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# Catalytic conversion of CO<sub>2</sub> to chemicals and fuels: the collective thermocatalytic/photocatalytic/electrocatalytic approach with graphitic carbon nitride

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Rapid industrial development and the excessive use of fossil fuels have produced a significantly large volume of CO<sub>2</sub> in the atmosphere. The efficient conversion of CO<sub>2</sub> to useful chemicals and fuels is an important step towards reducing the concentration of CO<sub>2</sub>. To overcome the thermodynamic and kinetic barriers involved in CO<sub>2</sub> activation and conversion, an effective heterogeneous catalyst and a suitable form of energy, such as thermal, photochemical, and electrochemical, are necessary. Considering a wide variety of catalytic materials for CO<sub>2</sub> conversion using all forms of energy resources discussed above, earth-abundant carbon materials have exhibited the capability to catalyze the thermal, photo-assisted, and electrochemically mediated conversion of CO<sub>2</sub>. Among carbon-based materials, graphitic carbon nitride (g-C<sub>3</sub>N<sub>4</sub>) is an inexpensive and sustainable catalyst that can be effectively used for CO<sub>2</sub> conversion. The chemical reactivity and optoelectronic properties of g-C<sub>3</sub>N<sub>4</sub> can be finely tailored for its use as a conventional catalyst or photocatalyst or photoelectrocatalyst. This article aims to summarize the synthetic strategies to prepare various types of g-C<sub>3</sub>N<sub>4</sub> materials as conventional catalysts/photocatalysts/electrocatalysts for the improved conversion of CO<sub>2</sub> to chemicals and fuels. This article reviews the utilization of CO<sub>2</sub> as a feedstock for chemical and fuel synthesis *via* insertion, carboxylation, and reduction reactions. Further, an in-depth understanding of the catalytic mechanism of various reactions discussed in this article will help in the design of a new generation of heterogeneous catalysts for CO<sub>2</sub> conversion. Special emphasis is placed on the various parameters influencing the photocatalytic and electrocatalytic CO<sub>2</sub> reduction. Finally, the underlying challenges and outlook for the future development of catalysts for CO<sub>2</sub> conversion to specialty chemicals and fuels are discussed.

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## 1. Introduction

CO<sub>2</sub> is one of the major variable components in the troposphere that significantly influence human life.<sup>1</sup> A large amount of CO<sub>2</sub> is present in the soil as CaCO<sub>3</sub>. However, only gaseous CO<sub>2</sub> present in the troposphere is responsible for global warming. The effect of carbon dioxide on Earth is being discussed at every forum and international meeting.<sup>2,3</sup> A closer look reveals that several activities by millions of living entities (plants, animals, and human beings) have raised the CO<sub>2</sub> level in the atmosphere by 280 ppm in the last few centuries. However, after the industrial revolution, the CO<sub>2</sub> level in the atmosphere has risen from 280 ppm to 400 ppm and it is expected to reach up to 500 ppm by the 22nd century, which would raise the mean

temperature of the Earth by 1.9 °C.<sup>4</sup> Today, each and every second of our life is dependent on energy resources. The energy dependency can be taken care of with the use of renewable and alternative resources such as wind, water, radioactive, and solar energy, which do not increase the CO<sub>2</sub> level in the atmosphere and provide a CO<sub>2</sub> emission-free environment.<sup>5</sup> In recent times, several alternative energy generation systems, such as fuel cells, are being developed which do not emit CO<sub>2</sub>.<sup>7</sup> However, the H<sub>2</sub> required in fuel cells is produced by steam reforming, which produces a large amount of CO<sub>2</sub> in the atmosphere.<sup>6</sup> Future energy demands and current technologies suggest that it is a long journey and by that time a significant amount of CO<sub>2</sub> will be released into the atmosphere. If the CO<sub>2</sub> concentration increases beyond 600 ppm it would be difficult for human beings to breathe because it is difficult for the human respiratory system to operate in the presence of a greater concentration of CO<sub>2</sub> even though an excess of O<sub>2</sub> is present in the atmosphere. Thus, there is a high priority in developing strategies to reduce the CO<sub>2</sub> concentration in the environment. This can be achieved *via* two

