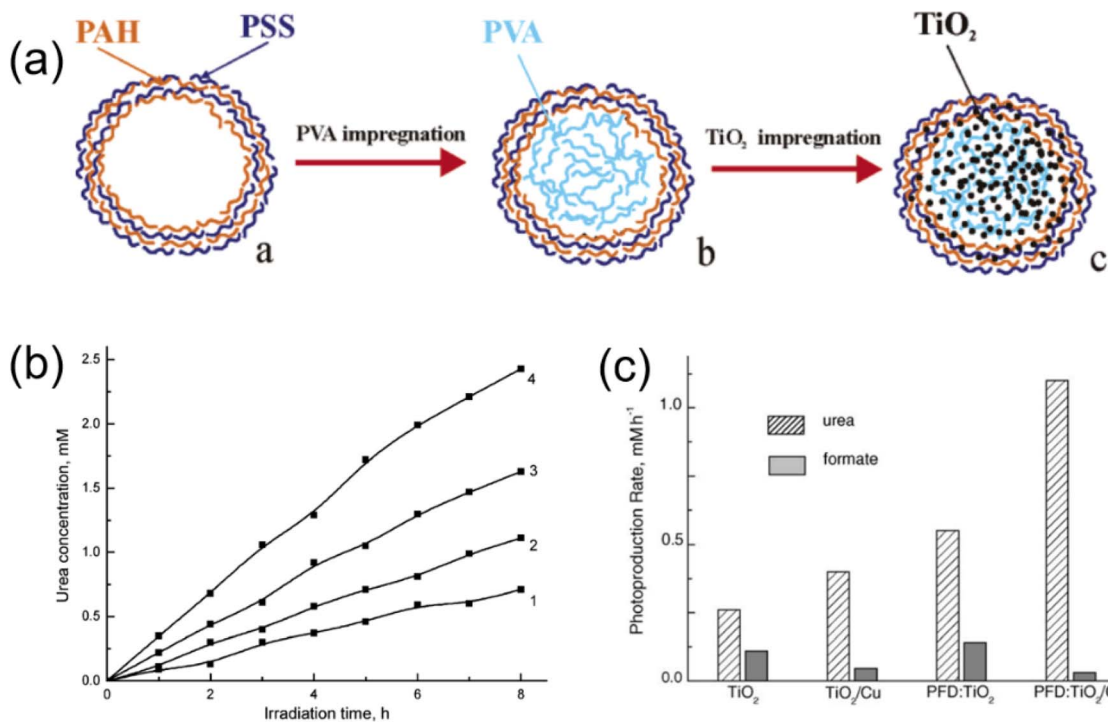


Fig. 4 (a) Schematic illustration of the assembly of a photocatalytic microreactor. (b) Time course of the photoproduction of urea in CO₂-saturated TiO₂/Cu suspension (1) and inside TiO₂/Cu-loaded polyelectrolyte capsules of 8.1 (2), 4.2 (3), and 2.2 μm (4) diameter. Reproduced with permission from ref. 51. Copyright 2005, American Chemical Society. (c) Initial rate of urea and formate photoproduction in CO₂-saturated TiO₂ suspension, suspension of Cu-modified TiO₂, PFD-in-water emulsion stabilized with TiO₂, and PFD-in-water emulsion stabilized with Cu-modified TiO₂. Aquatic phase initially contained 1 M NaNO₃ and 1 M isopropanol, pH 5.5. Reproduced with permission from ref. 37. Copyright 2005, Elsevier.

Huang *et al.* presented the novel Cs₂CuBr₄/TiO_x-Ar (CCBT-Ar) photocatalyst for urea generation through the simultaneous reduction of CO₂ and NO₃⁻.⁵³ The synthesis of the designed

CCBT-Ar catalysts was accomplished through a facile two-step method (as depicted in Fig. 6a). The X-ray diffraction (XRD) patterns showed that CCB was confined with amorphous TiO_x

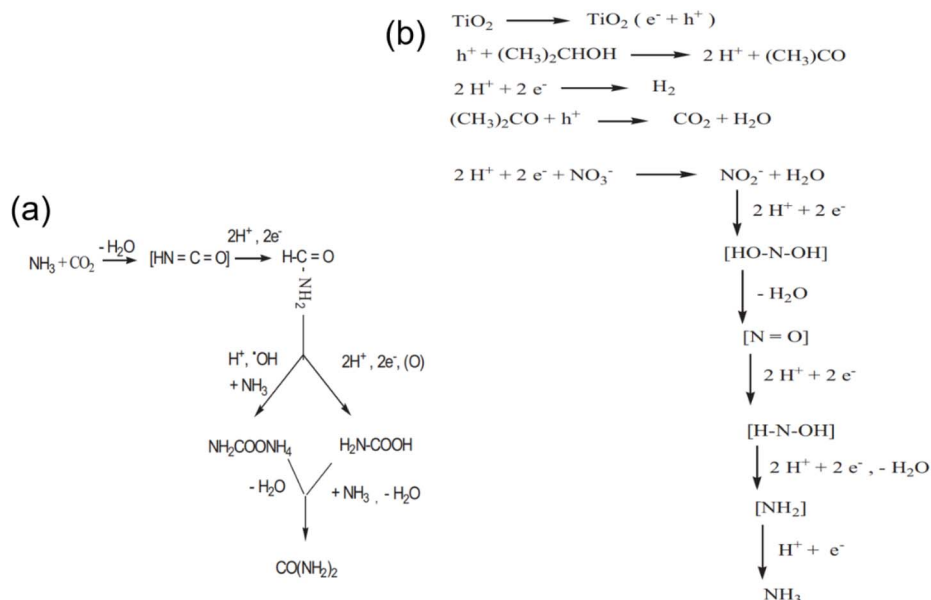


Fig. 5 (a) Speculative mechanism for urea formation. (b) Schematic representation and stabilization of intermediate species during nitrate reduction and prevention of ammonia oxidation over TiO₂-zeolite. Reproduced with permission from ref. 12. Copyright 2011, The American Society of Photobiology.



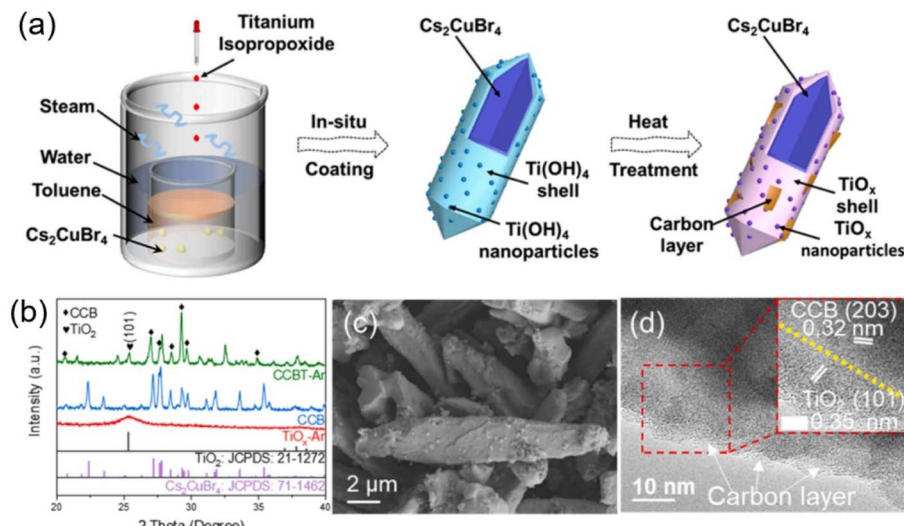


Fig. 6 (a) Schematic illustration of the synthesis procedure of CCBT-Ar/air catalysts. (b) XRD patterns of CCBT-Ar, CCB, and TiO_x-Ar. (c) SEM image of CCBT-Ar. (d) High-resolution TEM image of CCBT-Ar (with a 5 nm bar size inset). Reproduced with permission from ref. 53. Copyright 2025, Elsevier.

(Fig. 6b). The scanning electron microscopy (SEM) of the CCBT-Ar adopted a spindle-like shape (Fig. 6c). The research findings demonstrated that the introduction of oxygen vacancies (O_v) in TiO_x played a crucial role in reactant adsorption and the promotion of rate-determining steps in urea generation. However, the presence of O_v also led to significant carrier trapping effects, reducing the overall reaction activity. The well-designed CCBT-Ar catalyst exhibits outstanding photocatalytic urea generation activity, which attributes to its unique structural synergy. The TiO_x nanocrystals were interconnected by *in situ* grown carbon nanosheets (Fig. 6d), allowing for efficient electron extraction from TiO_x and acting as an electron reservoir, effectively suppressing O_v -induced carrier recombination. As electrons in TiO_x were consumed, those reserved in the carbon nanosheets were replenished, ensuring the sustainability of the reaction.

Li and coworkers engineered a TiO_2 nanoparticle modified Cu nanorod photocatalyst ($TiO_2@Cu$) for simultaneously promoting the NO_3^- reduction and CO_2 reduction reaction in the photocatalytic synthesis of urea (Fig. 7a).⁵⁴ The TiO_2 nanoparticles were uniformly covered onto the surface of the Cu nanorod *via* an impregnation-reduction method (Fig. 7b), and the well-integrated core-shell $TiO_2@Cu$ showed excellent efficiency in photocatalytic urea synthesis, reaching up to $72.8 \mu\text{mol g}^{-1} \text{ h}^{-1}$ of urea yield (Fig. 7c). The remarkable photoactivity was attributed to the unique $Ti-O-Cu$ bond in heterojunction interface of $TiO_2@Cu$ (Fig. 7d-f), and $Ti-O-Cu$ bond provided a favorable electron transfer pathway from TiO_2 to Cu, which accelerated the transfer of photogenerated charge and reduced the recombination of hole and electron. Meanwhile, the introduction of Cu altered the energy band structure of TiO_2 , resulting in a smaller band gap and further improving the utilization of light. The density functional theory calculation (Fig. 7g) indicated that the energy barrier of the C-N coupling

reaction in $Ti-O-Cu$ site of $TiO_2@Cu$ (-3.22 eV) was much lower than individual Cu site (1.21 eV).

It is hard to achieve the simultaneously catalytic reduction of two reactants CO_2 and nitrate at a single active site. Therefore, constructing bimetallic active sites is a feasible method to simultaneously catalyze the reduction of CO_2 and nitrate to C-N bonding.⁵⁵ However, due to the long distance between the coupling reaction intermediates generated at the bimetallic sites in the alloy, it is difficult to achieve efficient coupling between CO_2 and nitrate reduction intermediates. Hence, structuring bimetallic active site with short distance for the simultaneous catalytic co-reduction of CO_2 and nitrate is conducive to the C-N coupling between reduction intermediate of CO_2 and nitrate for urea synthesis. Zhao *et al.* engineered dual metal Cu and Ti active sites with a short distance of 2 \AA by single atom Cu anchored on TiO_2 toward photoelectrocatalytic urea synthesis from CO_2 and nitrate.³⁹ Cu and Ti dual active sites can efficiently catalyze the reduction of CO_2 to *CO and reduction of nitrate to *NH_2 intermediates, respectively. The relatively short distance of the Ti and Cu double site on SAC Cu- TiO_2 was conducive to the coupling of the two reaction intermediates *CO and *NH_2 to formation urea *via* C-N bonding by strong nucleophilic attack of *NH_2 . Density functional theory (DFT) calculations (Fig. 8) verified that compared with parallel competing reactions of *CO and *NH_2 such as *CO hydrogenation, *NH_2 hydrogenation and its dimerization, the coupling of *CO and *NH_2 had a lower energy barrier on dual metal active sites with short distance by single atom Cu anchored on TiO_2 .

