Recent advances in visible-light graphitic carbon nitride (g-C$_3$N$_4$) photocatalysts for chemical transformations

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Graphitic carbon nitride (g-C$_3$N$_4$) has emerged as a new research hotspot, attracting broad interdisciplinary attention in the form of metal-free and visible-light-responsive photocatalysts in the field of solar energy conversion and environmental remediation. These photocatalysts have evolved as attractive candidates due to their non-toxicity, chemical stability, efficient light absorption capacity in the visible and near-infrared regions, and adaptability as a platform for the fabrication of hybrid materials. This review mainly describes the latest advances in g-C$_3$N$_4$ photocatalysts for chemical transformations. In addition, the typical applications of g-C$_3$N$_4$-based photocatalysts involving organic transformation reactions are discussed (synthesis of heterocycles, hydrosulfonylation, hydration, oxygenation, arylation, coupling reactions, etc.).

Polymeric g-C$_3$N$_4$ contains the earth-abundant carbon and nitrogen elements. Polymeric carbon nitride (PCN) is a highly active photocatalyst. The history of C$_3$N$_4$ polymers and their precursors can be traced back to 1834, where the embryonic form of melon, which is a linear polymer of interconnected tri-triazine through secondary nitrogen, was first developed by Berzelius and named by Liebig. The utilization of g-C$_3$N$_4$ in the field of heterogeneous catalysis started in 2006. Wang and co-workers, in 2009, reported the discovery of g-C$_3$N$_4$ as a metal-free conjugated semiconductor photocatalyst for H$_2$ evolution, potentially shifting the research focus from inorganic to polymeric conjugated semiconductor photocatalysts.

1 Introduction

Currently, photocatalytic technology is considered as one of the most important approaches to address the global energy and environmental issues. Inspired by photosynthesis in plants (Fig. 1), synthetic organic chemists are always trying to develop new organic reactions that can be carried out under green, ecofriendly, and sustainable conditions.
The photocatalytic activities of g-C₃N₄ are greatly influenced by its structure, including its electronic structure, crystal structure, nanostructure and/or heterostructure. In recent years, several synthetic techniques and effective alternative processes have been suggested for the optimization of the photoactivity of g-C₃N₄ with respect to nanostructure design, electronic structure modulation, crystal structure engineering and heterostructure construction. Both doping and copolymerization have been considered to be an efficient technique to incorporate external impurities in g-C₃N₄ for the modulation of its electronic structure and photocatalytic activity. In the context of nanostructure design, nanostructures with controllable morphologies not only facilitate mass transfer in catalysis but also accelerate the collection and separation of electron–hole pairs to drive relevant reactions. Different synthetic pathways have been developed for g-C₃N₄ materials with nanopores and nanoscale geometrical shapes including hard/soft templating strategies, supramolecular preorganization, solvothermal technology and exfoliation strategies. With respect to crystal-structure engineering, ionothermal synthesis reveals great opportunities for the synthesis of highly crystalline, more completely condensed g-C₃N₄ with enhanced photocatalytic activity. The synthesis of hetero-structured composites by coupling g-C₃N₄ with other semiconductors not only suppresses the recombination of photoinduced charges, but also endows the composites with enhanced properties or novel features as a result of their synergistic effects. Accordingly, the enlightened design of its structure at different levels can provide new insights into the creation of high-performance g-C₃N₄ materials for the efficient conversion of solar energy. Polymeric g-C₃N₄ is an inexpensive and high-performance nanomaterial, exhibiting abundant scope for adaptable applications. Polymeric g-C₃N₄ nanocomposites have been extensively explored for photocatalytic degradation, environmental remediation, removal of heavy metals from contaminated water, high-dispersed photocatalyst for phenol mineralization and E. coli disinfection, photocatalytic hydrogen production and water detoxification, photocatalytic water splitting, photo-reduction of CO₂ and efficient removal of tetracycline hydrochloride. Polymeric g-C₃N₄ has been previously applied in the above-mentioned process but its application in the field of organic transformation has been limited. Visible light photocatalysis is a robust emerging field for organic transformation and highly appreciable. The combination of visible light photocatalysis with polymeric g-C₃N₄ has enabled organic chemists to carry out the functionalization of organic compounds. In continuation of our work on photocatalysed organic synthesis, this review mainly describes the latest advances in g-C₃N₄ photocatalysts for chemical transformations and may be particularly useful for those who are already involved in the area of heterocycles and interested in the use of visible light photocatalysis in combination with polymeric g-C₃N₄ for bioactive heterocyclic synthesis.

2 Graphitic carbon nitride photocatalyst

Several methods have been reported to synthesize g-C₃N₄, including chemical vapor deposition, plasma sputtering reaction deposition, solvothermal method, and solid-state reaction. However, these protocols are hindered by their strict conditions and the limited access to expensive instruments. Thus, due to its simplicity, high yield and cost effectiveness, thermal polycrystallization has become the most popular method for the preparation of g-C₃N₄. Given that g-C₃N₄ only contains earth-abundant elements (carbon and nitrogen), it can be prepared with accessible raw materials at low cost. Several inexpensive N-rich precursors with a C–N core structure, such as melamine, cyanamide, dicyandiamide, thiourea and mixtures thereof, have been widely used to prepare g-C₃N₄ via the thermal polycrystallization method. In the facile annealing process, the N-rich molecules are polymerized at the temperature of 450–660 °C to form g-C₃N₄. Yan et al. performed a thermogravimetric–differential scanning calorimetry (TGA–DSC) analysis to investigate the condensation processes of melamine. They observed that the sublimation and condensation of melamine occur at 297–390 °C, and the subsequent deamination and decomposition of the material happen at a higher temperature of 545–630 °C. The g-C₃N₄ synthesized from urea normally has a higher specific surface area than that synthesized from melamine, and g-C₃N₄ obtained from urea shows a larger bandgap than that prepared from thiourea. Zhang and co-workers studied the effect of annealing temperature on the properties of the obtained g-C₃N₄. They found that an increase of the temperature in the range of 450–600 °C could be favorable for the polycrystallization of g-C₃N₄, and thus could improve the electron delocalization in its aromatic structure. However, a further increase in the temperature to over 650 °C could result in the decomposition of g-C₃N₄ into small particles with increased bandgaps. Therefore, the annealing temperature is a key parameter in the synthesis of g-C₃N₄ nanosheets.