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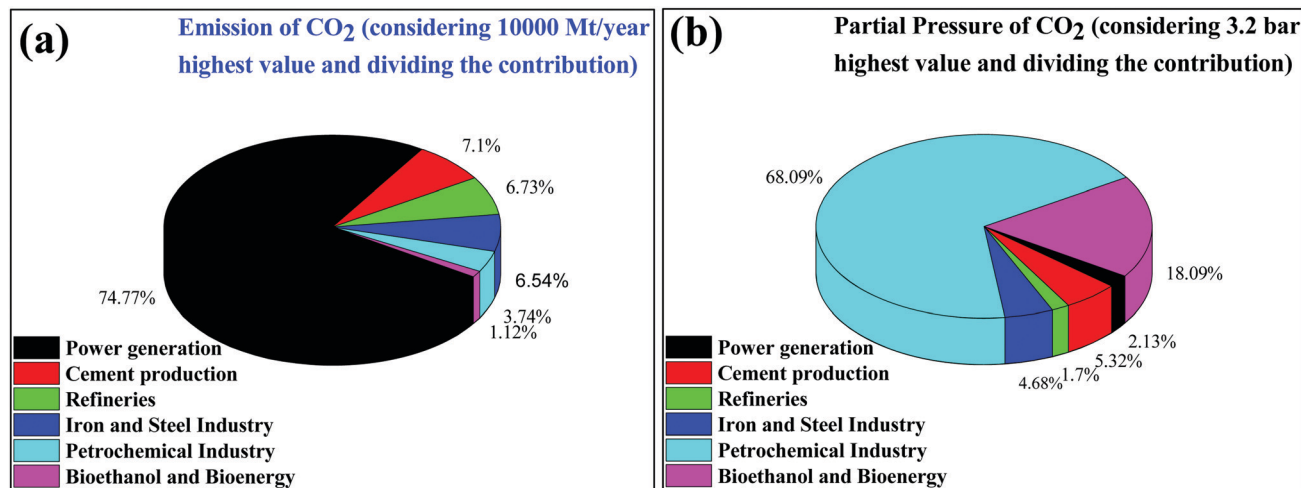


Fig. 1 (a) Emission of CO<sub>2</sub> and (b) partial pressure of CO<sub>2</sub> emitted from different stationary sources.

strategies: (1) CO<sub>2</sub> capture and storage (CCS) and (2) CO<sub>2</sub> capture and utilization (CCU) to useful chemicals and fuels.<sup>7</sup> This article is focused on the CO<sub>2</sub> conversion to useful chemicals and fuels; therefore, only a brief discussion is provided for CO<sub>2</sub> capture and storage.

The various types of stationary plants and their corresponding CO<sub>2</sub> emission (amount and partial pressure) are summarized in pie charts (Fig. 1). It is very easy to capture CO<sub>2</sub> from point sources where the CO<sub>2</sub> concentration is high but it is very difficult to capture CO<sub>2</sub> from low concentration sites, such as on city roads

or houses where the concentration ranges fall at the ppm level.<sup>8</sup> Besides being the most desirable task, direct CO<sub>2</sub> capture and storage are undoubtedly challenging.<sup>9,10</sup> In general, liquid CO<sub>2</sub> scrubbers and metal oxides are used to absorb CO<sub>2</sub> from flue gases.<sup>11–20</sup> Aqueous amine based liquid CO<sub>2</sub> scrubbers are commonly used but their associated energy consumption and cost remain a constraint for further development.<sup>9</sup> Similarly, solid materials such as alkali metal oxides, layered double hydroxides, solid amines, *etc.* are widely explored materials for CO<sub>2</sub> adsorption.<sup>14</sup> However, CO<sub>2</sub> absorption using the CO<sub>2</sub>



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**Rajendra Srivastava**

Dr Rajendra Srivastava joined the Department of Chemistry, IIT Ropar in 2009 as an Assistant Professor, headed the department from March 2015 to May 2018, and is currently working as an Associate Professor. He acquired his Ph.D. degree from CSIR-National Chemical Laboratory, Pune, India where his doctoral work won him the best thesis award of 2006, conferred by the Catalysis Society of India. Afterward, he worked as a

postdoctoral research fellow in KAIST, South Korea (2005–2007) and as a JSPS fellow in the Hokkaido University, Japan (2007–2009). He received the NASI-SCOPUS Young Scientist Award 2017, The Catalysis Society of India Young Scientist Award 2018, and Mid-career Faculty Research and Innovation Award 2019 by IIT Ropar. His fields of interest include designing and synthesis of nanostructured functional materials for catalytic applications, including electrocatalysis (actuators, sensors, supercapacitors and energy harvesters), and photocatalysis. He has published more than 130 peer-reviewed original research articles in reputed international journals and has filed 8 patents.