The photoelectrochemical (PEC) system can be a viable solution for green synthesis of urea by combining the light absorbers and the catalysts into a fully integrated electrode. Thus, exploring and developing the PEC device is significant for green and efficient synthesis of urea in aqueous solution under



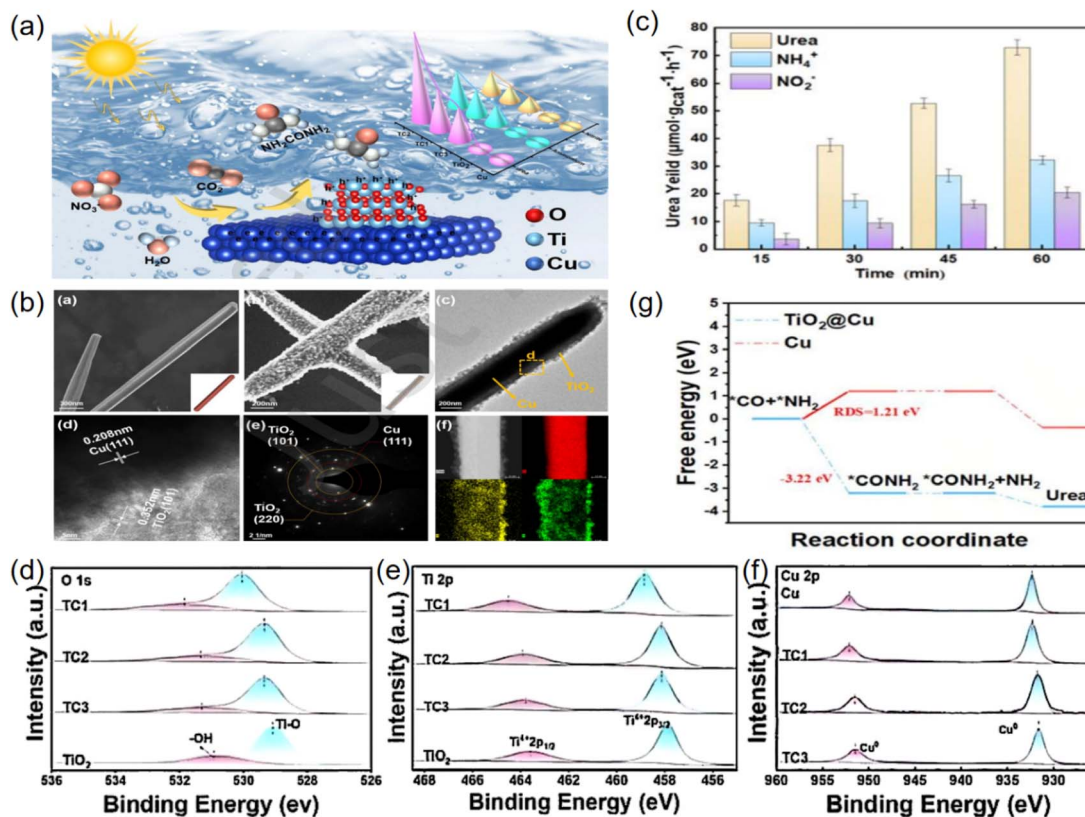


Fig. 7 (a) Schematic diagram of photocatalytic co-reduction of nitrate ions and carbon dioxide for urea synthesis over $\text{TiO}_2@\text{Cu}$ nanorod. (b) Structure characterization. (SEM image of Cu, the SEM image of TC2, TEM image of TC2, HRTEM image of TC2, SAED image of TC2; the elemental mapping of TC2). (c) The yield of urea over TC2. (d) O 1s (e) Ti 2p and (f) Cu 2p for TC1, TC2, TC3, and TiO_2 nanoparticle. (g) Free energy diagrams for the electrocatalytic C–N coupling of $^*\text{CO}$ and $^*\text{NH}_2$ on $\text{TiO}_2@\text{Cu}$ and Cu. Reproduced with permission from ref. 54. Copyright 2025, Tsinghua University Press.

mild conditions *via* solar energy. Wang *et al.* designed a hierarchical-structured Si-based photocathode to efficiently drive the solar-to-urea conversion by coupling the NO_3^- and CO_2 , in which nanostructured n^+p -Si serves as the light absorber decorated with TiO_2 layer and NiFe diatomic cocatalysts on N-doped carbon nanosheets (Fig. 9a).⁵⁶ The results demonstrated a remarkably high urea yield rate, faradaic efficiency (FE), and stable operation time of $81.1 \text{ mg h}^{-1} \text{ cm}^{-2}$, 24.2%, and 20 h at -1.0 V vs. RHE , respectively (Fig. 9b–d). The synergistic effect of NFDA, TiO_2 layer, and n^+p -Si enhances the charge-carrier dynamics of the photocathode. Duan *et al.* reported a photoelectrochemical method for urea synthesis by co-reduction of carbon dioxide and nitrate ion over a Cu_2O photocathode,⁶ delivering urea formation rate of $29.71 \pm 2.20 \mu\text{mol g}^{-1} \text{ h}^{-1}$ and faradaic efficiency (FE) of $12.90 \pm 1.15\%$ at low external potential ($0.017 \text{ V vs. Reversible hydrogen electrode}$) (Fig. 10a). The Cu_2O exhibited cubic morphology and exposes (100) facet (Fig. 10b and c), with suitable conduction band (CB) position (at about 1.1 V vs. SHE) to co-reduce CO_2 and NO_3^- (Fig. 10d–f). *In situ* Fourier transform infrared spectroscopy (Fig. 10h–j) and photo-assisted online differential electrochemical mass spectrometry (Fig. 10g) showed that CO^* and NO_2^* species were the key intermediates for the subsequent C–N coupling. Density functional theory (DFT) calculations

revealed that the first CN coupling during urea synthesis took place between CO^* and NO_2^* , which were the rate determining step, and the second C–N coupling occurred between CONH^* and NO_2^* . Zhao demonstrated an efficient GaN/Si photo-electrode for PEC urea synthesis from simultaneous NO_3^- and CO_2 reduction reactions under solar light.⁵⁷ The built-in potential in n^+p Si and the inherent catalytic activity of GaN nanowires (NWs) led to the selective synthesis of urea at a low overpotential.

3.2. Urea synthesis from CO_2 and N_2

The conversion of N_2 and CO_2 into urea through a photocatalytic C–N coupling reaction under ambient conditions presents a favorable approach. On the one hand, N_2 can serve as an abundant nitrogen source for urea photosynthesis, which accounts for 78% of the atmosphere. On the other hand, CO_2 , a main greenhouse gas leading to serious environmental concerns, can serve as a carbon source. Thus, the approach not only achieves effective energy conservation but also mitigates environmental concerns. Nevertheless, the photocatalytic urea production process still suffers from extremely serious challenges. Rationally designing photocatalysts that integrate the capture of inert gas molecules, bond cleavage and C–N coupling



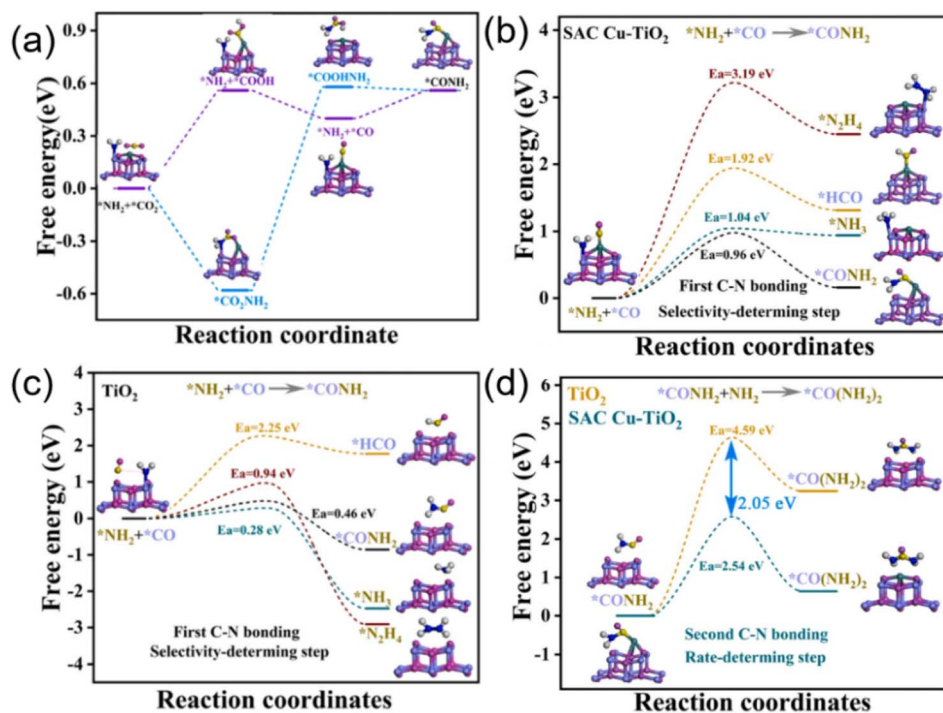


Fig. 8 (a) Free energy of formation $^*\text{CONH}_2$ from $^*\text{NH}_2$ and $^*\text{CO}_2$, diagram of free energy changes and activation barriers of $^*\text{CO}$ and $^*\text{NH}_2$ coupled to CONH_2 and other parallel $\text{CO}_2/\text{NO}_3^-$ reduction reaction on (b) SAC Cu-TiO₂, (c) TiO₂, (d) $^*\text{CO}(\text{NH}_2)_2$ formation from $^*\text{CONH}_2$ on TiO₂ and SAC Cu-TiO₂. Reproduced with permission from ref. 39. Copyright 2023, Elsevier.

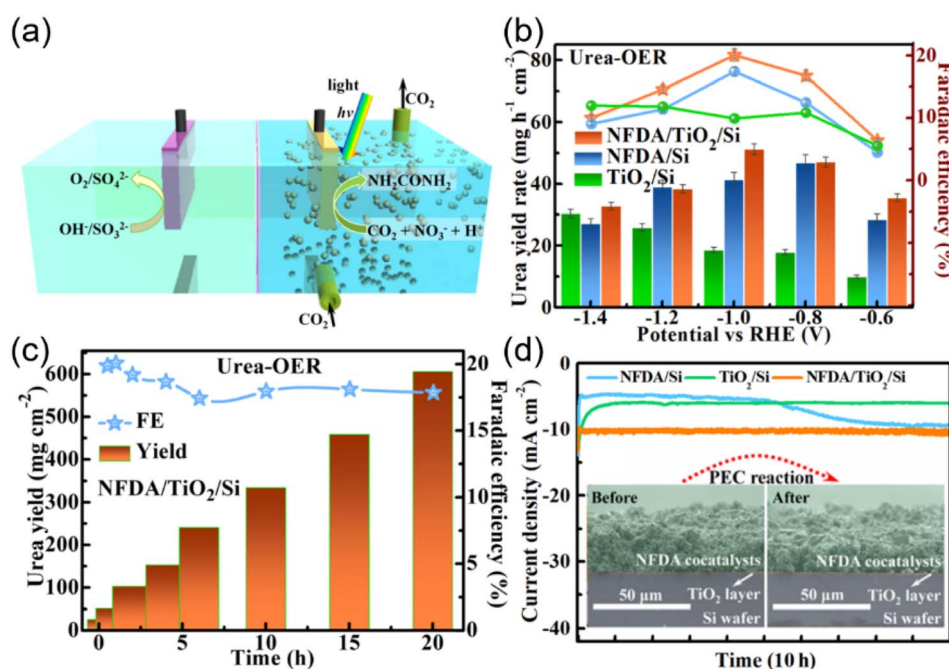


Fig. 9 PEC urea synthesis of the Si-based photocathodes in a 0.1 M KHCO_3 + 0.05 M KNO_3 electrolyte. (a) Diagrammatic sketch of PEC urea synthesis under 1 sun illumination. (b) Urea yield rate (column diagrams) and FE (point plots) of NFDA/TiO₂/Si (orange), NFDA/Si (blue), and TiO₂/Si (green) at various potentials for 0.5 h. (c) Time dependence of urea yield (column diagrams) and FE (point plots) acquired from NFDA/TiO₂/Si held at -1.0 V vs. RHE. (d) $J-t$ curves of NFDA/TiO₂/Si, NFDA/Si, and TiO₂/Si at -1.0 V vs. RHE for 10 h. The Insets are the cross-sectional FESEM images of NFDA/TiO₂/Si before and after 10-h PEC reactions. Reproduced with permission from ref. 56. Copyright 2024, PNAS.

