Recent advancements in g-C₃N₄-based photocatalysts for photocatalytic CO₂ reduction: a mini review

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Carbon dioxide (CO₂) is a very important micro-molecular resource. Using CO₂ captured from the atmosphere for high-output synthesis of chemicals as raw materials has great significance and potential for various industrial applications. Since the industrial revolution in the 18th century, manmade CO₂ emission has increased by 45%, which negatively impacts the planetary climate by the so-called greenhouse effect. Therefore, high-efficiency photocatalysis and photocatalysts for CO₂ conversion have become the most important challenges and milestones throughout the world. In consideration of this, various catalysts have been explored. Among these, graphitic carbon nitride (g-C₃N₄) as a semiconductor is emerging as a highly promising photocatalyst for removing CO₂ from the atmosphere. Moreover, due to its excellent chemical stability and unique band structure, g-C₃N₄ has exhibited significant application potential for photocatalysis. This review summarizes the advancements that have been made in the synthesis and photocatalytic applications of g-C₃N₄-based catalysts for CO₂ reduction in recent years and explains the future challenges and prospects in this vital area of research.

1. Introduction

Carbon dioxide (CO₂) is a major greenhouse gas that contributes approximately 20% to the greenhouse effect. Its emissions are increasing and are causing a significant negative influence on the overall climate and life on our planet. It was reported that the annual global carbon dioxide emissions have exceeded 30 billion tons, and only 1% of this amount is being removed annually. However, these emissions are not considered to be solely waste because CO₂ is a very important micro-molecular resource for many industrial chemicals, such as methane, methyl alcohol, and dimethylformamide (DMF). Artificial synthesis of industrial-scale chemicals with CO₂ captured from the atmosphere not only removes the harmful CO₂, but also produces valuable chemical products for industries.

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Since 1970, it has been a challenge worldwide to develop a facile and efficient method for the photocatalytic reduction of CO₂. Three types of CO₂ photoreduction systems have been developed, i.e., the semiconductor photocatalytic system, the metal complex photocatalytic system, and the enzyme photocatalytic system.² In 1979, Fujishima et al. reported photocatalytic CO₂ reduction using semiconductors TiO₂ and CdS as photocatalysts, and these were among the earliest photocatalytic systems for CO₂ reduction.³ In 1983, Lehn et al. reported for the first time a CO₂ reduction system using fac-Re[bpym]₂[CO]₃Cl as a photocatalyst, and it exhibited high selectivity and high quantum yield.⁴ In 1986, Willner et al. reported a photocatalytic CO₂ reduction system using enzymes as catalysts.⁵

Since their discoveries, all three types of CO₂ photoreduction systems have been significantly developed. Due to the high substrate specificity, high conversion rates, excessive sensitivity, high cost, and narrow applicability of the enzymatic catalysts, their broad utilization has been limited.⁶ Thus, most of the interest has been focused on semiconductor or metal complex-based catalysts. In general, metal complex systems are based on homogeneous organic photocatalysts. Because all the metal atoms can serve as active sites, the catalytic efficiency and selectivity of the metal complex systems are sensitive to energy distribution. Moreover, due to the homogeneous structure, there are few side reactions accompanying metal complex photocatalysis.⁷,⁸ However, it is difficult to separate reactants from products because of the homogeneous structure, which limits the cyclic utilization and can be disadvantageous for sustainable development. In comparison, semiconductor-based photocatalysts are mostly heterogeneous catalysts, and with their use, it is easier to separate products from the reaction system.⁹ Furthermore, the photocatalytic performance can be improved through various structural and chemical modifications of the photocatalyst, e.g., porous semiconductor photocatalysts, doping¹⁰–¹⁴ or compositing of a semiconductor photocatalyst with other materials.¹⁵–¹７ Thus, there has been great interest and widespread usage of semiconductor photocatalysis during recent years.

Among the various types of semiconductors, graphitic carbon nitride (g-C₃N₄) is considered promising because it is composed of triazine rings and tri-s-triazine (heptazine) rings as its basic unit, and it forms a two-dimensional lamellar structure (Fig. 1).¹⁸ Because of the structure, it is insoluble in most solvents such as water, ethanol, ether, toluene, DMF, and tetrahydrofuran (THF).¹⁹ g-C₃N₄ is a typical polymer semiconductor with a band gap of 2.7 eV that can absorb blue-violet light with a wavelength less than 475 nm in the solar spectrum,²⁰ and thus, it is a potential candidate for photocatalysis.²¹,²² In 2009, Wang et al. for the first time reported photocatalytic hydrogen production using g-C₃N₄ as a photocatalyst.²³ In 2013, Peng et al. used g-C₃N₄ in CO₂ photoreduction.²⁴ This review will focus on the present pathways for the applications of g-C₃N₄ in the field of photocatalytic CO₂ reduction. Finally, the conclusion of the article addresses the current challenges and future development of g-C₃N₄-based photocatalysts.