scrubbers and solid materials discussed above is energetically reversible due to the strong bonding of CO<sub>2</sub> to these materials. Several types of nanoporous materials based on zeolites,<sup>17,18</sup> metal-organic frameworks,<sup>17–19</sup> and carbon<sup>20,21</sup> have been developed for this purpose. All of those above-mentioned materials have functional amines and metal sites for the efficient adsorption of CO<sub>2</sub> over other gases present in flue gas. Apart from tuning the chemical properties, tailoring textural properties such as the surface area, pore size and porous architecture can improve the CO<sub>2</sub> adsorption capabilities. It is noteworthy that materials having a pore size greater than 0.33 nm and lower than 0.35 nm would be ideal for CO<sub>2</sub> adsorption over other gases present in flue gas (the kinetic diameters of CO<sub>2</sub>, O<sub>2</sub>, and N<sub>2</sub> are 0.33 nm, 0.345 nm, and 0.36 nm, respectively).<sup>22</sup> However, such a fine porous material with specific requirements is yet to be developed. Therefore, high surface area materials having dense CO<sub>2</sub> adsorption sites with pore sizes in the range of 0.3 to 2 nm are being prepared.<sup>20,23,24</sup> Thus the direct capture of CO<sub>2</sub> from the air (in which the O<sub>2</sub>/CO<sub>2</sub> ratio is more than 500 or even more) is very demanding but challenging for research scientists and engineers. However, plants do this job very efficiently when the present CO<sub>2</sub> level is < 400 ppm. Plants abstract CO<sub>2</sub> from the atmosphere and convert it to carbohydrate with the aid of sunlight and grow and liberate O<sub>2</sub>.