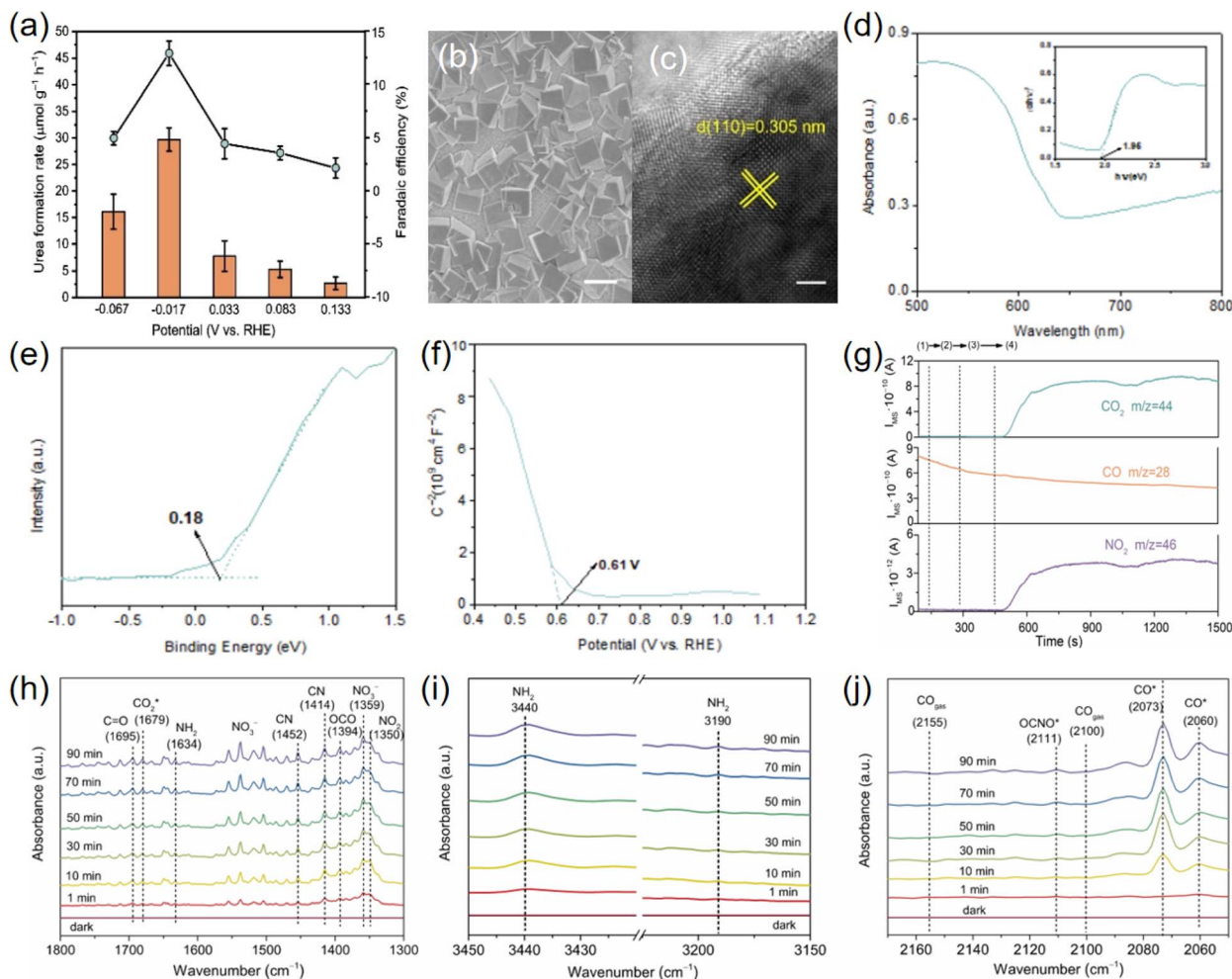


Fig. 10 (a) Urea formation rate and FE at different external potentials over Cu₂O. (b) SEM image of Cu₂O. (c) TEM image of Cu₂O. (d) Diffuse reflectance ultraviolet-visible spectrum and the corresponding $(\chi h\nu)^2$ versus photon energy plot of Cu₂O. (e) VB XPS spectrum of Cu₂O. (f) Mott–Schottky plots of Cu₂O. (g) Photo-assisted DEMS measurements over Cu₂O. Reaction conditions: (1) without external bias and CO₂ in darkness; (2) with cathodic bias but without CO₂ in darkness; (3) with cathodic bias but without CO₂ under light irradiation (AM 1.5 G irradiation); (4) with cathodic bias and CO₂ under light irradiation. *In situ* FTIR spectra in the range of (h) 1300–1800 cm^{−1}, (i) 3150–3450 cm^{−1} and (j) 2050–2170 cm^{−1}. Reproduced with permission from ref. 6. Copyright 2024, Wiley-VCH.

capabilities is a crucial prerequisite to promote the performance of urea photosynthesis.

In 2021, Maimaiti successfully established the photocatalytic synthesis of urea for the first time in the N₂/CO₂ system.⁵⁸ They utilized oxygen vacancy-rich TiO₂ loaded on carbon nanotubes with Fe cores (Ti³⁺-TiO₂/Fe-CNTs) as the catalyst and achieved the coreduction of N₂ and CO₂ into urea in water without the addition of a hole scavenger. The authors identified Ti³⁺ sites and the adjacent oxygen vacancy serves as the active center for N₂ and CO₂, respectively. Adsorbed N₂ and CO₂ are further activated by photogenerated electrons, forming six-membered cyclic intermediates (H₂NCONH₂)₂, which eventually evolved into urea (Fig. 11a).

It can be concluded from photocatalytic reactions involving multiple electrons (such as N₂ fixation and CO₂ reduction) that the efficient extraction of photogenerated electrons is an essential prerequisite for advanced photocatalysis. However, low availability of photogenerated electrons in intrinsic

photocatalysts seriously hinders its further application. Zhang *et al.* reported a photoinduced strategy based on a TiO₂ photocatalyst-immobilized reversible single-atom copper (denoted as Cu SA-TiO₂) for the photocatalytic synthesis of urea using N₂ and CO₂ molecules in the presence of pure H₂O (Fig. 11b).⁴³ The introduction of reversible single-atom Cu in the designed sample imparted additional electron-rich sites for the photoactivation of reactants (N₂, CO₂, and H₂O) and C–N coupling, together with the accelerated electron-transfer dynamics, ensuring the multi-electron supply for urea photosynthesis.

Theoretically, integrating N₂ reduction reaction semiconductors and CO₂ reduction reaction semiconductors to construct composite photocatalytic materials is expected to enable the conversion of CO₂ and N₂ into urea under photocatalytic conditions. For example, Wang *et al.* prepared 2D-CdS@3D-BiOBr S-scheme heterostructures by self-assembly of

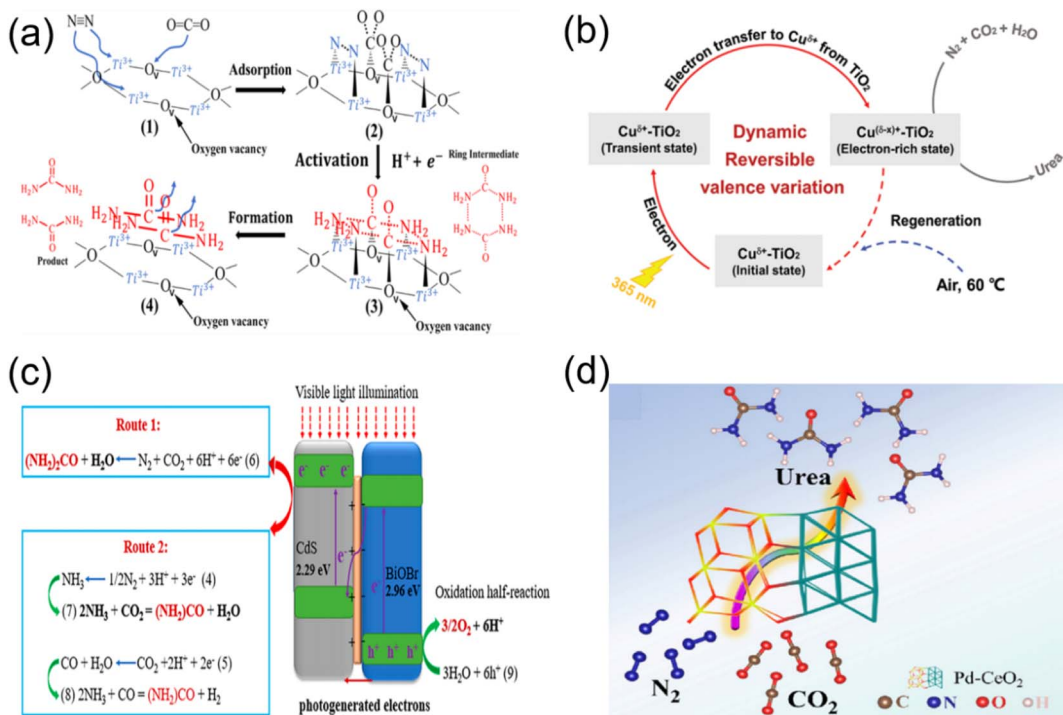


Fig. 11 (a) Mechanism of photocatalytic coreduction of N_2/CO_2 to form $\text{CO}(\text{NH}_2)_2$. Reproduced with permission from ref. 58. Copyright 2021, American Chemical Society. (b) Proposed mechanism of reversible and cooperative photocatalysis in Cu SA-TiO_2 . Reproduced with permission from ref. 43. Copyright 2022, Wiley-VCH. (c) Suggested mechanism for the photocatalytic synthesis of urea on 40% 2D-CdS@3D-BiOBr. Reproduced with permission from ref. 59. Copyright 2023, American Chemical Society. (d) Schematic diagram of the photocatalytic C–N coupling reaction in urea synthesis through the conversion of N_2 and CO_2 . Reproduced with permission from ref. 44. Copyright 2023, The Royal Society of Chemistry.