2. Fundamentals of photocatalytic CO₂ reduction on semiconductors

Photocatalytic CO₂ reduction is a process where light energy is used as the input energy, and the product can be controlled by changing the reaction conditions and types of catalysts.²⁵,²⁶ The common reaction pathways are listed as follows.

\[
\begin{align*}
\text{CO}_2 + 2\text{H}^+ + 2\text{e}^- & \rightarrow \text{HCOOH}, E^0 = -0.61 \text{ V} & (I) \\
\text{CO}_2 + 2\text{H}^+ + 2\text{e}^- & \rightarrow \text{CO} + \text{H}_2\text{O}, E^0 = -0.53 \text{ V} & (II) \\
\text{CO}_2 + 4\text{H}^+ + 4\text{e}^- & \rightarrow \text{HCHO} + \text{H}_2\text{O}, E^0 = -0.48 \text{ V} & (III) \\
\text{CO}_2 + 6\text{H}^+ + 6\text{e}^- & \rightarrow \text{CH}_3\text{OH} + \text{H}_2\text{O}, E^0 = -0.38 \text{ V} & (IV) \\
\text{CO}_2 + 8\text{H}^+ + 8\text{e}^- & \rightarrow \text{CH}_4 + 2\text{H}_2\text{O}, E^0 = -0.24 \text{ V} & (V)
\end{align*}
\]

The principle of photocatalytic CO₂ reduction consists of five main steps: light adsorption (i), charge separation (ii), CO₂ adsorption (iii), surface redox reaction (iv), and product desorption (v).²⁷ To trigger the CO₂ reduction reaction, the conduction band edge of the photocatalysts should be more negative than the reaction potential, and they are listed above (vs. NHE).²⁸ To achieve efficient generation and transfer of photo-excited electrons and holes, certain modifications of photocatalysts are required. To increase the effectiveness of the process, a semiconductor-based photocatalyst should be designed considering these issues.

First, a semiconductor with a band gap of 2.0–3.0 eV is ideal for visible light adsorption.²⁷ Second, the separation of
electron–hole pairs is sufficiently large so that the recombination of electrons and holes is limited.\textsuperscript{30} Third, to enhance the adsorption of CO\textsubscript{2}, the surface area of photocatalysts should be enlarged, which can be accomplished by providing more active sites or introducing alkaline components into the photocatalysts.\textsuperscript{31} Fourth, to improve the efficiency of surface redox reactions, one can introduce a cocatalyst or prepare composites to enhance the charge transfer rate. Finally, accurate timing of the release of the products is necessary to increase the conversion and reaction rate.

3. \textit{g-C\textsubscript{3}N\textsubscript{4} as photocatalyst}

A good photocatalyst requires a suitable energy band structure to ensure light harvesting ability and efficient charge transfer. Among varieties of photocatalysts, oxide semiconductors usually have too large band gap energies to sufficiently utilize incident light, while the chalcogenide semiconductors often have a narrow band gap, which results in easy recombination of the photogenerated charge carriers (Fig. 2). In comparison, \textit{g-C\textsubscript{3}N\textsubscript{4}} is a non-metal semiconductor with a band gap of 2.7 eV and suitable conduction band (CB) and valence band (VB) positions, enabling it to absorb visible light. Moreover, \textit{g-C\textsubscript{3}N\textsubscript{4}} can be prepared by facile methods, and the microstructure of \textit{g-C\textsubscript{3}N\textsubscript{4}} can be easily adjusted to create pores as active sites or enlarge the surface area for more optimal light absorption. Because of these advantages, \textit{g-C\textsubscript{3}N\textsubscript{4}} is a superior photocatalyst for visible-light-driven CO\textsubscript{2} reduction compared to TiO\textsubscript{2} and other common photocatalysts. However, because the 2.7 eV band gap of \textit{g-C\textsubscript{3}N\textsubscript{4}} is insufficiently large, the separated electron–hole pairs tend to recombine, thus limiting the effective separation of charges and subsequent redox reactions. Although it has numerous Lewis base sites, \textit{e.g.}, Bronsted base sites and nucleophile sites that are favourable for CO\textsubscript{2} adsorption, bulk \textit{g-C\textsubscript{3}N\textsubscript{4}} prepared by direct calcination has a relatively small surface area. Therefore, several modifications including structural tuning,\textsuperscript{31-37} elemental doping,\textsuperscript{38-45} addition of cocatalyst,\textsuperscript{46-48} and compositing\textsuperscript{41-70} have been used to increase the adsorption and obtain more effective charge separation.

3.1 \textit{g-C\textsubscript{3}N\textsubscript{4} with nanostructures and/or defects}

It is expected that \textit{g-C\textsubscript{3}N\textsubscript{4}} has good potential for CO\textsubscript{2} fixation and activation. However, the bulk form obtained from some precursors has a small surface area, and thus, has low adsorption efficiency and catalytic activity. To alleviate this drawback, researchers have taken specific measures to increase the surface area and improve the functionality of \textit{g-C\textsubscript{3}N\textsubscript{4}}. In this regard, Zhang \textit{et al.} applied porous structured \textit{g-C\textsubscript{3}N\textsubscript{4}} to a CO\textsubscript{2} photocatalytic reduction to CO under visible light irradiation.\textsuperscript{31} Moreover, the \textit{g-C\textsubscript{3}N\textsubscript{4}} was synthesized by calcination, using either melamine or melamine hydrochloride as the precursor. It was found that the latter has a surface area 39 times greater than bulk \textit{g-C\textsubscript{3}N\textsubscript{4}} due to the porous structure, but the performance of CO\textsubscript{2} photoreduction was not obviously improved. This occurred because the porous structure not only endows \textit{g-C\textsubscript{3}N\textsubscript{4}} with a higher surface area but also enlarges the band gap, which resulted in more difficult electron excitation and the lower yield of CO\textsubscript{2}. Although it failed to improve the performance of CO\textsubscript{2} photoreduction, there was still direct significance to this pioneering research in that the structural tuning of \textit{g-C\textsubscript{3}N\textsubscript{4}} resulted in great improvement of the photoreactivity.

![Fig. 2](https://example.com/fig2)  
**Fig. 2** Band gap energies and CB (green) and VB (red) edge positions of the selected photocatalysts with respect to the vacuum level and NHE. The two dashed lines indicate the water redox reaction potentials.\textsuperscript{29} Copyright 2016 WILEY-VCH.