With a graphite-like layered structure, g-C₃N₄ exists in the form of 2D nanosheets based on s-triazine (Fig. 2b) or tri-s-triazine (Fig. 2b) tectonic units interconnected by tertiary amines. In this case, given that the nitride pore size and the electronic environment of the N atom are different, the energetic stability of the allotropes is different. Among the allotropes of C₃N₄, tri-s-triazine-based g-C₃N₄ is the most stable under ambient conditions. Benefiting from the aromatic C–N...
heterocycles and their high condensation, g-C₃N₄ with tri-s-triazine rings is thermally stable at a temperature of up to 600 °C in the air, as confirmed by thermogravimetric analysis. Furthermore, the strong interlayer van der Waals interactions render g-C₃N₄ chemically stable, which can resist many organic solvents, as well as acid and alkaline solutions. The 2D g-C₃N₄ nanosheets also provide a large specific surface area due to their graphite-like layered structure, with a theoretical value for monolayer g-C₃N₄ of up to 2500 m² g⁻¹, which is favorable for the adsorption of reactants in solution, and thus can facilitate surface reactions, including photocatalysis.

The presence of sp²-hybridized C and N results in the unique established π-conjugated electronic structures of g-C₃N₄. Wang et al. performed density functional theory (DFT) calculations, which revealed that the valence band (VB) maximum and the conduction band (CB) minimum consist predominantly of the N and C pz orbitals of g-C₃N₄, respectively, which has a direct bandgap of ~2.7 eV. As depicted in Fig. 2c, the bandgap of g-C₃N₄ is around 2.7 eV, enabling the harvesting of visible light up to 460 nm. The conduction band (CB) position of g-C₃N₄ is approximately −1.1 eV and its valence band (VB) position is about +1.6 eV with reference to the normal hydrogen electrode, thereby meeting the requirements for photocatalytic water splitting to produce O₂ and H₂, respectively. Compared with TiO₂, which has a bandgap of 3.0–3.2 eV and can only respond to UV light, g-C₃N₄ possesses more suitable band positions to enable water splitting and shows photocatalytic activity in the visible-light region. For the first time, in 2009, g-C₃N₄ was used as a metal-free photocatalyst for water splitting to generate H₂ under visible-light irradiation. Since this pioneering work and because of its aforementioned intriguing properties, g-C₃N₄ has been attracting significant interest from the scientific community and has drawn worldwide attention in recent years. Furthermore, g-C₃N₄ exhibits photocatalytic activity for CO₂ reduction, pollutant degradation, and photocatalytic degradation via photoredox processes (Fig. 2d).

The design of g-C₃N₄-based materials includes zero-dimensional quantum dots (QDs); one-dimensional (1D) nanowires, nanorods, nanotubes, and nanobelts; two-dimensional (2D) nanosheets and nanoplates; and 3D microspheres, microflowers, sandwich-like structures, and porous structures. For instance, combining theoretical calculation results with practical experimental findings, the structure and electronic and optical properties of modified g-C₃N₄ QDs were systematically and comprehensively explored by Bandyopadhyay et al. The preparation method is relatively simple for single-component 3D g-C₃N₄. For example, Chen et al. designed a bottom-up supramolecular self-assembly pathway to synthesize 3D porous g-C₃N₄ assembled from highly crystalline and ultrathin porous g-C₃N₄, respectively, which has a high crystallinity and overall photocatalytic water splitting activity. The unique electronic structure of g-C₃N₄-based photocatalysts has prompted their applications in selective organic transformations, including oxidation, reduction, and coupling reactions. Given that these organic transformation reactions play a key role in industrial and fine chemical synthesis, it is very attractive for nontoxic g-C₃N₄-based photocatalysts driven by visible light to accomplish the corresponding reactions under mild conditions.

3 Functionalization of g-C₃N₄

3.1 Elemental doping of g-C₃N₄

Elemental doping involves the incorporation of heteroatoms or impurities (metal or nonmetal elements) in g-C₃N₄ to regulate...
its absorption, electronic, and physicochemical properties. There are two main types of elemental doping, i.e., nonmetal doping and metal doping. In the field of photocatalysis, bandgap engineering of g-C\textsubscript{3}N\textsubscript{4} via the incorporation of anions and cations or the co-doping of both plays a predominant role in modulating its light absorption and redox band potentials for targeted photocatalytic applications. Hitherto, numerous studies on the doping of anions such as O\textsuperscript{2−}, P\textsuperscript{5−}, S\textsuperscript{2−}, B\textsuperscript{3+}, I\textsuperscript{−}, P\textsuperscript{−} and some combinations thereof\textsuperscript{89–91} in g-C\textsubscript{3}N\textsubscript{4} have been extensively reported.

Besides metal elements, nonmetal elemental doping is also a promising way to govern the electronic structure of g-C\textsubscript{3}N\textsubscript{4}. Jiang and co-workers\textsuperscript{92} fabricated defective oxygen-doped porous C\textsubscript{2}N\textsubscript{4} via the polymerization of ammonium persulfate and dicyandiamide. Compared to g-C\textsubscript{3}N\textsubscript{4}, the bandgap of the oxygen-doped g-C\textsubscript{3}N\textsubscript{4} increased from 2.70 to 2.75 eV, and its CB decreased from −0.60 to −0.81 V. The more negative CB position indicates more powerful reduction ability, and hence the photocatalytic performance will increase significantly.