BiOBr microspheres (3D- BiOBr) and CdS nanosheets (2D- CdS). It was proposed that Cd^{2+} in CdS and oxygen vacancies in BiOBr of 2D- CdS @3D- BiOBr hybrids facilitate the adsorption and activation of N_2 and CO_2 , respectively, resulting in the formation of the $^*\text{HNCONH}$ intermediate.⁵⁹ Subsequently, their group synthesized the $\text{CdS}@\text{Bi}_2\text{WO}_6$ heterojunction for photocatalytic $\text{CO}_2\text{--N}_2\text{--H}_2\text{O}$ to urea with visible light. It was suggested that urea was synthesized on the surface of CdS through two mechanisms: (I) the reaction between the activated intermediates of CO_2 and N_2 (mainly) and (II) the reaction of the *in situ* formed NH_3 with the feed CO_2 (Fig. 11c).

The weak adsorption and activation ability of inert gases (CO_2 and N_2) on photocatalysts has been the main challenge hindering the development of this technology.^{60–62} Niu designed a Pd -decorated CeO_2 photocatalyst for the photoinduced coreduction of N_2 and CO_2 into urea,⁴⁴ enabling spontaneous electron transfer at the palladium–ceria interface (Fig. 11d). The investigations further endorsed that the emerged space-charge region in the $\text{CeO}_2(111)/\text{Pd}(111)$ interface not only effectively facilitates the targeted capture and activation of inert CO_2 and N_2 but also stabilizes the formation of key intermediates ($^*\text{NCON}$) (Fig. 12). Subsequently, they established mesoporous CeO_{2-x} nanorods with adjustable oxygen vacancy concentration by heat treatment in Ar/H_2 (90%:10%) atmosphere (Fig. 13a) and served as photocatalysts to convert both CO_2 and N_2 into urea under ambient conditions.⁶³ By introducing oxygen

vacancies to enhance the targeted adsorption and activation of N_2 and CO_2 (Fig. 13b), $\text{CeO}_2\text{-500}$ (CeO_2 nanorods heated treatment at 500 °C) revealed high photocatalytic activity toward the

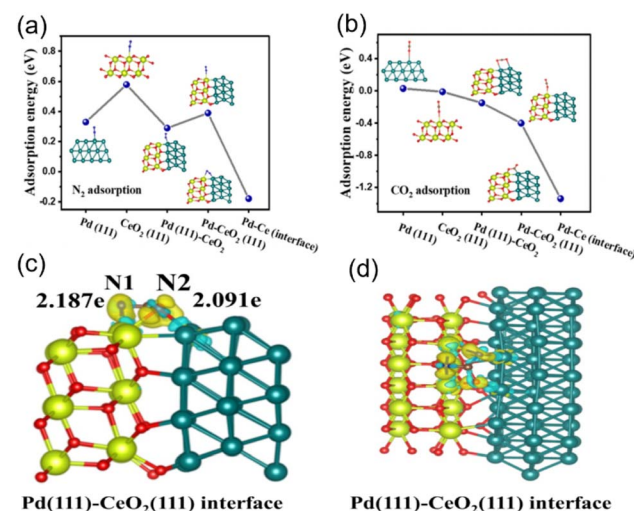


Fig. 12 (a) The adsorption energy of N_2 at different reaction sites. (b) The adsorption energy of CO_2 at different reaction sites. The differential charge density of $^*\text{NCON}$ at $\text{Pd}(111)\text{-CeO}_2(111)$ interface. (c) Side view, (d) top view. (The blue color represents electron consumption and the yellow color represents electron accumulation). Reproduced with permission from ref. 44. Copyright 2023, The Royal Society of Chemistry.



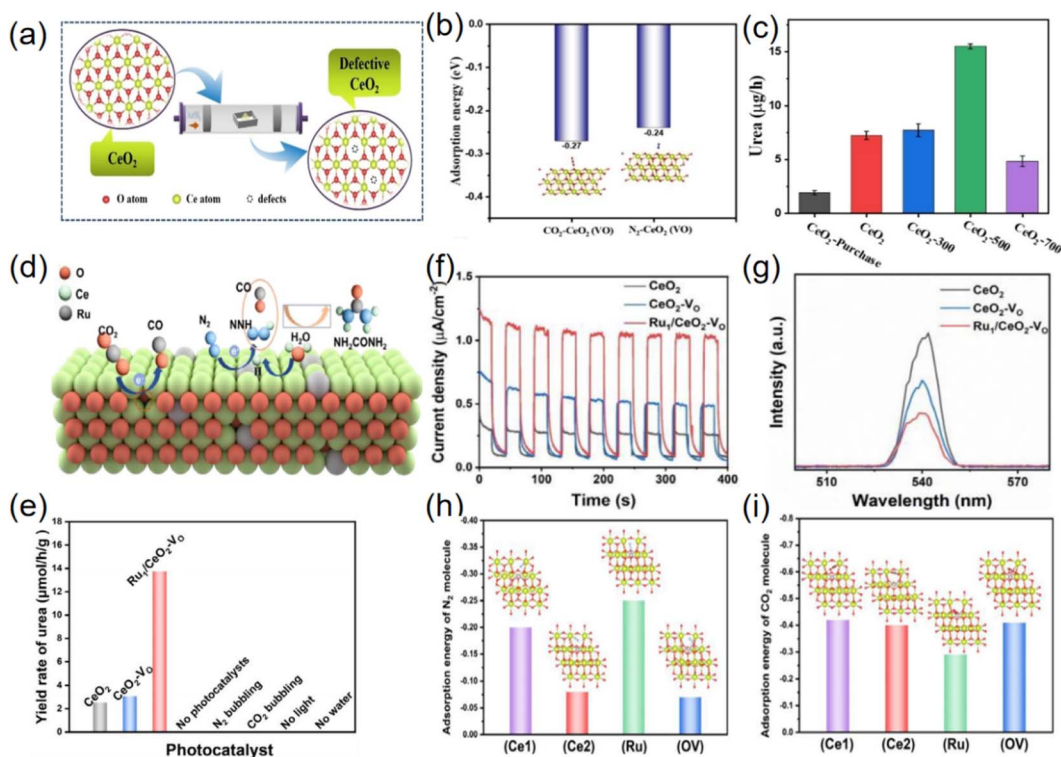


Fig. 13 (a) Schematic diagram of oxygen vacancy generation. (b) Adsorption energy of N_2 and CO_2 on CeO_2 with oxygen vacancy. (c) The urea formation rates of CeO_2 -purchase (commercial), CeO_2 , CeO_2 -300, CeO_2 -500 and CeO_2 -700 with all optical spectrum as light source. Reproduced with permission from ref. 63. Copyright 2023, Wiley-VCH. (d) Mechanism of the photocatalytic urea synthesis based on synergistic effects of the Ru_1/CeO_2 -VO. (e) The prepared materials and urea production efficiency under different conditions. (f) Photocurrent responses of CeO_2 , CeO_2 -VO, and Ru_1/CeO_2 -VO. (g) PL spectra of CeO_2 , CeO_2 -VO, and Ru_1/CeO_2 -VO. (h) N_2 adsorption energy of different sites. (i) CO_2 adsorption energy of different sites. Reproduced with permission from ref. 64. Copyright 2025, Wiley-VCH.

C–N coupling reaction for urea synthesis with a remarkable urea yield rate of $15.5 \mu\text{g h}^{-1}$ (Fig. 13c). In the most recent study, Meng synthesized a single-atom Ru and oxygen vacancies co-modified CeO_2 (Ru_1/CeO_2 -VO) by a photochemical strategy to achieve photocatalytic simultaneous reduction of CO_2 and N_2 for the synthesis of urea (Fig. 13d).⁶⁴ Benefiting from the synergistic effect of Ce^{3+} -VO and single atoms, Ru_1/CeO_2 -VO nanosheets delivered the highest urea yield rate of $13.73 \mu\text{mol g}^{-1} \text{h}^{-1}$ (Fig. 13e). Strong electron metal support interactions between the Ru single atoms and CeO_2 enabled effective separation of photoexcited carriers (Fig. 13f and g). Importantly, it was confirmed by experimental characterization and DFT calculations (Fig. 13h and i) that the vacancies can effectively adsorb CO_2 , and Ru single atoms can promote N_2 adsorption and activation and contribute to hydrolysis dissociation, supplying protons for hydrogenation of active species.

The use of metal elements as single active sites for the reduction of N_2 or CO_2 has been extensively studied and reported.^{65–68} However, urea synthesis requires the coupling reaction of the key intermediates $^*\text{CO}$ and $^*\text{N}$ to produce C–N bonds. Due to their high reactivity, these crucial reaction intermediates cannot exist in a stable form within the reaction system.^{42,69–71} Therefore, maintaining an optimal distance between the two reaction sites to enable the interaction of intermediates is an excellent strategy. Luo proposed a catalyst

design strategy with dual active sites to meet the needs of urea synthesis (Fig. 14a). A series of composites ($\text{SiW}_{12-X}\text{Mo}_X$ @MIL101(Cr), $X = 0, 3, 6, 9, 12$) were obtained and applied for photocatalytic urea synthesis from N_2 and CO_2 .⁷² The urea production rate of SiW_6Mo_6 @MIL-101(Cr) reaches $1148 \mu\text{g h}^{-1} \text{g}_{\text{cat}}^{-1}$ under the optimal experimental conditions (Fig. 14b). The performance from both experimental (Fig. 14c and d) and DFT calculation results (Table 1) indicated that the W site in SiW_6Mo_6 was assigned for the activation of nitrogen, whereas the Mo site was assigned for the activation of CO_2 . Zheng developed a multi-site photocatalyst, consisting of CeO_2 nanorods decorated with Ru nanoparticles and Cu single atoms ($\text{Ru-Cu}/\text{CeO}_2$), for the purpose of synthesizing urea at high yield.⁷³ The incorporation of Ru and Cu sites was crucial not only to generate high-density photogenerated electrons, but also to facilitate N_2 and CO_2 adsorption and conversion. The *in situ* formed local nitrogen-rich area at Ru sites increased the encounter possibility with the carbon-containing species generated from Cu sites, substantially promoting C–N coupling (Fig. 14e). Lu reported a Cu single-atom-decorated porous Fe_2O_3 nanorod catalyst with a Cu–O–Fe configuration for the direct artificial photocatalytic synthesis of urea from N_2 and CO_2 in pure water.³⁷ Because the d-orbitals of the Cu/Fe sites were close to the molecular orbitals of CO_2/N_2 , the Cu/Fe dual active sites