![Fig. 3](https://example.com/fig3)  
**Fig. 3** Synthetic process for helical nanorod-like graphitic carbon nitride (HR-CN) based on chiral mesoporous silica (CMS) as the template and cyanamide (CY) as the precursor.\textsuperscript{32} Copyright 2014, Wiley-VCH.
Thus far, different varieties of g-C3N4 nanostructures have been developed, such as porous structures, 1D nanorods, nanowires, 2D nanosheets, and 3D nanostructures.\textsuperscript{12-37} As a typical example, Zheng and Wang \textit{et al.} imprinted helical g-C3N4 nanorods with chiral silicon dioxide as templates (Fig. 3).\textsuperscript{32} This helical structure caused multiple reflections of incident light, leading to an improved light-harvesting capability across the entire optical spectrum. Although the band gap increased from 2.66 eV to 2.75 eV, the helical g-C3N4 still exhibited an overall enhanced optical absorption compared to bulk g-C3N4. A lower photoluminescence (PL) intensity and a stronger PL quenching was measured for the helical g-C3N4 nanorods as compared to the bulk g-C3N4, which indicated that the recombination of photogenerated charge carriers in the helical g-C3N4 nanorods was suppressed.\textsuperscript{71} The CO production and selectivity of the helical g-C3N4 system during visible-light CO\textsubscript{2} reduction reached 8.9 \textmu mol and 96.7\%, respectively. In recent years, there has been much experimentation with 2D g-C3N4 nanosheets because of their large specific surface area, which is advantageous for light harvesting and gas adsorption. Cao \textit{et al.} prepared ultra-thin g-C3N4 nanosheets through a stepwise NH\textsubscript{3}-mediated thermal exfoliation approach.\textsuperscript{33} The resulting nanosheets were approximately 3 nm thick and were produced with a hierarchical structure due to the amine-induced assembly. This structure endowed g-C3N4 nanosheets with a much higher specific surface area, abundant active sites, shorter diffusion distance of charge carriers, and increased CO\textsubscript{2} adsorption. Therefore, the yields of CH\textsubscript{4} and CH\textsubscript{3}OH with g-C3N4 nanostructures reached 8.9 \textmu mol and 96.7\%, respectively. In recent years, there has been much experimentation with 2D g-C3N4 nanosheets because of their large specific surface area, which is advantageous for light harvesting and gas adsorption. Cao \textit{et al.} prepared ultra-thin g-C3N4 nanosheets through a stepwise NH\textsubscript{3}-mediated thermal exfoliation approach.\textsuperscript{33} The resulting nanosheets were approximately 3 nm thick and were produced with a hierarchical structure due to the amine-induced assembly. This structure endowed g-C3N4 nanosheets with a much higher specific surface area, abundant active sites, shorter diffusion distance of charge carriers, and increased CO\textsubscript{2} adsorption. Therefore, the yields of CH\textsubscript{4} and CH\textsubscript{3}OH with g-C3N4 nanosheets as catalyst were 9.93 and 5.34 times higher than that with bulk g-C3N4 as catalyst, respectively.

Apart from structural tuning, the introduction of defects is also a useful method for promoting the photocactivity of g-C3N4. Zhang \textit{et al.} introduced carbon vacancies into synthesized g-C3N4 with urea and heat treatment under an NH\textsubscript{3} atmosphere to enhance CO photogeneration.\textsuperscript{34} The g-C3N4 with enriched C vacancies exhibited a CO evolution rate of 4.18 \textmu mol g\textsuperscript{-1} h\textsuperscript{-1}, which was 2.3 times higher than the evolution rate of pristine g-C3N4. The improved CO\textsubscript{2} reduction performance was ascribed to greater CO\textsubscript{2} adsorption and activation, an upshifted conduction band, increased charge carrier concentration, and prolonged lifetime. Furthermore, the reactive oxygen species (ROS) and electrochemical impedance spectroscopy (EIS) indicated that the C vacancies could weaken the exciton effect and elevate the charge carrier generation.

In another study, Chai \textit{et al.} introduced N vacancies into ultrathin g-C3N4 nanosheets using NH\textsubscript{4}Cl as a dynamic gas template.\textsuperscript{35} The NH\textsubscript{4}Cl decomposed into NH\textsubscript{3} and HCl gases upon heating, which formed a bubble film on the surface of the polymerized g-C3N4 nanosheets. As the \pi-\pi interlayer interactions in the g-C3N4 nanosheets became diminished with the expansion of the bubble film, an ultrathin bubble g-C3N4 film formed, and N vacancies were introduced afterwards by a post-treatment under a reducing atmosphere. The few-atomic-layered structure enlarged the exposed surface area, and this benefited charge carrier transportation. Additionally, N vacancies induced a mid-gap state, which facilitated multi-electron excitation in a two-step process and extended the optical absorption of the g-C3N4 nanosheets to the near infrared region, and thus, the utilization of solar energy was enhanced (Fig. 4). Furthermore, N vacancies could also result in an aftereffect where the decay lifetime was prolonged because the mid-gap state retarded the radiative recombination rate. The g-C3N4 nanosheets with N vacancies exhibited a 5.14-fold higher CH\textsubscript{4} evolution rate than pristine g-C3N4. This pioneering work was intriguing and inspiring because it highlighted the synergistic effect of morphology control and vacancy modulation on photocatalytic CO\textsubscript{2} reduction.

Both structural adjustment and defect tuning are meant to change the energy band structure of g-C3N4 to enhance the light-harvesting ability and charge transfer efficiency, and they are the most convenient methods to improve the photocatalytic performance of g-C3N4.