### 3.2 Copolymerization of g-C\textsubscript{3}N\textsubscript{4}

The photocatalytic reactivity of g-C\textsubscript{3}N\textsubscript{4} is attributed to its π-conjugated framework because of its sp\textsuperscript{2}-hybridization system. However, the aromatic π system of g-C\textsubscript{3}N\textsubscript{4} in its pristine form generally encounters intrinsic drawbacks, namely, the rapid recombination of charge carriers, inadequate absorption and utilization of sunlight and low surface areas as a result of the defective polymerization of its organic framework, considerably restricting its photochemical functions.\textsuperscript{93−95} Thus, employing the merits of the polymeric π-conjugated system, it is envisaged that the delocalization of the π electrons can be further extended by modifying and anchoring the existing molecular structure of g-C\textsubscript{3}N\textsubscript{4} with other structure-matching aromatic groups or organic additives during the copolymerization process to modulate its intrinsic properties.\textsuperscript{96−100} The copolymerization approach is regarded as typical molecular doping to adjust the conventional π systems, electronic properties, optical absorption, band structure and importantly photocatalytic performance of g-C\textsubscript{3}N\textsubscript{4}.\textsuperscript{101−105}

Che and co-workers\textsuperscript{106} investigated a thermal-conjugate in-plane polymerization approach using glucose as the starting material to integrate \textit{C\textsubscript{ring}} into the triazine units of g-C\textsubscript{3}N\textsubscript{4}. The synthetic route for (\textit{C\textsubscript{ring}})\textsubscript{n}g-C\textsubscript{3}N\textsubscript{4} including the dehydration and polymerization process is shown in Fig. 3a. The successful fusion of \textit{C\textsubscript{ring}} in the C\textsubscript{3}N\textsubscript{4} matrix was confirmed by the TEM and HRTEM analysis (Fig. 3b and c). The lattice fringe in orange ellipses (Fig. 3c) is 0.21 nm, in accordance with the (100) plane of graphite. The enrichment of C (elliptic region) can be observed in the TEM and HRTEM analysis (Fig. 3b and c). The lattice fringe in orange ellipses (Fig. 3c) is 0.21 nm, in accordance with the (100) plane of graphite. The enrichment of C (elliptic region) can be observed in Fig. 3d, further suggesting the presence of \textit{C\textsubscript{ring}}. In addition, the integration of \textit{C\textsubscript{ring}} in the matrix of g-C\textsubscript{3}N\textsubscript{4} can significantly broaden its visible-light absorption, facilitate charge carrier separation, and thus promote its photo-catalytic overall water splitting (OWS) activity. The \textit{C\textsubscript{ring}}–g-C\textsubscript{3}N\textsubscript{4} photocatalyst accelerated the surface reaction kinetics and enhanced the separation of photocarriers in the photoconversion of CO\textsubscript{2} to CH\textsubscript{4} and adsorption and photodegradation of methylene blue\textsuperscript{107} (Fig. 3e).

It is expected that the fusion of conjugated aromatic groups in g-C\textsubscript{3}N\textsubscript{4} units is a favourable route to regulate the electronic features of g-C\textsubscript{3}N\textsubscript{4} to furnish materials with desirable physical and chemical properties, consequently delivering highly efficient photocatalysts. The current research on the copolymerization of co-monomer-modified g-C\textsubscript{3}N\textsubscript{4} is expected to offer a new path for the sustainable conversion of solar energy due to its fascinating novel properties.

### 3.3 Crystalline g-C\textsubscript{3}N\textsubscript{4}

g-C\textsubscript{3}N\textsubscript{4} derived via direct polymerization exhibits an amorphous or hypocrysalline structure due to the incomplete polymerization of the precursors, resulting in a relatively poor photocatalytic performance. Due to kinetic hindrance problem in the solid-state route, the formation of highly crystalline g-C\textsubscript{3}N\textsubscript{4} is a major challenge. However, the insertion of suitable metal salts can enhance the mass-transfer efficiency and improve the crystallinity, thus enabling a new route to synthesize highly crystalline g-C\textsubscript{3}N\textsubscript{4}. Bhunia and co-workers\textsuperscript{108} demonstrated that highly crystalline g-C\textsubscript{3}N\textsubscript{4} (PTF-0.13) can be synthesized via the polymerization of melamene, 2,4,6-triaminopyrimidine, and a mixture of LiCl and KCl in air at 550 °C.

### 3.4 Surface-engineered g-C\textsubscript{3}N\textsubscript{4}

Surface engineering of the interface of g-C\textsubscript{3}N\textsubscript{4} can enhance the amount of adsorption sites by increasing the absorption of protons and inhibit carrier recombination, ultimately resulting in an improved photocatalytic performance. Yu and co-workers\textsuperscript{109} proposed a surface engineering strategy to gradually dope graphic carbon rings in g-C\textsubscript{3}N\textsubscript{4}. By changing the amount of 1,3,5-cyclohexanetriol, graphic carbon rings were incorporated at different depths in g-C\textsubscript{3}N\textsubscript{4}. Gradual surface doping furnished g-C\textsubscript{3}N\textsubscript{4} with suitable electronic band structures and built-in electric fields, resulting in accelerated carrier separation and improved photocatalytic performance. In another case, Wang’s group\textsuperscript{110} fabricated proton- and –OH-modified g-C\textsubscript{3}N\textsubscript{4} spheres and evaluated the effects of the introduction of protons and –OH on the photocatalytic degradation and HER activity. The results suggested that proton functionalization has a lower influence on the photocatalytic hydrogen process, while –OH can dramatically enhance the photocatalytic hydrogen production rate.