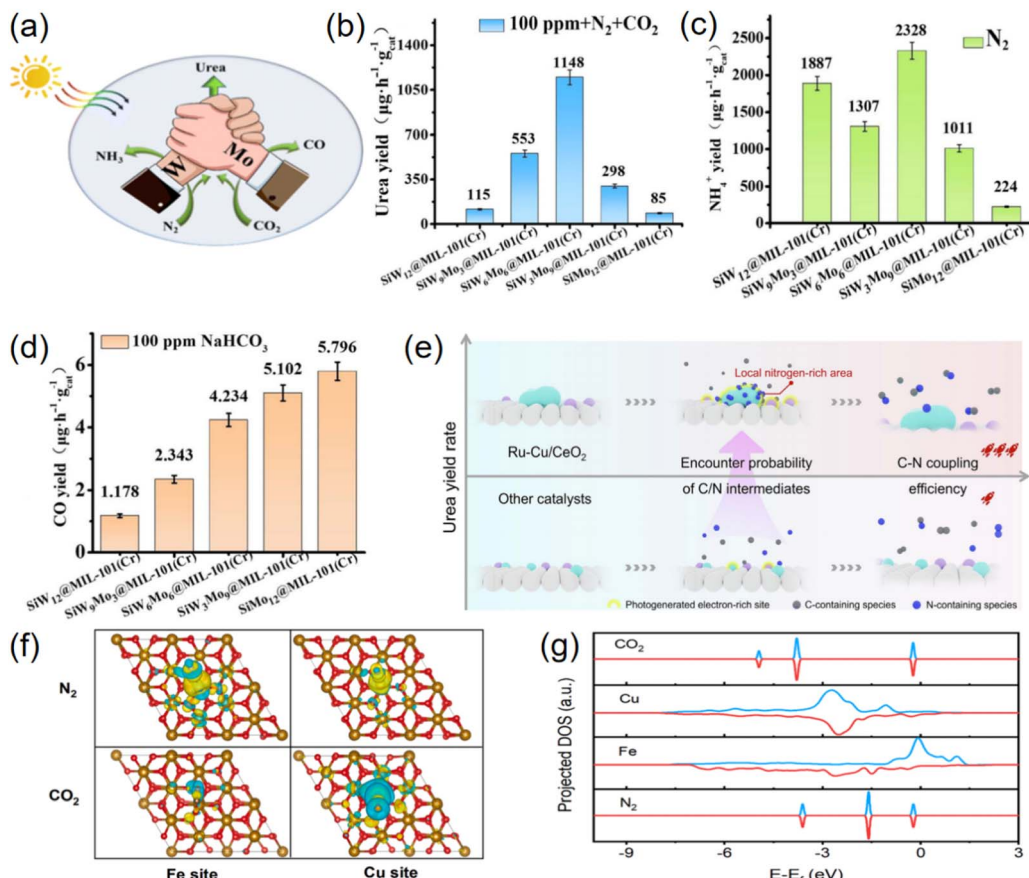


Fig. 14 (a) Schematic diagram of the design strategy for dual-active site catalysts in urea synthesis; (b) the urea production rate of composites in 100 ppm NaHCO₃; (c) the NH₄⁺ production rate of the composites in N₂; (d) the CO production rates of the composites in 100 ppm NaHCO₃. Reproduced with permission from ref. 72. Copyright 2024, The Royal Society of Chemistry. (e) The comparison between the designed Ru–Cu/CeO₂ catalysts (above) and the reported multi-site catalysts (below) during urea synthesis. Reproduced with permission from ref. 73. Copyright 2025, Elsevier. (f) Difference charge-density stereograms of adsorbed N₂ and CO₂ on the Cu and Fe sites, respectively. Yellow and cyan represent electron accumulation and depletion, respectively. (g) Calculated projected density of states of the d-orbital of Cu and Fe in CuL-Fe₂O₃, and CO₂/N₂ molecular orbitals. Reproduced with permission from ref. 37. Copyright 2025, Elsevier.

Table 1 Preparation of the catalysts under different conditions. Reproduced with permission from ref. 72. Copyright 2024, The Royal Society of Chemistry

Species	<i>E</i> _{total} (eV)	<i>E</i> _{mol-gas} (eV)	<i>E</i> _{sub-SiW₆Mo₆} (eV)	<i>E</i> _{ads} (eV)
Mo–CO ₂	471.519	−22.958	−448.034	−0.527
W–CO ₂	−471.303	−22.958	−448.034	−0.311
Mo–N ₂	−465.620	−17.096	−448.034	−0.490
W–N ₂	−465.968	−17.096	−448.034	−0.838

can selectively adsorb and activate N₂/CO₂, thereby facilitating efficient C–N coupling (Fig. 14f and g).

Photocatalysts with high reduction activity are necessary for overcoming the unfavorable energy barrier.^{42,74} It is well known that the Z-scheme photocatalyst can retain the high reduction capability of materials in the heterojunction.^{75–78} The high reduction activity favors boosting kinetics for co-reduction of N₂ and CO₂ and promoting C–N coupling reaction. Meanwhile, the Z-scheme photocatalyst has good charge separation ability, which can inhibit charge recombination and accelerate electron

transfer.^{79–81} In this regard, the Z-scheme photocatalyst could be more reasonable for improving photocatalytic urea synthesis by co-reducing CO₂ and N₂. SrTiO₃–FeS–CoWO₄ Z-scheme photocatalyst was designed to promote urea synthesis by N₂ and CO₂ co-reduction in water (Fig. 15a).⁸² Its fast electron transfer overcomes the bottleneck of sluggish kinetics for C–N coupling reaction, and dual active sites for the adsorption and activation of N₂ and CO₂ enhanced the kinetics for urea synthesis, resulting in enhanced urea yield of 8054.2 μg g_{cat}^{−1}·h^{−1} on SrTiO₃–FeS–CoWO₄ (Fig. 15b and c).

The disparity in physical properties, structure, and catalytic kinetics between N₂ and CO₂ poses stringent demands for the design of key catalysts, reaction control, mass transfer, and other aspects. Considering the unique advantages of nitrogen oxidation activation and the bilateral redox of photocatalytic reactions, Ding *et al.* synthesized a Ru–TiO₂ photocatalyst for urea photo-synthesis through a nitric acid-mediated pathway combining nitrogen oxidation and subsequent kinetically advantageous nitrate and CO₂ co-reduction (Fig. 15d).⁴⁸ The remarkable photo-activity was attributed to its unique oxygen



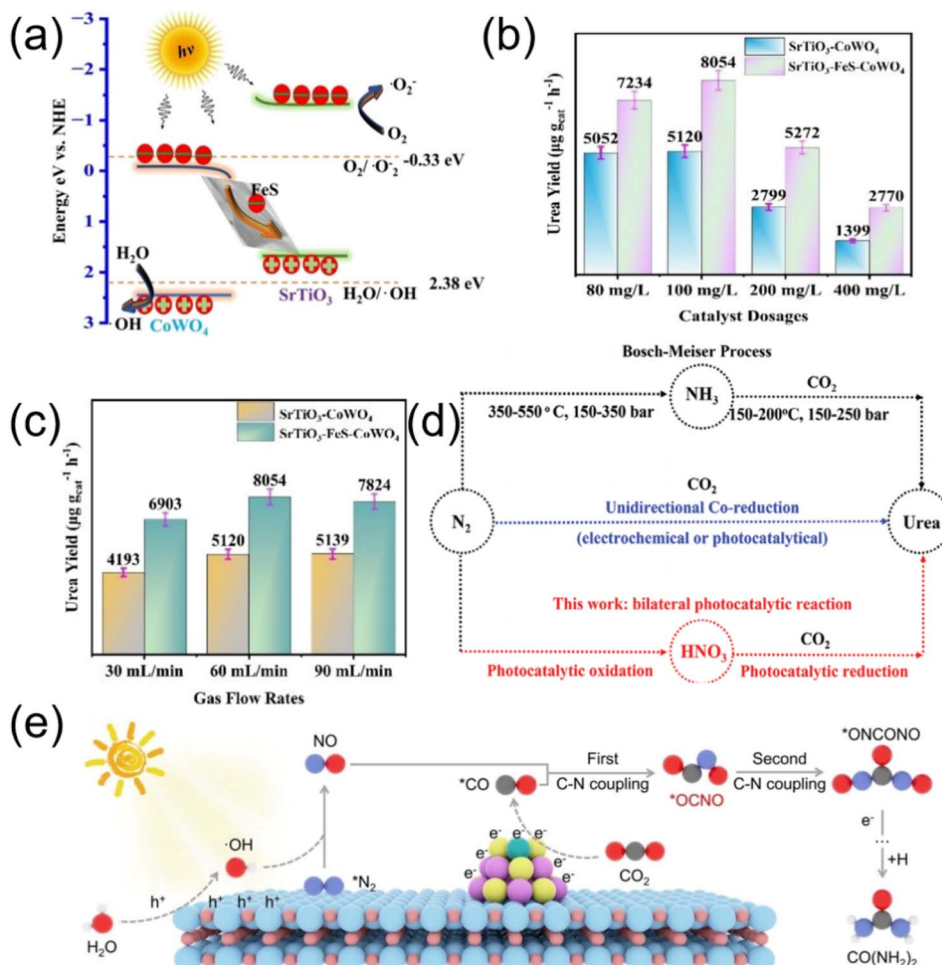


Fig. 15 (a) Energy band diagram of $\text{SrTiO}_3\text{-FeS-}\text{CoWO}_4$ Z-scheme heterojunctions. (b) Urea yields of $\text{SrTiO}_3\text{-FeS-}\text{CoWO}_4$ with different catalyst dosages; (c) urea yields of $\text{SrTiO}_3\text{-FeS-}\text{CoWO}_4$ with different gas-flow rates. Reproduced with permission from ref. 82. Copyright 2024, Wiley-VCH. (d) Comparison of the reaction conditions of different routes for urea synthesis. Reproduced with permission from ref. 48. Copyright 2024, Wiley-VCH. (e) Schematic illustration of photocatalytic urea synthesis mechanism over $\text{Ni}_1\text{-CdS}/\text{WO}_3$ catalyst. Reproduced with permission from ref. 45. Copyright 2024, Wiley-VCH.

vacancy-anchored Ru nanostructure ($\text{Ru-O}_4\text{Ti}_1$), which effectively activates inert N_2 molecules to minimize the disparity of orbital energy levels, facilitated the formation of crucial $^*\text{NN(OH)}$ intermediates, and served as an “electronic pump” to avoid electronegativity effect for facilitating electron transfer from nitrogen to TiO_2 support for urea photosynthesis.

Typically, the urea synthesis from N_2 and CO_2 is regarded as a co-reduction process. Current works mainly focused on the photogenerated electrons, the role of photogenerated holes has been commonly ignored. If photogenerated holes could be employed in activating reactant, it would not only mitigate the competition between N_2 and CO_2 in photogenerated electrons, but also improve the utilization efficiency of photogenerated carriers toward urea product, thereby boosting the performance of photocatalytic urea synthesis. Zheng *et al.* constructed a redox heterojunction consisting of WO_3 and Ni single-atom decorated CdS ($\text{Ni}_1\text{-CdS}/\text{WO}_3$),⁴⁵ enabling photogenerated electrons and holes to participate in the conversion of CO_2 and N_2 respectively (Fig. 15e). Specifically, the N_2 was activated by

photogenerated holes and CO_2 was converted by photogenerated electrons during photocatalytic urea synthesis, which mitigated the competition of photogenerated electrons between N_2 and CO_2 , enhanced the utilization efficiency of photogenerated carriers for urea products, and ultimately promoted the synthesis of urea.

3.3. Urea synthesis from CO_2 and NH_3

Compared with the stable $\text{N}\equiv\text{N}$ bond in N_2 , the lone pair electrons in NH_3 are naturally reactive centers, in favor of fast reaction dynamics.^{18,83} As a total result, photosynthesis of urea from NH_3 and CO_2 appears more promising for future practical industrial applications.⁸³⁻⁸⁷ Jiang *et al.* designed and synthesized a series of 3D N-heterocyclic covalent organic frameworks for urea photosynthesis from NH_3 and CO_2 .³⁸ Three isomeric three-dimensional (3D) COFs with two fold interpenetrated ffc topology (namely 3D-TPT-COF, 3D-PDDP-COF, and 3D-TBBD-COF) were functionalized by benzene, pyrazine, and tetrazine active cores, respectively, to modulate the catalytic micro-

environment through the change in the number of heterocyclic N atoms on the active cores (Fig. 16a). The geometrical and electronic structural advantages endowed 3D-TBBD-COF with superior photocatalytic activity towards urea production with the yield of $523 \mu\text{mol g}^{-1} \text{h}^{-1}$, 40 and 4 times higher than that for 3D-TPT-COF and 3D-PDDP-COF, respectively (Fig. 16b), indicating the heterocyclic N microenvironment dependent catalytic performance for these COFs photocatalysts.