### 3.2 Non-metal doped g-C3N4

Pristine g-C3N4 exhibits a poor performance due to low CO\textsubscript{2} absorption and poor utilization of photo-excited charge carriers because of the deficiency of electron donation sites and unoptimized electronic structure.\textsuperscript{36} To address these challenges, heteroatom doping was developed to tune the electronic structure of g-C3N4. The doped elements included mainly B, C, O, P, and S.\textsuperscript{38-41} In g-C3N4, N atoms are active sites for photocatalytic reactions, and the electrons are mainly localized around N atoms. Under light irradiation, transfer of excited electrons from N atoms to C atoms is difficult due to the localized electron structure.\textsuperscript{72-74} Moreover, the excited electrons are expected to transfer back for redox reaction, which increases the chances of recombination of the charge carriers. To address these problems, Liu \textit{et al.} prepared boron-doped g-C3N4 by a one-step calcination using boric acid and urea as precursors.\textsuperscript{38} B atoms were doped into cavities between adjacent tri-s-triazine units through coordination with N atoms. The density functional theory (DFT) studies demonstrated that doping of B introduced a new electron excitation pathway from N (2p\textsubscript{z}, 2p\textsubscript{z}) to B (2p\textsubscript{x}, 2p\textsubscript{y}), which improved the charge transfer and localization, and
thus the reaction dynamics. Furthermore, the doped B atoms altered the gas adsorption on the surface of g-C3N4 so that they could subsequently act as active sites. Consequently, the optimal B-doped g-C3N4 resulted in a 32-fold higher yield of CH4 than that of pure g-C3N4.

In another work, Tian et al. synthesized phosphorus-doped g-C3N4 nanotubes by calcinating a mixture of melamine and sodium hypophosphate monohydrate. The nanotube structure attained a much larger surface area than that of g-C3N4 nanosheets. The doping of phosphorus resulted in a shifting down of the conduction band and valence band of g-C3N4, and the band gap was narrowed by 0.12 eV (Fig. 4), which enhanced the optical absorption. The phosphorus doping also created additional amino groups in the surface of g-C3N4, which increased CO2 adsorption due to the acid–base interaction. As a result, the gas yields were 9.48 μmol g⁻¹ for CO and 7.24 μmol g⁻¹ for CH4, which were 3.12 and 13.9 times the yield of pristine g-C3N4, respectively. Similarly, Yu et al. fabricated sulfur-doped g-C3N4 (TCN) by a facile calcinating method using thiourea as the precursor. According to UV-Vis diffuse reflection spectra (UV-Vis DRS), the band gap of TCN was 2.63 eV, which was 0.07 eV narrower than that of un-doped g-C3N4 (MCN). This narrow band gap enabled TCN to absorb more solar energy and produce more charge carriers, thus improving the photocatalytic performance. According to the theoretical calculation, however, the band gaps of TCN and MCN were the same, but an impurity level was introduced due to sulfur doping on TCN. The photoelectrons could jump from the VB to the impurity level or from the impurity level to the CB. As a result, the CH3OH yield of TCN was 2.5 times greater than that of MCN.

It is worth noting that many studies have been recently conducted on metal and non-metal co-doped g-C3N4 for CO2 photoreduction. Metal elements such as K, Na, and Co can improve the photocatalytic performance through a similar mechanism. Zheng et al. prepared N vacancy-rich g-C3N4 co-doped with B and K atoms through a one-step method. In this system, K functioned as an electron donor and promoted interlayer electron transfer, B assisted in maintaining a high reduction potential and counteracted the drawbacks of K, and N vacancies lowered the conduction band minimum (CBM) and promoted CO2 absorption. The synergistic effect of the multiple modifications significantly improved the CO2 reduction performance. The CH4 and CO produced in 5 h were 5.93 μmol g⁻¹ and 3.16 μmol g⁻¹, respectively, and 161% and 527% of the production of the pristine g-C3N4 occurred under the same conditions. It further suggested that the combination of doping and vacancies enhanced the photocatalytic performance of g-C3N4, and provides a new strategy for photocatalyst modification.

Overall, the non-metal doping increased the CO2 photoreduction activity of g-C3N4 by tuning the energy band structure, which manifested as a narrower band gap that provided more effective light absorption in a wider range of spectra, especially in the visible light region. The non-metal doping could also introduce an impurity level or a new electron excitation pathway to g-C3N4, which facilitated the charge transfer. Moreover, the doping process might introduce surface groups, which improved CO2 adsorption due to the acid–base interaction.

3.3 g-C3N4 with co-catalyst
In photocatalysis, suitable co-catalysts can serve as active sites for surface reactions and improve charge separation and charge utilization. For a semiconductor such as g-C3N4, frequently used co-catalysts include metal, carbon materials, reduction co-catalyst, and double co-catalyst. Generally, a metal co-catalyst refers to noble metals such as Au, Ag, and Pt. By depositing the noble metals on a photocatalyst, a Schottky barrier can be formed on the interface, and the metal nanoparticles can become electron traps to facilitate the separation of photogenerated charges. Pt is one of the most common co-catalysts used in CO2 reduction because of its great methane selectivity in a gas–solid system. Yu et al. prepared g-C3N4 by direct calcination with thiourea as a precursor and Pt nanoparticles that were deposited for CO2 photoreduction. The deposited Pt nanoparticles absorbed nearly all the incident light, and thus increased the visible light absorption without changing the band gap of g-C3N4. Moreover, the introduction of Pt nanoparticles promoted the generated photoelectrons to transfer from g-C3N4 to Pt and react with CO2. The yield of CH4 initially increased as the amount of deposited Pt increased from 0 to 1%, and then decreased as the loading of Pt increased to 2% or higher. Similar observations have also been reported by Chai et al. (Fig. 6). This relative reduction in the production of CH4 was attributed to the excess Pt that increased the negative charges on the surface of the g-C3N4 and promoted the recombination of electrons and holes. Another possibility is that excessive or redundant Pt covers the active sites on the surface of g-C3N4 and shields them from light. In another study, Du et al. substituted Pt nanoparticles with individual Pt atoms and found that the Pt/g-C3N4 catalyst preferred to reduce CO2 to CH4, with a rate-determining barrier of 1.16 eV and greatly enhanced visible light absorption.
Another noble metals are also frequently employed to modify g-
C3N4.49,50 Currently, there are few CO2 reduction systems using
non-noble metals.71,72 However, according to a DFT calculation,
Mo could be an efficient co-catalyst on g-C3N4 for CO2
conversion.53