### 3.5 Single-atom engineering of g-C\textsubscript{3}N\textsubscript{4}

In the conversion of solar energy, single-atom photocatalysts are of tremendous importance due to their maximum utilization of metal active sites, high catalytic activity, and selectivity. Accordingly, g-C\textsubscript{3}N\textsubscript{4} is an excellent support for single-metal atoms (SMAs) because the lone electron pairs present in the N atoms can coordinate with SMAs. To date, various SMAs such as Pt\textsuperscript{110−112}, Ni\textsuperscript{113} and Cu\textsuperscript{114,115} have been investigated for the photocatalytic HER. Zhou and co-workers\textsuperscript{116} demonstrated a low-temperature in situ photocatalytic reduction approach to synthesize a high-density Pt single atom (Pt\textsubscript{ISA}/g-C\textsubscript{3}N\textsubscript{4}). Zhang and co-workers\textsuperscript{117} selected single-atom Pt/g-C\textsubscript{3}N\textsubscript{4} nanorods (S-Pt–g-C\textsubscript{3}N\textsubscript{4}) as a model material and employed synchronous
illumination X-ray photoelectron spectroscopy to monitor the bond evolution process (Fig. 4).

4  g-C₃N₄-based heterojunction photocatalysts

g-C₃N₄ exhibits a bandgap energy, also known as the void region, covering the top of the filled VB to the bottom of the vacant CB. Initially, the absorption of a photon with energy equal to or greater than the bandgap results in the excitation of an electron from the VB to CB, leaving behind an empty state, which constitutes a positive hole.¹¹⁶,¹¹⁷ Once spatially separated, the charge carriers in the excited states, which then migrate to the surface of g-C₃N₄, are responsible for initiating the reduction and oxidation processes for the photocatalytic conversion of reactant molecules (Fig. 5).

The location of the reactive sites may be either on the g-C₃N₄ surface, where the photoexcitation occurs, or across the interface of another semiconductor or cocatalyst.¹¹¹ In this case, the main role of g-C₃N₄ is to absorb light, produce electron–hole pairs and transfer them to its surface or to a cocatalyst. Indeed, g-C₃N₄ is chemically active only when the photo-induced electron–hole pair is consumed simultaneously before recombination occurs in a fraction of a nanosecond.¹¹⁹–¹²¹

The excitation of the bandgap upon the absorption of light and the formation of electron–hole pairs can also be rapidly accompanied by a few deexcitation pathways (Fig. 5). It is noteworthy that the successful migration of charge carriers to the acceptor molecules, which results in reduction and oxidation processes (pathways 1 and 2, respectively), also competes with the recombination processes.¹²² For example, the recombination process can take place via two different paths, as follows: (1) on the surface of the particle depicted in pathway 3...
(surface recombination) and (2) in the bulk of g-C₃N₄ illustrated in pathway 4 (volume recombination). These recombination processes are disadvantageous to the photocatalytic efficiency. When the recombination process occurs, the excited electrons revert to the VB, dissipating the energy as heat without reacting with the adsorbed species on the surface of g-C₃N₄. Therefore, various strategies are employed to prevent the recombination of charge carriers in g-C₃N₄ and upgrade its photocatalytic performance.

The construction of heterojunctions is another feasible approach to enhance the properties of g-C₃N₄. Especially, its charge separation efficiencies can be dramatically accelerated via the construction of heterojunctions. According to the component, the heterojunction can be classified into five types, i.e., type II, Z-scheme, S-scheme, g-C₃N₄/metal, and g-C₃N₄/carbon heterojunctions.

### 4.1 g-C₃N₄-based type II heterojunction

A g-C₃N₄-based type II heterojunction can be formed with TiO₂, ZnO, WO₃, CdSe, ZnIn₂S₄, BiVO₄, Ag₃PO₄, and many others. Xiao and co-workers proposed an in situ exfoliation and conversion strategy to synthesize a holey C₃N₄ nanosheet/TiO₂ nanocrystal heterojunction (Fig. 6a–e). In type-II heterojunctions, the CB and VB of semiconductor A are higher than that of semiconductor B and their different chemical potentials show band bending at the junction. Type-II heterojunctions are less effective given that they provide low reduction and oxidation potentials, and the transfer of charges is difficult due to electrostatic repulsion between similar charges and their quantum efficiency has been found to be low. For the type-II heterostructure, reduction and oxidation reactions occur at semiconductor B with lower reduction ability, while semiconductor A has low oxidation ability. Consequently, the redox potential of type-II heterojunctions significantly decreases. The electrons in the CB of semiconductor B and holes in the VB of semiconductor A migrate owing to the electrostatic repulsion between the hole–hole and electron–electron pairs. A novel type of structure named Z-scheme, which has a new charge migration mechanism, has been developed to overcome the drawbacks of type-II heterojunctions.
4.2 g-C₃N₄-based Z-scheme heterojunction

In the traditional type II heterojunctions, electrons tend to inject in the CB with mild reduction ability, while holes tend to inject in the VB with weak oxidation ability because of the thermodynamic effect. Therefore, after their separation and transfer, both the reduction and oxidation ability would be weakened. Accordingly, the exploration of heterojunctions that simultaneously promote carrier separation and maintain the oxidation and reduction ability is of practical importance. In principle, the transport direction of the electrons and holes of green plants are opposite to the traditional type II heterojunction, which is named Z-scheme heterojunction. Z-scheme systems can be divided into three types, i.e., traditional Z-scheme, all-solid-state Z-scheme, and direct Z-scheme. Their detailed differences can be seen in ref. 131. The Z-scheme photocatalytic system, as a new type of heterostructure, can improve the separation of electron–hole pairs and enhance the redox ability compared with the corresponding pristine materials, resulting in wide opportunities in various application areas, especially in terms of tremendous potential in improving the photocatalytic reaction activity.