Besides, Sheng *et al.* presented a photocatalytic pathway for the selective urea synthesis through the oxidative coupling between CO and NH_3 .⁴⁶ They uncovered that the O_2 concentration played a crucial role in controlling both the urea production rate (Fig. 16c) and its selectivity by effectively controlling the generation of oxidative species (Fig. 16d–g), such as photogenerated holes (h^+), superoxide radicals (O_2^\cdot) and hydroxyl radicals (OH^\cdot). Using oxygen-deficient TiO_2 under an air-level (20%) O_2 atmosphere, a urea generation rate of $54.31 \text{ mg g}_{\text{cat}}^{-1} \text{ h}^{-1}$ with 100% selectivity can be achieved (Fig. 16h). Mechanistic studies reveal that the process began

with the oxidation of NH_3 to NH_2^\cdot through oxidative radicals generated on TiO_2 , especially the oxygen-derived O_2^\cdot . This NH_2^\cdot radicals then coupled with CO to form urea.

3.4. Urea synthesis from other raw materials

The clean-energy-driven synthesis of urea from carbon- and nitrogen-containing small molecules has garnered significant interest but remained great challenges to achieve with high selectivity.

A Pt cluster-modified TiO_2 (Pt cluster/ TiO_2) catalyst was designed through the impregnation reduction method to facilitate the photocatalytic synthesis of urea by promoting the simultaneous N_2 reduction and CH_3OH oxidation reactions (Fig. 17a).⁴⁰ The prepared Pt cluster/ TiO_2 catalyst exhibited outstanding efficiency in urea synthesis, achieving a rate of $105.68 \mu\text{mol g}^{-1} \text{ h}^{-1}$ with N selectivity of $97.29 \pm 0.79\%$ (Fig. 17b). Further analysis with density functional theory (DFT) calculation revealed that the “ σ – π^* ” donor–acceptor interaction occurred between Pt clusters and N_2 (Fig. 17c–e), efficiently reducing the N_2 hydrogenation barrier. EPR experiments

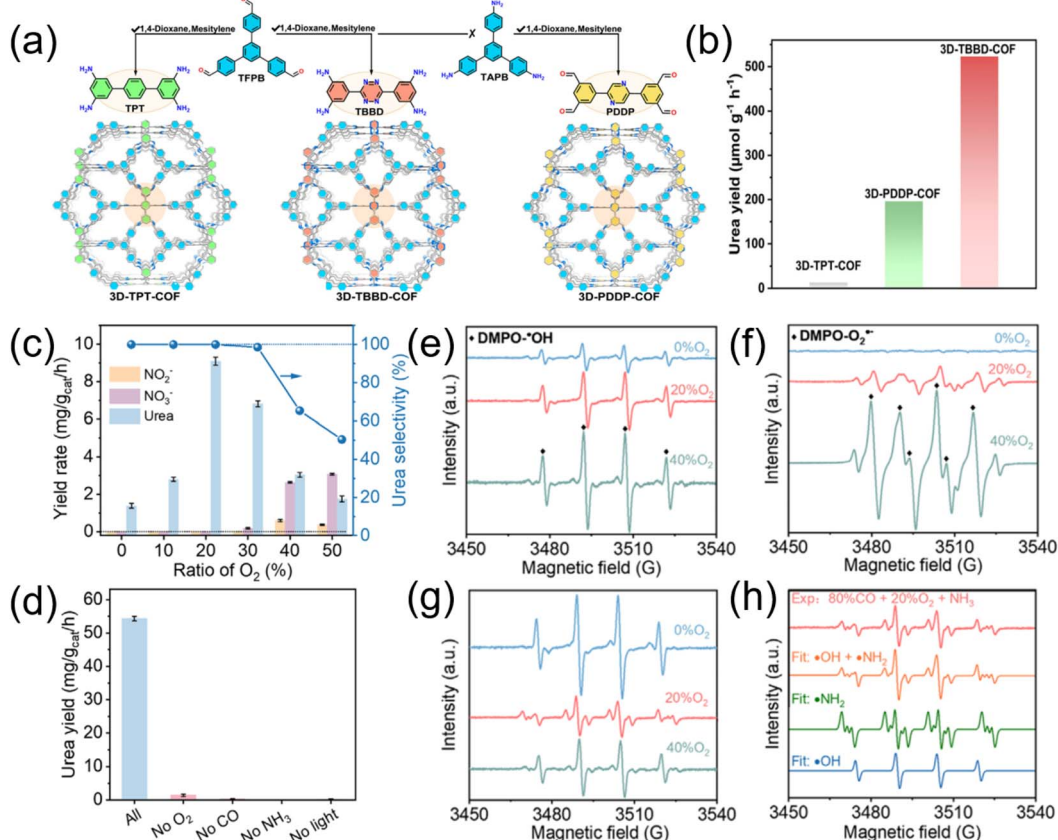


Fig. 16 (a) Schematic synthesis of 3D-TPT-COF (green), 3D-TBBD-COF (red) and 3D-PDDP-COF (yellow); (b) urea formation rate using 3DTPT-COF, 3D-PDDP-COF, and 3D-TBBD-COF as photocatalyst. Reproduced with permission from ref. 38. Copyright 2025, Nature. (c) Dependence of urea yield and selectivity on the ratio of O_2 in the atmosphere; EPR measurements of reactive radicals generated during photocatalysis on P25-4 h using DMPO as the trapping agent: (d) detection of OH^\cdot in pure aqueous solution under different atmospheric O_2 ratios; (e) detection of O_2^\cdot in methanol solution under different atmospheric O_2 ratios; (f) detection of both OH^\cdot and NH_2^\cdot in 2 M ammonia aqueous solution under different atmospheric O_2 ratios. (g) Comparisons of EPR signal obtained in 2 M ammonia under 20% O_2 (red trace in (f)) with theoretically fitted EPR signals. (h) Performance comparison under the most optimized conditions (20% O_2 , 80% CO and pH = 9) and control systems lacking O_2 , CO, ammonia or light. Reproduced with permission from ref. 46. Copyright 2025, Wiley-VCH.

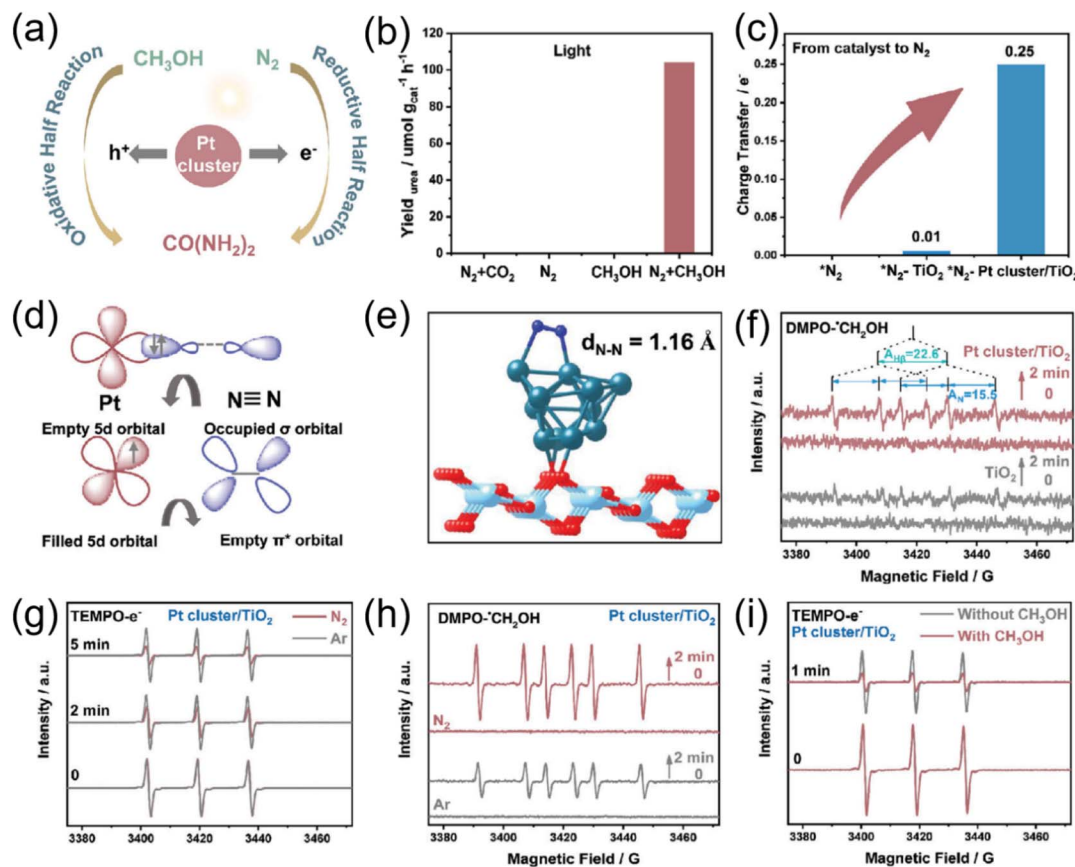


Fig. 17 (a) Schematic diagram of the photocatalytic synergistic redox synthesis system; (b) blank control experiments of N_2 , CH_3OH , and light for urea synthesis; (c) bader calculation of N_2 on TiO_2 and Pt cluster/ TiO_2 ; (d) a simplified schematic diagram of N_2 bonding to a Pt center; (e) schematic diagram of N_2 on the Pt cluster/ TiO_2 surface; (f) EPR results of the captured oxidative DMPO- CH_2OH on the TiO_2 and Pt cluster/ TiO_2 ; (g) EPR results of reductive TEMPO- e^- generation on the Pt cluster/ TiO_2 under N_2 or Ar atmosphere; (h) EPR results of the captured oxidative DMPO- CH_2OH on the Pt cluster/ TiO_2 under N_2 and Ar atmosphere; (i) EPR results of reductive TEMPO- e^- generation on the Pt cluster/ TiO_2 with or without CH_3OH during the photoredox reaction. Reproduced with permission from ref. 40. Copyright 2024, Wiley-VCH.

Table 2 Comparison of the urea production performance of reported photocatalysts. Reproduced with permission from ref. 23. Copyright 2025, Wiley-VCH

Photocatalysts	Reactants	Urea formation rate [$\mu\text{mol g}^{-1} \text{h}^{-1}$]	Light source	AQY	Cyclic stability
Cu SA-TiO ₂	N_2 , CO_2 , H_2O	7.2	365 nm	—	10 h
Ni ₁ -CdS/WO ₃	N_2 , CO_2	7.8	385 nm	0.15%	10 h
2D-CdS@3D-BiOBr	N_2 , CO_2 , H_2O	15	Visible light	3.93%	2 h
TiO ₂ /10 wt%–Fe ₂ TiO ₅	NH_3 , CO_2	3	UV	—	12 h
SrTiO ₃ –FeS–CoWO ₄	N_2 , CO_2	134.24	UV-vis	—	4 h
Pt cluster/TiO ₂	N_2 , CH_3OH	105.68	—	—	9 h
Ru-TiO ₂	N_2 , CO_2	24.95	380 nm	4.7%	3.75 h
Pd-CeO ₂	N_2 , CO_2	9.2	UV-vis	—	15 h
CeO _{2-x}	N_2 , CO_2	15.5	200–400 nm	3.93%	15 h
15%NiCoP-ZnIn ₂ S _{4-x}	N_2 , CO_2 , H_2O	19.6	420 nm	9.2%	9 h
Ti ³⁺ -TiO ₂	N_2 , CO_2 , H_2O	177.53	UV-vis	—	12 h

demonstrate that photogenerated electrons (e^-) and hole (h^+) were synchronously consumed through N_2 reduction and CH_3OH oxidation (Fig. 17f–i), thereby accelerating urea synthesis. The crucial step of C–N coupling was initiated by the reaction between $^*\text{NH}-\text{NH}$ and $^*\text{CHO}$ intermediate, facilitated

by the low energy barrier on Pt cluster/TiO₂. For comparative analysis, the key performance parameters of photocatalysts employed in urea synthesis, including such metrics as urea yield, stability, and reaction conditions, are summarized in Table 2.