3.3.2 Carbon materials. Carbon materials, e.g., carbon
nanodots, graphene, graphene oxide, and other similar mate-
rials, have also been employed as co-catalysts to facilitate
the electron transfer on g-C3N4. Chai et al. fabricated sandwich-like
graphene-g-C3N4 nanocomposites (GCN) by one-pot
impregnation-thermal reduction.54 The covalent bonds formed
between the graphene and g-C3N4 endow the GCN sample with
a slight redshift of the absorption band edge. The GCN sample
exhibited a 2.3-fold enhancement of CH4 yield over pure g-C3N4
due to the inhibited electron–hole pair recombination and
increased charge transfer.

In another study, 3D porous g-C3N4/carbon nanosheets were
prepared by simple pyrolysis and subsequent carbothermal
activation.55 Gas-phase CO2 photoreduction experiments under
simulated solar irradiation showed that the photocatalytic activity of the 3D g-C3N4 was 2 times that of the bulk g-C3N4 due
to the introduction of carbon and the 3D porous structure. First,
the hierarchical 3D porous architecture promoted CO2 trans-
portation to active sites and accelerated the diffusion of prod-
ucts. Second, the 3D porous structure could have increased the
light-harvesting ability. Furthermore, the photogenerated elec-
trons could have effectively drifted from g-C3N4 to the hybrid
carbon under the action of the inner electric field due to the
quantum confinement effect, which inhibited the electron–hole
pairs from combining and prolonged the lifetime of the pho-
togenerated charge carriers. Finally, according to their X-ray
photoelectron spectroscopy (XPS) analysis, there were partial
electron-withdrawing carbon–oxygen groups on the surface of
this composite, e.g., C–O, C=O, and O–C=O groups, which
could have facilitated the electron flow on the external surface.
Thus, the recombination of electron–hole pairs was suppressed.

Carbon quantum dots (CQDs) doped onto g-C3N4 can also
offer many advantages in CO2 photoreduction, such as reducing
the band-gap, the electron-withdrawing effect, and the up-
conversion effect.56 Recently, Tang et al. reported the hole-
accepting effect of carbon dots in promoting selective CO2
reduction to methanol.57 The microwave-synthesized carbon
dots have a pure graphite structure that facilitates efficient hole
transfer to carbon dots and favours electron accumulation on the
surface of g-C3N4.49 Moreover, the hole-accepting carbon dots
can prevent the surface adsorption of methanol, which subse-
quently prevents the re-oxidation of produced methanol. As
a result, the carbon dot-decorated g-C3N4 produced methanol
from water and CO2 with nearly 100% selectivity to methanol.
This work exhibited the great potential of carbon quantum dots
in efficient and highly selective CO2 photoreduction.

3.3.3 Metal oxide/sulfide/phosphide materials. Apart from carbon
materials and metals, other co-catalysts are emerging, e.g., metal oxide, metal sulfide, and metal phosphide. Ye et al.
loaded chainmail Ni/NiO co-catalyst onto g-C3N4 to increase the
photocatalytic CO2 reduction.58 The NiO shell over the surface of
Ni particles protects Ni from being oxidized to ensure increased
stability and high photoactivity of elemental Ni. Additionally,
the XPS spectra of g-C3N4 and Ni/NiO/g-C3N4 indicated that g-
C3N4 interacted with Ni/NiO via π–π interactions to facilitate charge carrier transfer at the interface, which can also be
demonstrated by photocurrent responses, electrochemical
impedance spectra, and PL spectra. The surface photovoltage
(SPV) spectra and transient photovoltage (TPV) spectra of the
samples indicated that the introduction of Ni/NiO endowed the
catalyst with stronger generation and more efficient separation
of the photogenerated charge carriers. With an adequate
amount of Ni/NiO loading, the CO yield on Ni/NiO/g-C3N4
reached 13.95 μmol g−1 h−1, which was approximately 9-fold
higher than that on pristine g-C3N4.

Zhou et al. loaded molybdenum phosphide (MoP) onto g-
C3N4.59 The introduction of MoP facilitated the separation and
transfer of electron hole pairs, and increased light absorption
throughout the UV and visible light spectrum without changing
the absorption edge of g-C3N4, thus endorsing the photocatalyst
with a 4.5-fold higher production of CO compared with pristine
g-C3N4. Pan et al. prepared NiS2 quantum dot-modified g-C3N4
photocatalysts by a hydrothermal method.60 The lower Fermi
level and surficial metallicity resulted in the ability of NiS2 to
become an ideal electron acceptor, which accelerated the
separation of electron–hole pairs and inhibited their recombi-
nation. Additionally, the introduction of NiS2 quantum dots
provided more active sites and contributed to close interface
contact. Therefore, the CO yield on NiS2/g-C3N4 reached 10.68
μmol g−1 h−1, which was 3.88 times higher than that on pristine
g-C3N4.