Fig. 6  (a) PL spectra (360 nm excitation), (b) time-resolved fluorescence spectra, (c) Nyquist plots from electrochemical impedance spectroscopy, and (d) transient photocurrent response of TiO₂, g-C₃N₄, g-C₃N₄/TiO₂, and holey C₃N₄/TiO₂. (e) Proposed synergistic effect of surface reactions and charge separation in holey C₃N₄/TiO₂. Adapted with permission from ref. 124a. Copyright 2020, Elsevier. Z-scheme mechanism for the photocatalytic oxidation of 5-HMF to DFF over WO₃/g-C₃N₄. Adapted with permission from ref. 132b. Copyright 2019, Elsevier.

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of g-C₃N₄/RGO/Fe₂O₃ suggested the inhibited recombination of carriers. Therefore, g-C₃N₄/RGO/Fe₂O₃ exhibited efficient solar light-induced OWS activity. Wu et al.³¹ used a WO₃/g-C₃N₄ composite, which was prepared via a direct calcination method with ammonium tungstate hydrate and melamine as raw materials, for the photocatalytic oxidation of 5-hydroxymethylfurfural (5-HMF) to 2,5-diformylfuran (DFF) (Fig. 6f).

4.3 g-C₃N₄-based S-scheme heterojunction

Yu’s group proposed the S-scheme heterojunction because of the great fundamental problems of type II heterojunctions and the complexity and misunderstanding of Z-scheme photocatalysts.¹³¹ Similar to the type II heterojunction, the S-scheme system contains an oxidation photocatalyst and a reduction photocatalyst with staggered band structures. However, the charge transfer route of the S-scheme system is completely different from the type II heterojunction.¹²⁹ The major advantages of S-scheme photocatalysts are the reservation of photo-generated holes and electrons in the VB and CB of the oxidation and reduction photocatalysts. The simultaneous recombination of useless electrons and holes generated by the internal electrical field imparts strong redox ability.¹²⁹ As discussed before, in type II heterojunctions, the redox abilities in both the CB and VB are weakened. In contrast, the strong redox potential is maintained in the S-scheme heterojunction due to the powerful photoinduced electrons and holes accumulated in the CB of the reduction photocatalyst and VB of the oxidation photocatalyst, respectively, with the recombination of the extra carriers. The disadvantages and limitations of S-scheme photocatalysts include the fact that the scope of their application is mainly applicable to powder photocatalysts and they do not apply to

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**Fig. 7** (a) Procedure for the formation of 2D/2D WO₃/g-C₃N₄ heterojunctions by electrostatic interaction. Electrostatic potentials of (b) WO₃ (001) surface and (c) g-C₃N₄ (001) surface. Insets show the structural models of the materials for DFT calculation. EPR spectra of (d) DMPO•°OH in aqueous and (e) DMPO•°O₂− in methanol dispersion in the presence of g-C₃N₄, WO₃ nanosheets and 15% WO₃/g-C₃N₄. Adapted with permission from ref. 134. Copyright 2018, Elsevier. S-scheme α-Fe₂O₃/g-C₃N₄ nanocomposite as heterojunction photocatalyst for the degradation of antibiotics. Adapted with permission from ref. 134b. Copyright 2022, the American Chemical Society.
photoelectrochemistry and solar cells with external circuits. The two selected semiconductors constructing S-scheme photocatalysts are mainly n-type semiconductors and should have suitable band structures and obvious Fermi level differences. For example, Fu et al. employed an electrostatic self-assembly method to design an ultrathin WO3/g-C3N4 S-scheme heterogeneous composite. Cao et al. described that S-scheme Fe3O4/g-C3N4 photocatalysts can degrade commercial cefalexin and amoxicillin (20 mg L−1) under visible light, with five- and nine-times higher performances than that of g-C3N4, respectively. Furthermore, the detailed evidence to propose S-scheme Fe3O4/g-C3N4 heterojunctions and comparison of their photocatalytic performance in the degradation of antibiotics were also included in this protocol (Fig. 7f).

4.4 g-C3N4/metal heterojunction

Many studies have demonstrated that the formation of a g-C3N4/metal heterojunction can effectively enhance the activity of g-C3N4. The mechanism of g-C3N4/metal is different from g-C3N4. When a metal contacts a semiconductor, an energy barrier difference is produced at the interface due to their different work functions and this energy barrier is denoted as the Mott–Schottky energy barrier. The Mott–Schottky energy barrier can greatly promote the electron transfer, and hence promote charge-carrier separation. Specifically, noble metals such as Ag and Au exhibit surface plasmon resonance effects upon visible-light irradiation, which can dramatically promote the light absorption and near field enhancement. However, the incorporation of noble metals increases the expense, and thus the exploration of green and low-cost systems is of significant importance.

5 Synthetic applications of g-C3N4-based photocatalysts

5.1 g-C3N4-catalyzed reactions of quinoxalin-2(1H)-ones with N-aryl glycines under visible light

Yu et al. reported the visible light-induced heterogeneous g-C3N4-catalyzed decarboxylative reaction of the low-cost and easily available quinoxalin-2(1H)-ones with N-aryl glycines as starting materials (Scheme 1). The simplicity of this synthesis and excellent regioselectivity make this protocol a promising tool for building molecules with diverse skeletons and discovering bioactive compounds.

The authors proposed a tentative mechanism for these divergent transformations (Scheme 2). Upon irradiation with visible light, the g-C3N4 semiconductor is excited to generate electrons and holes, which play important roles in organic transformations, acting as single-electron acceptors and donors, respectively. Initially, radical intermediate A is formed through a photooxidation process by releasing a CO2 molecule and proton. Then, radical A is added to the C==N bond of 1 to afford intermediate C. Then, intermediate C undergoes a single-electron photo-reduction to generate anion D. Then intermediate D can be converted into product 3 after protonation. Alternatively, the oxidation of radical A affords iminium species B. The combination of iminium species B and anion D gives intermediate E, followed by intramolecular cyclization to produce product 4 by releasing ArNH2.
5.2 Synthesis of thiocyanated heterocycles

B. Yu and co-workers reported a metal-free photocatalytic strategy for the preparation of thiocyanated heterocycles from the inexpensive NH4SCN using carbon nitride (g-C3N4) as a general heterogeneous catalyst in a green solvent under an air atmosphere and irradiation from a blue LED (Scheme 3). This general method showed advantages such as having a metal-free procedure and an external-oxidant-free procedure, utilizing a recyclable catalyst and a green solvent, wide applicability, applied at room temperature, and easy to scale up.