4 Conclusions

Photocatalytic synthesis of urea *via* converting CO_2 and $\text{N}_2/\text{NH}_3/\text{NO}_3^-$ into high-value urea chemical under ambient conditions is a green and sustainable technology driven by clean and renewable energy. In this review, we systematically describe the basic principle details for photocatalytic urea synthesis, including fundamental mechanisms (light absorption by photocatalyst, separation and migration of the photogenerated charge carriers, surface redox reactions-activation of carbon and nitrogen reactants and C–N coupling), identification of key intermediates and product identification and quantification. We emphatically review experimental study, catalyst design and theoretical research progress on urea photosynthesis. Various systems, including CO_2-N_2 , $\text{CO}_2-\text{NO}_3^-$ and CO_2-NH_3 systems, *etc.*, have been systematically summarized. Urea has been successfully produced on photocatalyst-like metal oxides (TiO_2 , CeO_2), single-atom (Cu SA-TiO_2), MOF (Ce-BTC), inorganic composites ($\text{Ti}^{3+}-\text{TiO}_2/\text{Fe-CNTs}$) from the conversion of CO_2 and types of nitrogen sources. To enhance the selectivity and efficiency of photosynthesis of urea, various photocatalysts have been designed and developed, including single-atom catalysts, bimetallic site catalysts, heterojunction catalysts and defect engineering, *etc.* The C–N coupling is a key step in urea synthesis, involving the activation and coupling of CO_2 and nitrogen sources. The variation of catalysts and reaction conditions can lead to different reaction mechanisms. For instance, direct C–N coupling, where CO_2 and N_2 are directly coupled on the catalyst surface to form urea. In addition, indirect C–N coupling means that the nitrogen source is first reduced to intermediates (such as $^*\text{NH}_2$, $^*\text{NO}$, *etc.*), and then coupled with $^*\text{CO}$ produced by the reduction of CO_2 to form urea. In some cases, the photogenerated electrons and holes respectively activate CO_2 and N_2 , promoting C–N coupling. Despite the diversity in raw materials, high-performance photocatalysts universally rely on a synergistic combination of three core aspects: (i) optimal adsorption and activation: the catalyst must possess active sites that can effectively adsorb and activate both N_2 (or other N-sources) and CO_2 (or other C-sources), often requiring dual or multiple active sites. (ii) Efficient charge separation and migration: a well-designed heterostructure or energy band alignment is crucial to spatially separate photogenerated electrons and holes and direct them to the respective reduction and oxidation sites. (iii) Facilitated C–N coupling: the local microenvironment and electronic structure of the active sites should be tuned to lower the energy barrier for the key C–N coupling step, which is often the rate-determining step. Although some exciting progress has been made, research in this area is still at the infant stage and requires further works. Firstly, the efficiency of C–N coupling is beyond satisfactory due to the difficulty in achieving the co-adsorption, activation and coupling processes of multiple reactants, such as the large difference in activation energy between CO_2 and $\text{N}_2/\text{NH}_3/\text{NO}_x$. Additionally, there is still controversy regarding the reaction mechanism, particularly concerning the activation of reactants, the formation of key intermediates, and the reaction pathway of

the coupling step. Besides, the currently developed photocatalysts for urea photosynthesis mainly response to ultraviolet region. The inherently contradiction between more visible light absorption and the thermodynamic feasibility of urea photosynthesis is largely challenge due to a relatively high potential of the CO_2RR to form $^*\text{CO}$. An ideal future photocatalyst for urea synthesis should embody the following key properties: (i) broad-spectrum solar energy harvesting: the ability to utilize a wider range of the solar spectrum, including visible and even near-infrared light. (ii) Atomic-level precision in active sites: catalysts with well-defined, single-atom or cluster sites tailored for specific reactant adsorption and C–N coupling. (iii) Unprecedented selectivity: near 100% selectivity towards urea, effectively suppressing competing reactions (*e.g.*, NH_3 emission, H_2 evolution). (iv) Exceptional long-term stability: robustness against photocorrosion, poisoning, and structural degradation over prolonged operation. Therefore, the design of multifunctional visible response photocatalysts, the development and application of advanced characterization techniques, and the exploration of novel reaction pathways require further research to address these challenges.

In summary, we hope that this review provides an overview of the current status and inspires greater interest in the development of alternative photocatalytic urea synthesis. Meanwhile, it is expected that through continuous innovation in photocatalyst design, optimization of reaction conditions, and in-depth understanding of reaction mechanisms, efficient photocatalytic urea synthesis can be achieved, making contributions to addressing energy crises and environmental issues.

Author contributions

Miss Peixia Li: writing original draft and conceptualization. Dr Zhidong Yang: supervision and writing review & editing.

Conflicts of interest

There are no conflicts of interest to declare.

Data availability

No primary research results, software or code have been included and no new data were generated or analysed as part of this review.

Acknowledgements

This project was financially supported by the Scientific Research Program Funded by Education Department of Shaanxi Provincial Government (No. 24JK0280), High-level Talents Foundation of Ankang University (No. 2023AYQDZR17).

References

- 1 Y. Wang, D. Chen, C. Chen and S. Wang, *Acc. Chem. Res.*, 2023, 57, 247–256.
- 2 H. Wang, Z. Xin and Y. Li, *Top. Curr. Chem.*, 2017, 375, 49.

3 P. Xing, S. Wei, X. Chen, H. Luo, L. Dai and Y. Wang, *Chem.-Eng. J.*, 2024, **494**, 153135.

4 C. Lv, L. Zhong, H. Liu, Z. Fang, C. Yan, M. Chen, Y. Kong, C. Lee, D. Liu, S. Li, J. Liu, L. Song, G. Chen, Q. Yan and G. Yu, *Nat. Sustain.*, 2021, **4**, 868–876.

5 D. Jiao, Y. Dong, X. Cui, Q. Cai, C. R. Cabrera, J. Zhao and Z. Chen, *J. Mater. Chem. A*, 2023, **11**, 232–240.

6 M. Li, Q. Shi, Z. Li, M. Xu, S. Yu, Y. Wang, S. M. Xu and H. Duan, *Angew. Chem., Int. Ed.*, 2024, **63**, e202406515.

7 P. Li, Q. Zhu, J. Liu, T. Wu, X. Song, Q. Meng, X. Kang, X. Sun and B. Han, *Chem. Sci.*, 2024, **15**, 3233–3239.

8 Z. Zeng, C. Liao and L. Yu, *Chin. Chem. Lett.*, 2024, **35**, 109349.

9 C. Chen, X. Zhu, X. Wen, Y. Zhou, L. Zhou, H. Li, L. Tao, Q. Li, S. Du, T. Liu, D. Yan, C. Xie, Y. Zou, Y. Wang, R. Chen, J. Huo, Y. Li, J. Cheng, H. Su, X. Zhao, W. Cheng, Q. Liu, H. Lin, J. Luo, J. Chen, M. Dong, K. Cheng, C. Li and S. Wang, *Nat. Chem.*, 2020, **12**, 717–724.

10 D.-S. Huang, X.-F. Qiu, J.-R. Huang, M. Mao, L. Liu, Y. Han, Z.-H. Zhao, P.-Q. Liao and X.-M. Chen, *Nat. Synth.*, 2024, **3**, 1404–1413.

11 P. Xing, S. Wei, Y. Zhang, X. Chen, L. Dai and Y. Wang, *ACS Appl. Mater. Interfaces*, 2023, **15**, 22101–22111.

12 B. Srinivas, V. D. Kumari, G. Sadanandam, C. Hymavathi, M. Subrahmanyam and B. R. De, *Photochem. Photobiol.*, 2011, **88**, 233–241.

13 H. Cai, J. Ding, T. Hou, T. Wei, Q. Liu, J. Luo, L. Feng, W. Liu and X. Liu, *Chem. Synth.*, 2024, **4**, 100378.

14 N. Meng, Y. Huang, Y. Liu, Y. Yu and B. Zhang, *Cell Rep. Phys. Sci.*, 2021, **2**, 100378.

15 Y. Luo, K. Xie, P. Ou, C. Lavallais, T. Peng, Z. Chen, Z. Zhang, N. Wang, X.-Y. Li, I. Grigioni, B. Liu, D. Sinton, J. B. Dunn and E. H. Sargent, *Nat. Catal.*, 2023, **6**, 939–948.

16 Z. Mei, Y. Zhou, W. Lv, S. Tong, X. Yang, L. Chen and N. Zhang, *ACS Sustain. Chem. Eng.*, 2022, **10**, 12477–12496.

17 C. Mao, J. Byun, H. W. MacLeod, C. T. Maravelias and G. A. Ozin, *Joule*, 2024, **8**, 1224–1238.

18 Y. Lou, H. Chen, L. Wang, S. Chen, Y. Song, Y. Ding, Z. Hao, C. He, D. Qiu, H. Li, J. Wang, D. Liu and X. Cui, *ACS Sustain. Chem. Eng.*, 2025, **13**, 151–164.

19 L. Kuang, Z. Chen, Y. Yan, F. Guo and W. Shi, *Int. J. Hydrogen Energy*, 2024, **87**, 20–49.

20 J. Yuan, L. Hu, J. Huang, Y. Chen, S. Qiao and H. Xie, *Appl. Catal., B*, 2023, **339**, 123146.

21 B. Bhattacharyya, R. M. Sarhan, Y. Lu and A. Taubert, *ChemCatChem*, 2024, **16**, 202400638.

22 H. Song, D. A. Chipoco Haro, P.-W. Huang, L. Barrera and M. C. Hatzell, *Acc. Chem. Res.*, 2023, **56**, 2944–2953.

23 Z. Lu, R. Chen, G. Liu, B. Xia, K. Fan, T. Liu, Y. Xia, S. Liu and B. You, *Adv. Funct. Mater.*, 2025, 2500944.

24 Z. Zeng, Y. Chen, X. Zhu and L. Yu, *Chin. Chem. Lett.*, 2023, **34**, 107728.

25 J.-Y. Zeng, X.-S. Wang, X.-H. Liu, Q.-R. Li, J. Feng and X.-Z. Zhang, *Natl. Sci. Rev.*, 2023, **10**, nwad142.

26 C. Liu, F. Gong, Q. Zhou and Y. Xie, *Energy Fuels*, 2024, **38**, 8951–8959.

27 S. Zhang, Y. Zhao, R. Shi, G. I. N. Waterhouse and T. Zhang, *EnergyChem*, 2019, **1**, 100013.

28 Z. Li, P. Zhou, M. Zhou, H. Jiang, H. Li, S. Liu, H. Zhang, S. Yang and Z. Zhang, *Appl. Catal., B*, 2023, **338**, 122962.