Co-catalysts significantly improve the photocatalytic perfor-
mance of g-C3N4 by facilitating the charge carrier transfer and
suppressing the recombination of electron–hole pairs. It is
worth noting that some metal oxides, metal phosphides, and
metal sulfides can form heterojunctions together with g-C3N4,
which is further discussed below.
3.4 **g-C\textsubscript{3}N\textsubscript{4}-based composites**

Combining g-C\textsubscript{3}N\textsubscript{4} with other semiconductors to form heterojunctions can facilitate electron transfer, light utilization, and even CO\textsubscript{2} adsorption. Hence, it is a highly recommended strategy for CO\textsubscript{2} photocatalytic reduction using g-C\textsubscript{3}N\textsubscript{4}.

### 3.4.1 **g-C\textsubscript{3}N\textsubscript{4} absorbent composites.** It is well known that the adsorption/activation of CO\textsubscript{2} is the first step in photocatalytic CO\textsubscript{2} reduction. However, pristine g-C\textsubscript{3}N\textsubscript{4} has a limited surface area and a small number of active sites due to its stacked structure, which impedes the adsorption of g-C\textsubscript{3}N\textsubscript{4}. One strategy to change this is to combine g-C\textsubscript{3}N\textsubscript{4} with CO\textsubscript{2} adsorbents.

Layered double hydroxides (LDHs) are a class of compounds made up of positively charged brucite-like layers with an interlayered region containing charge-compensating anions and solvation molecules (Fig. 7).\textsuperscript{61} By using a general formula \([\text{M}^{2+}\text{M}^{3+}(\text{OH})_2]^x\cdot[A^{m+}]_{n/m}\text{H}_2\text{O} \quad (\text{M} = \text{metal}, A = \text{interlayer anion})\), it was determined that LDHs exhibited some physicochemical properties that are in favour of CO\textsubscript{2} adsorption, e.g., large surface areas, positive surface charges, and compositional flexibilities.\textsuperscript{62} Xu et al. used Mg-Al-LDH nanosheets to enrich CO\textsubscript{2} on the surface of g-C\textsubscript{3}N\textsubscript{4} by the exploitation of its anion-exchange capacity.\textsuperscript{63} LDH/g-C\textsubscript{3}N\textsubscript{4} was assembled via electrostatic interaction combined with Pd co-catalyst, and it showed a remarkably enhanced CH\textsubscript{4} yield compared to the yield without LDH. Moreover, enriched CO\textsubscript{2} existed in the form of interlayered CO\textsubscript{3}\textsuperscript{2−} in the LDH, which could have been more efficiently reduced. Apart from LDH, metal organic frameworks (MOFs) have unique structural properties such as high surface area, high porosity, and low crystal density, as well as high thermal and chemical stability, and thus, they can also be utilized to enhance CO\textsubscript{2} adsorption on g-C\textsubscript{3}N\textsubscript{4}.\textsuperscript{62}

### 3.4.2 **g-C\textsubscript{3}N\textsubscript{4}-based heterojunctions.** From previous studies, it is known that the most frequently used g-C\textsubscript{3}N\textsubscript{4}-based photocatalysts are composites of g-C\textsubscript{3}N\textsubscript{4} with other semiconductors, and are called g-C\textsubscript{3}N\textsubscript{4}-based heterojunctions.\textsuperscript{76}

#### (a) **g-C\textsubscript{3}N\textsubscript{4}-based type II heterojunction.** The heterojunction is a type of p-n junction. It refers to two semiconductor materials that have been combined through surface assembly or internal crystal interface crosslinks, which accelerates the separation of electron–hole pairs by the construction of a built-in electric field.

In terms of the energy band, heterojunctions fall into three categories: (i) type I heterojunction, where the forbidden gap of one component is completely covered by that of the other component; (ii) type II heterojunction, where the forbidden bands of the two components are staggered, and the semiconductor has a more negative CB and a less positive VB; (iii) type III heterojunction, where the forbidden bands of the two components are completely separated. The most investigated photocatalysis systems are usually type II heterojunctions. When a photocatalytic reaction takes place, the photogenerated electrons transfer from the more negative CB to the CB of the other semiconductor and generate a reduction reaction, while the holes transfer from the more positive VB to VB of the other semiconductor and generate an oxidation reaction.

One can construct a type II heterojunction system with g-C\textsubscript{3}N\textsubscript{4} and another semiconductor with a more negative CB position or a more positive VB position (Fig. 8). Commonly, there are few semiconductors whose CB position is more negative than that of g-C\textsubscript{3}N\textsubscript{4}, and thus, it is easy to construct a g-C\textsubscript{3}N\textsubscript{4}-based heterojunction, as illustrated in Fig. 8b. Many metal oxides, sulfides, halides, and other semiconductors are used for such a system.\textsuperscript{21} The construction of the heterostructure plays a significant role in its photocatalytic performance and stability.\textsuperscript{82,83}

#### (b) **g-C\textsubscript{3}N\textsubscript{4}-based Z-scheme heterojunction.** The Z-scheme heterojunction, mimicking the photosynthesis of plants, was first proposed to solve the problem that a single photocatalyst could not split water to simultaneously produce both hydrogen and oxygen. A classic Z-scheme photocatalytic system consists of two photoluminescence processes and a series of redox reactions, and the electron transfer takes place in two optical systems (SI and SII).\textsuperscript{84} Different from type II heterojunctions, the electrons excited on SI transfer to SII through an electronic mediator and combine with holes generated on SII (Fig. 9a). Consequently, the electron–hole pairs generated in each semiconductor are separated, while the high reduction and oxidation abilities of the electrons and holes, respectively, are maintained.\textsuperscript{51,85} This mechanism involved with a mediator is known as the indirect Z-scheme. There is another type of mediator-free charge transfer mechanism, namely, the direct Z-scheme mechanism. In the direct Z-scheme mechanism, the photogenerated electrons on
SI are directly transferred to SII and combined with the holes in SII (Fig. 9b).\textsuperscript{66,67} This results in direct contact between the two semiconductors and accelerates the charge transfer between the two components, which not only increases the reaction efficiency but also reduces the cost.