They proposed the mechanism for this visible light-promoted reaction, which was catalyzed by g-C3N4 (Scheme 4).

Firstly, under the irradiation of visible light, g-C3N4 absorbs photons and generates holes in its valence band (VB) and electrons in its conduction band (CB). Then, the holes generated in the VB ($E_{VB} = +1.2$ V vs. SCE) obtain electrons from NH4SCN via single-electron transfer (SET) to generate the SCN radical. Simultaneously, the electrons in the CB are transferred to O2 (air) through SET to generate O2•−. Then, the SCN radical is added to the C=C bonds of 5a to generate radical intermediate A, which is then subjected to intramolecular cyclization to obtain radical intermediate B. Furthermore, B is oxidized by O2 (air) to form carbocation C through an SET process. Finally, C is transformed into the target product 9a by deprotonation.

5.3 Selective oxidative cleavage of C=C bonds in aryl olefins

S. Das and co-workers reported the use of polymeric carbon nitride (PCN) as a photocatalyst for the selective cleavage of C=C bonds in aryl olefins. This method was selective, scalable in the presence of sunlight and the photocatalyst was easily recyclable. Excellent substrate scope with high selectivity is the main advantage of this recyclable catalyst. In addition, they could perform late-stage C=C bond cleavage in aryl olefins and degradation of complex molecules to achieve highly valuable products and pharmaceuticals (Scheme 5).

5.4 Aerobic benzylic C–H oxygenations

Y. Cai and co-workers reported a metal-free heterogeneous photocatalytic system for highly efficient benzylic C–H oxygenations using oxygen as an oxidant. This visible light-mediated
The oxidation reaction utilized graphitic carbon nitride (g-C3N4) as a recyclable, nontoxic and low cost photocatalyst. The mild reaction conditions allowed the generation of synthetically and biologically valuable materials from readily accessible alkyl aromatic precursors in good yields (Scheme 6).

The probable reaction mechanism was proposed by the authors (Scheme 7). Due to the structural and electronic properties of g-C3N4 with a band gap of 2.7 eV, visible-light excitation leads to the efficient separation of photogenerated electron hole pairs. The photoelectrons (CBM of g-C3N4 located at 0.88 V vs. RHE) can activate O2 to the superoxide radical anion O2\(^{−}\) (reduction potential at 0.56 V vs. SCE). The photogenerated hole of g-C3N4 further undergoes single-electron transfer (SET) with the substrate including isochromans, benzylarenes and N-acyl-derived tetrahydroisoquinolines or isoindolines to afford benzylic radical A. Regarding the substrate N-aryl-derived tetrahydroisoquinolines or isoindolines, SET leads to the formation of the corresponding nitrogen radical cation B, followed by base-promoted deprotonation to generate benzylic radical C. Furthermore, A and C react with the peroxide radical \(\cdot\)OOH from superoxide anions to generate intermediates E and F. Finally, they transform into the products with the simultaneous formation of water as the side product.

5.5 **Aerobic cyclization of thiobenzanilides**

C. Y. Zhou and co-workers\(^{163}\) reported the use of a metal-free heterogeneous photocatalyst for the synthesis of benzothiazoles via intramolecular C–H functionalization/C–S bond formation of thiobenzanilides from benzothioamide 17 by the inexpensive graphitic carbon nitride (g-C3N4) under visible light irradiation. This reaction provides access to a broad range of 2-substituted benzothiazoles 18 in high yields under an air atmosphere at room temperature (Scheme 8).

The plausible reaction mechanism was proposed by the authors (Scheme 9). Initially, N-substituted benzothioamide 17 adsorbs on the surface of g-C3N4 to form complex A with the formation of a new donor level above the valence band of g-C3N4. Visible-light irradiation of A drives the promotion of an electron to the conduction band of g-C3N4 with the generation of radical cation B. The photogenerated electron quenching by O2 and loss of a proton from B produce radical C. The sulfur-centered radical of C intramolecularly attacks the benzene ring to form aryl radical D, which releases an electron to the conduction band of g-C3N4 to generate cation E under visible-light irradiation. Then, E undergoes deprotonation and rearomatization to provide the final product 18.
5.6 Graphitic carbon nitride for α-aminoalkyl radical additions, allylations, and heteroarylations

M. Rueping and co-workers\textsuperscript{164} reported a protocol for the photooxidative activation of α-silylamines and α-amino acids for desilylative and decarboxylative additions, allylations, and heteroarylations in the presence of graphitic carbon nitride (g-C\textsubscript{3}N\textsubscript{4}). This procedure has broad scope and provides the desired products in high yields (Scheme 10). In this protocol, the heterogeneous nature of the reaction system enables the recovery and reuse of the catalyst without loss in its reactivity. The reaction can also be conducted in a continuous flow fashion, even on the gram-scale using a simple and reusable continuous-flow photoreactor, which illustrates the practicability of this heterogeneous photocatalysis protocol.

5.7 Meerwein hydration reaction of alkenes

T. Niu and co-workers\textsuperscript{165} reported a green and efficient visible light-induced Meerwein hydration reaction of alkenes in aqueous medium using mpg-C\textsubscript{3}N\textsubscript{4} as a recyclable photocatalyst. This protocol provides a direct approach for the preparation of racemic alcohols via a free radical mechanism (Scheme 11). In this reaction, water acts as both a solvent and reagent without any additives or co-solvents.