29 Z. Yang, H. Zhang, J. Zhao, H. Shi, Y. Liu, H. Yang and P. Yang, *ChemSusChem*, 2022, **15**, e202200260.

30 Z. Yang, Y. Zhang, H. Zhang, J. Zhao, H. Shi, M. Zhang, H. Yang, Z. Zheng and P. Yang, *J. Catal.*, 2022, **409**, 12–23.

31 S. N. Habisreutinger, L. Schmidt-Mende and J. K. Stolarczyk, *Angew. Chem., Int. Ed.*, 2013, **52**, 7372–7408.

32 S. Lian, M. S. Kodaimati, D. S. Dolzhnikov, R. Calzada and E. A. Weiss, *J. Am. Chem. Soc.*, 2017, **139**, 8931–8938.

33 P. Yang and Z. Yang, *J. Energy Chem.*, 2020, **50**, 365–377.

34 Z. Guo, P. Zhou, L. Jiang, S. Liu, Y. Yang, Z. Li, P. Wu, Z. Zhang and H. Li, *Adv. Mater.*, 2024, **36**, 2311149.

35 W. Yang, R. Godin, H. Kasap, B. Moss, Y. Dong, S. A. J. Hillman, L. Steier, E. Reisner and J. R. Durrant, *J. Am. Chem. Soc.*, 2019, **141**, 11219–11229.

36 J. J. Leung, J. A. Vigil, J. Warnan, E. Edwardes Moore and E. Reisner, *Angew. Chem., Int. Ed.*, 2019, **58**, 7697–7701.

37 Y.-F. Mu, J.-L. Zhou, S.-X. Yuan, M.-R. Zhang, H. Pang, M. Zhang and T.-B. Lu, *Chem Catal.*, 2025, **5**, 101433.

38 N. Li, J. Zhang, X. Xie, K. Wang, D. Qi, J. Liu, Y.-Q. Lan and J. Jiang, *Nat. Commun.*, 2025, **16**, 1106.

39 J. Zheng, S. Xu, J. Sun, J. Zhang, L. Sun, X. Pan, L. Li and G. Zhao, *Appl. Catal., B*, 2023, **338**, 123056.

40 W. Yang, L. Xiao, W. Dai, S. Mou and F. Dong, *Adv. Energy Mater.*, 2024, **14**, 2303806.

41 B. Hu, B.-H. Wang, L. Chen, Z.-J. Bai, W. Zhou, J.-K. Guo, S. Shen, T.-L. Xie, C.-T. Au, L.-L. Jiang and S.-F. Yin, *ACS Catal.*, 2022, **12**, 11860–11869.

42 S. Lin, X. Zhang, L. Chen, Q. Zhang, L. Ma and J. Liu, *Green Chem.*, 2022, **24**, 9003–9026.

43 D. Li, Y. Zhao, Y. Miao, C. Zhou, L. P. Zhang, L. Z. Wu and T. Zhang, *Adv. Mater.*, 2022, **34**, 2207793.

44 S. Yang, J. Deng, J. Chen, Q. Tan, T. Liu, K. Chen, D. Han, Y. Ma, M. Dai and L. Niu, *Catal. Sci. Technol.*, 2023, **13**, 1855–1865.

45 Y. Zhang, Y. Sun, Q. Wang, Z. Zhuang, Z. Ma, L. Liu, G. Wang, D. Wang and X. Zheng, *Angew. Chem., Int. Ed.*, 2024, **63**, e202405637.

46 X. Huang, S. Xie, B. Sheng, B. Xiao, C. Chen, H. Sheng and J. Zhao, *Angew. Chem., Int. Ed.*, 2025, **64**, e202505630.

47 C. Shi, K. Xia, L. Zhang, M. Guo, X. Guan, C. Gu, X. Yang, Y. Wang, X. Liu and X. Ding, *Angew. Chem., Int. Ed.*, 2024, **14**, 2400201.

48 M. Zhou, Y. Zhang, H. Li, Z. Li, S. Wang, X. Lu and S. Yang, *Angew. Chem., Int. Ed.*, 2025, **64**, e202414392.

49 H. Y. S. Kuwabata and H. Yoneyama, *Langmuir*, 1998, **14**, 1899–1904.

50 T. T. B. J. Liu and H. Yoneyama, *J. Photochem. Photobiol., A*, 1998, **115**, 227–230.

51 H. M. h. D. G. Shchukin, *Langmuir*, 2005, **21**, 5582–5587.

52 E. A. Ustinovich, D. G. Shchukin and D. V. Sviridov, *J. Photochem. Photobiol., A*, 2005, **175**, 249–252.

53 H. Sun, Z. Lin, R. Tang, Y. Liang, S. Zou, X. Zhang, K. Chen, R. Zheng and J. Huang, *Appl. Catal., B*, 2025, **360**, 124511.

54 R. Tan, S. Meng, P. Wang, C. Yang, J. Yao, H. Li, T. Zhang and Z. Li, *Nano Res.*, 2025, **18**, 94907647.

55 Z. Ma, X. Xia, B. Song, R. Li, X. Wang and Y. Huang, *ACS Appl. Mater. Interfaces*, 2024, **16**, 46323–46331.

56 Y. L. X. Zhang, C. Chena, J. Zheng, S. P. Jiang and S. Wang, *Proc. Natl. Acad. Sci. U. S. A.*, 2024, **121**, e2311326121.

57 W. J. Dong, J. P. Menzel, Z. Ye, I. A. Navid, P. Zhou, K. R. Yang, V. S. Batista and Z. Mi, *ACS Catal.*, 2024, **14**, 2588–2596.

58 H. Maimaiti, B. Xu, J.-Y. Sun and L.-R. Feng, *ACS Sustain. Chem. Eng.*, 2021, **9**, 6991–7002.

59 Y. Wang, S. Wang, J. Gan, J. Shen, Z. Zhang, H. Zheng and X. Wang, *ACS Sustain. Chem. Eng.*, 2023, **11**, 1962–1973.

60 M. Yuan, J. Chen, Y. Bai, Z. Liu, J. Zhang, T. Zhao, Q. Wang, S. Li, H. He and G. Zhang, *Angew. Chem., Int. Ed.*, 2021, **60**, 10910–10918.

61 Y. Zhang, T. Hou, Q. Xu, Q. Wang, Y. Bai, S. Yang, D. Rao, L. Wu, H. Pan, J. Chen, G. Wang, J. Zhu, T. Yao and X. Zheng, *Adv. Sci.*, 2021, **8**, 2100302.

62 Y. He, L. Yin, N. Yuan and G. Zhang, *Chem.-Eng. J.*, 2024, **481**, 148754.

63 S. Yang, W. Zhang, G. Pan, J. Chen, J. Deng, K. Chen, X. Xie, D. Han, M. Dai and L. Niu, *Angew. Chem., Int. Ed.*, 2023, **62**, e202312076.

64 G. Ren, X. Chen, Z. Zhao, Z. Li and X. Meng, *Adv. Funct. Mater.*, 2025, 2506296.

65 H. Liang, C. Ye, Y. Wu, Y. Li, R. Long, J. Xiong, W. Jiang and J. Di, *Mater. Today*, 2025, **86**, 96–103.

66 H. Sun, X. Qin, Y. Zhang, Y. Xu, Y. Jiang, F. Pan and Q. Liu, *Small*, 2025, **21**, 2503761.

67 F. Yang, H. Han, H. Duan, F. Fan, S. Chen, B. Yu Xia and Y. L. He, *Adv. Energy Mater.*, 2025, **15**, 2405726.

68 Y. Zhang, B. Xia, J. Ran, K. Davey and S. Z. Qiao, *Adv. Energy Mater.*, 2020, **10**, 1903879.

69 Y. Jiao, H. Li, Y. Jiao and S.-Z. Qiao, *J. Am. Chem. Soc.*, 2023, **145**, 15572–15580.

70 D. Anastasiadou and M. Costa Figueiredo, *ACS Catal.*, 2024, **14**, 5088–5097.

71 W.-Y. Lin, Z.-X. Chen, H. Xiong, H.-C. Li, Y.-S. Ho, C.-T. Hsieh, Q. Lu and M.-J. Cheng, *ACS Catal.*, 2023, **13**, 11697–11710.

72 S. Su, X. Li, W. Ding, Y. Cao, S. Yuan, Z. Liu, Y. Yang, Y. Ding and M. Luo, *J. Mater. Chem. A*, 2024, **12**, 15300–15310.

73 Q. Wang, Y. Wan, Q. Liu, Y. Zhang, Z. Ma, Z. Xu, P. Sun, G. Wang, H.-L. Jiang, W. Sun and X. Zheng, *Sci. Bull.*, 2025, **70**, 1118–1125.

74 J. Long, D. Luan, X. Fu, H. Li and J. Xiao, *ACS Catal.*, 2024, **14**, 14678–14687.

75 K. Qi, B. Cheng, J. Yu and W. Ho, *Chin. J. Catal.*, 2017, **38**, 1936–1955.

76 J. Li, H. Yuan, W. Zhang, B. Jin, Q. Feng, J. Huang and Z. Jiao, *Carbon Energy*, 2022, **4**, 294–331.

77 Q. Xu, L. Zhang, J. Yu, S. Wageh, A. A. Al-Ghamdi and M. Jaroniec, *Mater. Today*, 2018, **21**, 1042–1063.

78 X. Li, H. Sun, Y. Xie, Y. Liang, X. Gong, P. Qin, L. Jiang, J. Guo, C. Liu and Z. Wu, *Coord. Chem. Rev.*, 2022, **467**, 214596.

79 R. Sun, Z. Zhu, N. Tian, Y. Zhang and H. Huang, *Angew. Chem., Int. Ed.*, 2024, **63**, e202408862.

80 Z. Li, J. Xiong, Y. Huang, Y. Huang, G. I. N. Waterhouse, Z. Wang, Y. Mao, Z. Liang and X. Luo, *Chem.-Eng. J.*, 2024, **486**, 150304.

81 X. Fu, H. Huang, G. Tang, J. Zhang, J. Sheng and H. Tang, *Chin. J. Struct. Chem.*, 2024, **43**, 100214.

82 M. I. Ahmad, Y. Liu, Y. Wang, P. Cao, H. Yu, H. Li, S. Chen and X. Quan, *Angew. Chem., Int. Ed.*, 2024, **64**, 167328.

83 X. Xiang, L. Guo, X. Wu, X. Ma and Y. Xia, *Environ. Chem. Lett.*, 2012, **10**, 295–300.

84 M.-A. Mohajer, P. Basuri, A. Evdokimov, G. David, D. Zindel, E. Miliordos and R. Signorell, *Science*, 2025, **388**, 1426–1430.

85 I. P. Moura, A. C. Reis, A. E. Bresciani and R. M. B. Alves, *Renew. Sustain. Energy Rev.*, 2021, **151**, 111619.

86 J. Ding, R. Ye, Y. Fu, Y. He, Y. Wu, Y. Zhang, Q. Zhong, H. H. Kung and M. Fan, *Nat. Commun.*, 2023, **14**, 4586.

87 C. Zhao, Y. Jin, J. Yuan, Q. Hou, H. Li, X. Yan, H. Ou and G. Yang, *J. Am. Chem. Soc.*, 2025, **147**, 8871–8880.