Li et al. synthesized an indirect Z-scheme BiOI/g-C\textsubscript{3}N\textsubscript{4} heterojunction using a simple deposition method.\textsuperscript{64} The composite at an optimal fraction of 7.4 wt\% BiOI exhibited a significantly enhanced CO yield of 17.9 \textmu mol g\textsuperscript{-1} in a CO\textsubscript{2} photoreduction test. The indirect Z-scheme charge transfer mechanism was confirmed through a series of characterizations including UPS, XPS, and DRS. A contrast test was executed to confirm the roles of intermediate $I_3^-/I^-$ pairs. In another study, an Al–O-bridged g-C\textsubscript{3}N\textsubscript{4}/\alpha-Fe\textsubscript{2}O\textsubscript{3} heterojunction was prepared using a two-step wet chemical method with AlCl\textsubscript{3} aqueous solution as the Al–O source.\textsuperscript{64} The Al–O bridges significantly promoted charge transfer and separation and consequently enhanced the CO\textsubscript{2} photoreduction performance (Fig. 10). The composite exhibited an approximately 4-fold enhancement of CO production (24 \textmu mol g\textsuperscript{-1} h\textsuperscript{-1}). Due to the existence of a charge transfer mediator, this type of Z-scheme heterojunction requires well-designed structures to achieve high charge separation and transfer efficiencies. In contrast, the charge transfers in direct Z-scheme heterojunctions are accelerated by the interface contacts, and there are high reaction efficiencies and low cost with direct Z-scheme heterojunctions. Therefore, there has been increasing interest in direct Z-scheme heterojunctions in recent years.

Wong et al. also fabricated a direct Z-scheme heterojunction consisting of urchin-shaped \alpha-Fe\textsubscript{2}O\textsubscript{3} and a g-C\textsubscript{3}N\textsubscript{4} nanosheet.\textsuperscript{65} Without using any sacrificial agent or co-catalyst, the Z-scheme hybrid had a 2.2-fold higher CO evolution rate as compared to pristine g-C\textsubscript{3}N\textsubscript{4}, consisting of up to 27.2 \textmu mol g\textsuperscript{-1} h\textsuperscript{-1}, which is also higher than that of an Al–O-bridged g-C\textsubscript{3}N\textsubscript{4}/\alpha-Fe\textsubscript{2}O\textsubscript{3} heterojunction. The introduction of 3D \alpha-Fe\textsubscript{2}O\textsubscript{3} not only narrows the band gap, increases the light absorption, and results in a high usage of optical energy, but it also increases the binding energy and CO\textsubscript{2} adsorption so that additional CO\textsubscript{2} molecules can participate in the reaction over a longer time span. Additionally, the direct Z-scheme structure facilitates the electron–hole separation in both \alpha-Fe\textsubscript{2}O\textsubscript{3} and g-C\textsubscript{3}N\textsubscript{4}, thus improving the reduction ability of photoelectrons in the conduction band of g-C\textsubscript{3}N\textsubscript{4}. The improved light harvesting, CO\textsubscript{2} adsorption, and separation of charge carriers all contributed to the improved CO\textsubscript{2} photoreduction performance.

There are many other materials that can be compounded with g-C\textsubscript{3}N\textsubscript{4} to fabricate direct Z-scheme heterojunctions for photocatalytic CO\textsubscript{2} reductions, e.g., ZnO, NiAl-LDH, and Bi\textsubscript{2}S\textsubscript{3}. Most of these are also well designed in terms of structure for further enhancements in CO\textsubscript{2} photoreduction yields.\textsuperscript{66–68} For instance, Ogale et al. synthesized a 2D g-C\textsubscript{3}N\textsubscript{4}/NiAl-LDH heterojunction with strong electrostatic interactions between positively charged NiAl-LDH and negatively charged g-C\textsubscript{3}N\textsubscript{4}.\textsuperscript{66} The enhancement of the photocatalytic activity was mainly ascribed to the excellent interfacial contacts at the 2D/2D interface. The large contact area between the NiAl-LDH and g-C\textsubscript{3}N\textsubscript{4} subsequently suppressed charge carrier recombination and improved the transfer and separation of photogenerated charge carriers. Guo et al. synthesized Bi\textsubscript{2}S\textsubscript{3} quantum dots/g-C\textsubscript{3}N\textsubscript{4} composites, where the Bi\textsubscript{2}S\textsubscript{3} quantum dots were uniformly dispersed on g-C\textsubscript{3}N\textsubscript{4}, and this increased the light absorption and the separation of electron–hole pairs. A series of characterizations demonstrated the Z-scheme mechanism. The heterojunction exhibited 4-fold higher CO yield than the pristine g-C\textsubscript{3}N\textsubscript{4}.\textsuperscript{67}