The plausible reaction mechanism proposed by the authors (Scheme 12). Initially, mpg-C\textsubscript{3}N\textsubscript{4} is excited upon irradiation with visible light, which undergoes single-electron transfer (SET) to generate aryl radical A from diazonium salt 23. Adding A to alkene 24 generates the corresponding benzyl radical B, which is oxidized by the valence band of mpg-C\textsubscript{3}N\textsubscript{4} to form intermediate C. Then, H\textsubscript{2}O as a nucleophilic reagent attacks intermediate C, followed by dehydrogenation to afford the corresponding product 25.

5.8 Multicomponent hydrosulfonylation of alkynes

T. Niu and co-workers\textsuperscript{166} reported the synthesis of β-keto-sulfones 30 via the facile and efficient graphitic carbon nitride (p-g-C\textsubscript{3}N\textsubscript{4})-photocatalyzed hydrosulfonylation of alkynes with the insertion of sulfur dioxide under aerobic conditions at room temperature. In this protocol, isotope experiments confirmed that the oxygen atom of the products comes from H\textsubscript{2}O, while the O\textsubscript{2} plays an important role in the reaction by quenching the DABCO radical cation (Scheme 13).
The plausible mechanism was proposed by the authors (Scheme 14). Initially, aryldiazonium cation 26 with DABCO(SO$_2$)$_2$ 29 provides complex A through electrostatic interactions. Subsequent homolytic cleavage of an N-S bond and single-electron transfer generate an aryl radical, sulfur dioxide, and radical cation intermediate B$^+$. Then, the aryl radical reacts with sulfur dioxide to give sulfonyl radical C. Meanwhile, electrons and holes are produced upon the excitation of p-g-C$_3$N$_4$ under visible light irradiation. Consequently, an electron reduces molecular oxygen to produce a superoxide radical, which is quenched immediately with DABCO radical cation B$^+$. Simultaneously, the obtained sulfonyl radical C reacts with alkynes 27 to form intermediate D. Subsequent SET oxidation of intermediate D forms alkenyl carbocation E, which undergoes hydrolysis the to produce the desired β-keto sulfones 30.

5.9 Mizoroki–Heck-type reactions and synthesis of 1,4-dicarbonyl compounds

B. König and co-workers$^{167}$ reported the synthesis of 1,4-dicarbonyl compounds and substituted alkenes (Mizoroki–Heck type coupling) starting from secondary and tertiary alkyl halides and vinyl acetate or styrene derivatives using visible-light photocatalysis (Scheme 15). This protocol uses mesoporous graphitic carbon nitride (mpg-CN) as a heterogeneous organic semiconductor photocatalyst and Ni(II) salts as Lewis acid catalysts.

The plausible mechanism was proposed by the authors (Scheme 16). The exact role of the nickel(II) salt is not clear; however, the authors speculated that the nickel salt acts as a Lewis acid to coordinate with the ester group of alkyl bromide 32, and thereby lowers the reduction potential of alkyl bromide 32. Alternatively, it may be possible that it can activate the double bond by coordinating with olefin 31 given that the formation of the product was observed in the presence of other Lewis acids. However, reactions involving non-activated alkyl bromides did not yield any product, which ruled out the typical cross-coupling mechanism. Based on these observations, they believed that mpg-CN under photochemical illumination generates two-dimensional surface redox centers as electron–hole pairs. The photogenerated electron effectively reduces alkyl bromide 32 and generates alkyl radical A. Then, alkyl radical A...
addition with an aryl halide, delivering Ni(II) oxidative addition (Scheme 18). Initially, the Ni(0) complex serves as a light-absorbing antenna undergoing an energy transfer process (EnT) (singlet–triplet band gap of ca. 0.39 eV) to produce electronically excited Ni(II) species VII. Homolysis of the Ni(II)–X bond and HAT followed by a rebound of the resulting carbon-centered radical with VIII generates Ni(n) species IX. Reductive elimination from the electronically excited species X, promoted by EnT with mpg-CN, provides the final product and regenerates the Ni(0) species, thus completing the catalytic cycle.

5.11 Synthesis of phenols from aryl halides
F. Zhang and co-workers<sup>39</sup> reported the synthesis of phenols via the facile and efficient hydroxylation reaction of aryl halides<sup>38</sup> with water through the merger of a heterogeneous organic semiconductor graphitic carbon nitride (g-C<sub>3</sub>N<sub>4</sub>) photocatalyst and a homogeneous nickel(iii) bipyridine organometallic catalyst under visible-light irradiation at room temperature (Scheme 19).

The reaction mechanism was proposed by the authors (Scheme 20). In the proposed mechanism for the NiDabpy/g-C<sub>3</sub>N<sub>4</sub> dual catalytic system, g-C<sub>3</sub>N<sub>4</sub> acts as a photocatalyst and Et<sub>3</sub>N provides electrons. After visible light illumination, g-C<sub>3</sub>N<sub>4</sub> is excited to enable the effective separation of the photogenerated electron–hole pairs. The photogenerated holes can cause the oxidation of Et<sub>3</sub>N to form a cationic radical. Then, the Et<sub>3</sub>N radical undergoes a hydrogen atom transfer process with water to generate a hydroxyl radical. At this stage, the concurrent oxidative addition of aromatic halides to Ni(0) species should deliver the Ni(n) species, which would be trapped by the hydroxyl radical to form an Ni(n) organometallic adduct. Thereafter, reductive elimination would deliver the desired phenols product and Ni(n) species. Finally, the photogenerated electron by semiconductor excitation is utilized for the reduction of the Ni(n) species to Ni(0) species via a single-electron transfer. Then, concurrent oxidative addition of aromatic halides to Ni(0) species delivers the Ni(n) species to complete the Ni catalytic cycle.