The photocatalytic mechanism of Z-scheme heterojunctions can be revealed by PL analysis and DFT calculations. Yu et al. prepared a g-C\textsubscript{3}N\textsubscript{4}/ZnO binary composite for photocatalytic CO\textsubscript{2} reduction to CH\textsubscript{3}OH.\textsuperscript{68} The photocatalytic activity exhibited 2.3-fold enhancement compared with pure g-C\textsubscript{3}N\textsubscript{4}. The possible photocatalytic mechanism was verified by carrying out a PL analysis of hydroxyl radicals (’OH) produced on pure g-C\textsubscript{3}N\textsubscript{4}, pure ZnO, and the composite. In terms of thermodynamics, the ’OH and ’O\textsubscript{2} can only be produced by photogenerated holes from the VB of the ZnO and electrons from the CB of the g-C\textsubscript{3}N\textsubscript{4}, respectively. According to the double-transfer mechanism, neither ’OH nor ’O\textsubscript{2} could be produced in the composite system, which implied that it was likely that the charge transfer followed the Z-scheme mechanism. This observation was supported by calculating the charge carrier effective mass and separating tendency of photogenerated electron–hole pairs.
using a DFT calculation. It was found that ZnO has a much higher relative effective mass of electrons and holes \(D = m_{\text{e}}/m_{\text{h}^+}\), indicating that the excited electrons of ZnO have a higher tendency to transfer, and thus, the charge transfer is more likely to originate from the ZnO to g-C3N4 instead of the opposite direction. These findings support the rationality of the Z-scheme mechanism.

### 3.4.3 Photocatalyst based on g-C3N4 and metal complexes.

As mentioned before, compared to traditional semiconductor catalysts, metal complexes possess higher efficiency and selectivity, but poor oxidation ability and recycling ability, which limit their utilization. Therefore, g-C3N4 and metal complexes are combined in some studies, where the metal complex and g-C3N4 act as active sites and light-harvesting units, respectively, to combine their advantages.

Maeda et al. synthesized a heterogeneous photocatalyst consisting of a ruthenium [Ru] complex and g-C3N4. The former served as the catalytic unit and the latter as a light-harvesting unit, respectively. The electronic interactions between the two components enable the electrons to transfer from g-C3N4 into ruthenium. With trans-[Cl]-[Ru[4,4′-(CH2PO3H2)2-2,2′-bipyridine](CO)3Cl2] (RuP) as catalyst and a DMA : TEOA mixture (4 : 1, v/v) as solvent, CO2 was reduced to HCOOH and CO with a selectivity of 80% for HCOOH, and the turnover number (TON) reached 1061 after 24 h of irradiation. The apparent quantum yield was 5.7% at 400 nm, which is higher than the performance reported in earlier studies. Jain et al. combined g-C3N4 with cobalt(II) phthalocyanine tetracarboxylate (CoPc-OOH) in various ratios to form numerous complexes. The CO2 photoreduction activity of the synthesized CoPc-OOH, g-C3N4, and g-C3N4/CoPc-OOH were tested in a water/DMF environment using triethylamine (TEA) as a sacrificial agent. A higher tendency to transfer, and thus, the charge transfer is more likely to originate from the ZnO to g-C3N4 instead of the opposite direction. Therefore, g-C3N4 and metal complexes are combined in some studies, where the metal complex and g-C3N4 act as active sites and light-harvesting units, respectively.

The improved photocatalytic efficiency was ascribed to the higher CO2 concentration resulting from the binding ability of CoPc-OOH to CO2.

For the g-C3N4/metal complex photocatalytic system, the interaction between g-C3N4 and metal complex plays an important role in the enhancement of charge transfer migration and suppression of charge recombination. To maximize the interaction, either π–π interaction between the tri-s-triazine unit of g-C3N4 and organic ligands or hydrogen bonds/covalent bonds between the functional groups can make positive contributions.

### 4. Conclusions

We reviewed the recent progress on g-C3N4-based materials for photocatalytic reduction of CO2. Furthermore, the basic properties of g-C3N4, and the principles and main steps of CO2 photoreduction have been critically elaborated. Additionally, different types of g-C3N4-based composites were compared, e.g., pristine g-C3N4, non-metal doped g-C3N4, g-C3N4 with cocatalysts, g-C3N4-based semiconductors, and g-C3N4/metal complex. Among the g-C3N4-based photocatalysts for CO2 reduction, g-C3N4-based semiconductors are most frequently used and have attracted the most interest. The Z-scheme structure, especially direct Z-scheme heterojunctions, realize charge transfer through interface contact, which accelerates charge transfer and leads to increased reaction efficiency and reduced cost. Hence, direct Z-scheme heterojunctions have attract more interest in recent years. Moreover, combined with other treatments such as elemental doping and structural fabrication, the photocatalytic efficiency of g-C3N4-based materials can be further increased. However, compared to g-C3N4/metal complex systems, a heterogeneous photocatalysis system based on semiconductor photocatalysts exhibits relatively lower utilization of light and depressed photocatalytic efficiency. By improving the interaction between the active sites and light-harvesting unit and optimizing the reaction conditions, g-C3N4/metal complex photocatalysts can increase the conversion efficiency greater than 10 fold, but this can complicate separation and recycling.

In view of the above challenges, significant efforts have been made to combine homogeneous and heterogeneous photocatalysts to obtain high efficiency and high oxidation ability. For the heterogeneous photocatalytic system, strategies were mainly focused on three approaches: (i) tuning the surface properties of g-C3N4 to enhance the adsorption or activation of CO2, (ii) tuning the band structure for more optimal light utilization and photogenerated charge transfer, and (iii) reasonable design of a reaction pathway to accelerate the interface reaction. With laboratory-grade reactions, greater than 90% yield and selectivity of a product could be attained, which is much better than an industrial grade CO2 conversion. Due to financial constraints and industrial scale, these systems are not widely applicable to industrial production. However, future recommended pathways will enable the development of more efficient and sustainable technologies for the g-C3N4 photoreduction of CO2 so that a cleaner environment will exist for future generations.

### Conflicts of interest

There are no conflicts to declare.

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