5.12 Sonogashira coupling synthesis of ynones
H. Kaur and co-workers<sup>37m</sup> reported the use of copper nanoparticles supported on graphitic carbon nitride (Cu<sub>2</sub>O NPs@ g-
C$_3$N$_4$) as a photoactive catalyst for the synthesis of ynones under mild and sustainable conditions (Scheme 21). This methodology is simple and efficient and allows the reaction to progress more efficiently under visible light at room temperature.

The authors proposed a plausible reaction mechanism for Cu$_2$O NPs/g-C$_3$N$_4$ (Scheme 22). In this mechanism, they proposed that the reaction proceeds via the Csp-H activation of alkyne molecules under visible light irradiation given that Cu$_2$O NP/g-C$_3$N$_4$ has the ability to promote the formation of the copper acetylide complex more effectively. The copper acetylide complex reacts with the preformed iminium ion to form a 2-pyridinyl propargyl amine intermediate. Subsequently, the cycloisomerization of the intermediate affords the desired product.

### 5.13 Anaerobic oxidation of alcohols to ketones

J. Liu and co-workers$^{71}$ reported the use of a platinum nanocluster/graphitic carbon nitride (Pt/g-C$_3$N$_4$) composite...
solid catalyst with photocatalytic anaerobic oxidation function for the highly active and selective transformation of alcohols 43 to ketones 44. The desired products were successfully obtained in good to excellent yield from various functionalized alcohols at room temperature (Scheme 23).

The authors proposed a possible two-electron oxidation mechanism for the photocatalytic anaerobic dehydrogenation of alcohols (Scheme 24). In this mechanism, firstly, holes ($h^+$) on the valence band (VB) and electrons ($e^-$) on the conduction band (CB) are generated under the irradiation of solar light. Then, the Pt-mediated $h^+$ extracts hydrogen from substrate 43 to release $\alpha$-alcohol radicals. Subsequently, the $\alpha$-alcohol radical intermediate is oxidized to afford the corresponding product 44. Simultaneously, the electrons in the CB band transfer to the Pt nanoclusters through the Schottky-junction to drive the formation of $H_2$ molecules. Furthermore, both byproduct 45 and $H_2$ provide evidence for the reaction mechanism for the photocatalytic anaerobic dehydrogenation of alcohols.

5.14 Photocatalytic decarboxylative alkyl/acyl radical addition and reductive dimerization of para-quinone methides

Y. Cai and co-workers$^{172}$ reported a modified carbon nitride-based heterogeneous photocatalytic system for both the decarboxylative addition and reductive dimerization of para-quinone methides (Scheme 25). The potassium-intercalated carbon nitride (CN–K) was facilely prepared via the direct KCl-induced structure remodeling of bulk $g$-C$_3$N$_4$, exhibiting remarkable catalytic activity. The heterogeneous nature and mild reaction
conditions of this protocol allow good catalyst recyclability, broad substrate scope, scale-up in a continuous flow, and applications for the target synthesis.

The authors proposed the mechanism (Scheme 26), in which, firstly, visible-light absorption by CN–K leads to efficient charge separation, yielding two-dimensional surface redox centers as electron–hole pairs. The photo-electrons reduce \( p \)-QMs to the corresponding diaryl methyl radical \( A \) via the PCET process. For the reductive dimerization reaction, radical 1,6-addition of \( A \) to \( p \)-QMs generates radical \( B \). The photogenerated hole of CN–K with a valence band potential of +0.84 eV undergoes single-electron oxidation with HCOONa to form HCOO\(^{-}\), followed by hydrogen atom transfer (HAT) to \( B \), affording the target dimerization products. For decarboxylative 1,6-addition reaction, the photogenerated hole of CN–K undergoes single-electron transfer (SET) with carboxylic acid anions (+0.91 to +1.25 V vs. SCE for benzylic acid anions) to deliver the corresponding alkyl radical \( C \) or acyl radical \( D \). The direct coupling of radical \( A \) with radical \( C \) or \( D \) affords the target 1,6-addition adducts.

**Scheme 26** Proposed reaction mechanism for potassium-modified carbon nitride for heterogeneous photocatalytic decarboxylative alkyl/acyl radical addition and reductive dimerization of \( \text{para-quinone methides} \).

6 Conclusions and perspectives

In the field of photocatalysis, \( \text{g-C}_3\text{N}_4 \)-based photocatalytic materials are considered one of the best candidates in terms of their properties, preparation, and applications. \( \text{g-C}_3\text{N}_4 \)-based photocatalysts have significant advantages and simple strategies can be used to change the properties of these photocatalysts, while their overall structure composition remains unchanged. The outstanding properties of \( \text{g-C}_3\text{N}_4 \)-based photocatalysts make them the preferred form of visible light photocatalysts for a variety of applications compared to other photocatalysts. This review includes the application of \( \text{g-C}_3\text{N}_4 \) via numerous synthetic strategies in an expanded manner. The application of \( \text{g-C}_3\text{N}_4 \) in chemical functionalization via the covalent or noncovalent approach using a wide range of chemical modifiers has been proven to be a useful strategy to amplify and introduce new properties in this photocatalyst. The properties and utilization of \( \text{g-C}_3\text{N}_4 \) will give researchers some new insights, ideas, and momentum, thus enhancing the production of clean energy and their application in efficient visible light-driven catalysts in sustainable development to next-level perspectives and encourage researchers for the future development in this research field. Although significant efforts have already been devoted to the modification of \( \text{g-C}_3\text{N}_4 \) materials and optimizing their photocatalytic activity, the potential of \( \text{g-C}_3\text{N}_4 \) materials is yet to be fully exploited. In summary, we believe that with the great efforts of researchers in synthesis, characterization, and mechanistic understanding, we can finally obtain an ideal photocatalyst that achieves the aim of utilizing \( \text{g-C}_3\text{N}_4 \) to carry out various chemical transformations.

**Conflicts of interest**

“There are no conflicts to declare”.

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