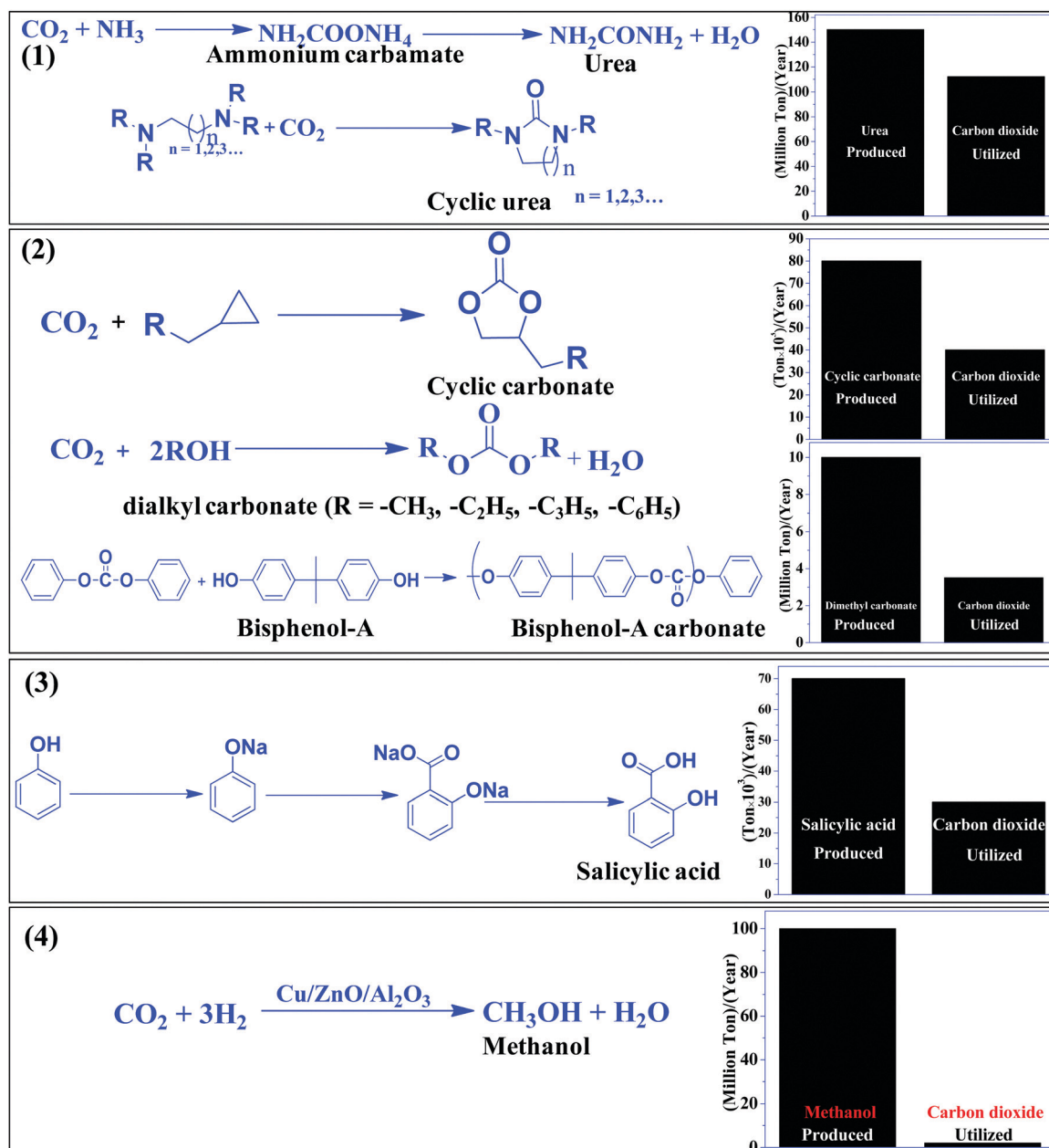
The most efficient strategy to reduce the concentration of CO<sub>2</sub> in the atmosphere is to convert CO<sub>2</sub> to useful chemicals and fuels. CO<sub>2</sub> is a thermodynamically inert molecule with a high C=O bond dissociation energy of 750 kJ mol<sup>−1</sup> which is much higher than the C–O bond (327 kJ mol<sup>−1</sup>) and even higher than the C–C (336 kJ mol<sup>−1</sup>) and C–H (441 kJ mol<sup>−1</sup>) bonds.<sup>25</sup> Due to such high stability, it is difficult to transform CO<sub>2</sub> into useful chemicals. The energy gap between its highest occupied molecular orbital and lowest unoccupied molecular orbital is 13.7 eV and the electron affinity of CO<sub>2</sub> is (−0.6 ± 0.2 eV), which leads to a very negative redox potential for one-electron transfer CO<sub>2</sub> + e<sup>−</sup> → CO<sub>2</sub><sup>−</sup> (*E*<sub>redox</sub><sup>0</sup> = −1.90 eV vs. NHE).<sup>26,27</sup> Such a high energy requirement for electron transfer to generate active species for CO<sub>2</sub> conversion makes photocatalytic and electrocatalytic processes highly challenging tasks. Due to such a high energy requirement, only a few chemicals have been commercialized, which utilizes a large amount of CO<sub>2</sub> to give useful chemicals such as urea, methanol, salicylic acid, cyclic carbonates, and dimethyl carbonate (Scheme 1). Synthesis of urea and cyclic urea is not a 100% atom economic process because water is formed as a by-product. In an atom economic process, the total number of atoms present in the reactants is incorporated into the product. Synthesis of cyclic carbonates takes place by an insertion reaction in which CO<sub>2</sub> is inserted into a three-membered epoxide ring to produce a five-membered cyclic carbonate *via* a 100% atom economic process.<sup>28,29</sup> Dimethyl carbonate (DMC), a solvent, can be produced by the reaction of methanol and CO<sub>2</sub>.<sup>30,31</sup> DMC reacts with phenol using a Pb(OPh)<sub>2</sub> catalyst to produce diphenyl carbonate (DPC). Then DPC reacts with bisphenol-A to produce a polycarbonate *via* a melt polymerization method (Scheme 1).<sup>32</sup> Salicylic acid is manufactured

by the reaction of phenol and CO<sub>2</sub> in a strongly basic medium.<sup>33</sup> Another bulk chemical is methanol, which is produced annually at a rate of 100 million tons by utilizing 2 million tons of CO<sub>2</sub>.<sup>34</sup> (Scheme 1). Industrial processes for chemical synthesis using CO<sub>2</sub> as a reactant are operated at high temperature and pressure due to the thermodynamic stability and inertness of CO<sub>2</sub>. Such reactions can be catalyzed under mild conditions if the CO<sub>2</sub> is efficiently activated. Since CO<sub>2</sub> is an acidic molecule, therefore it can be activated using basic catalysts such as alkali and alkaline earth metal-based oxides and hydroxides. Various organic amines (N-heterocyclics, DBU, guanidine, melamine, phthalocyanine, and porphyrins) and polyamines have also been used in chemical synthesis utilizing CO<sub>2</sub> as the reactant. In general, organic amines are more reactive than inorganic bases and thus various ionic liquids, metal-organic frameworks, silica, and zeolite based materials with functional amine groups have been utilized for chemical synthesis involving CO<sub>2</sub> as a reactant. Nitrogen-containing carbon materials, such as carbon nitrides, are a unique class of materials which can be prepared with economical nitrogen and carbon-containing precursors. Since these materials have a large nitrogen content, they can be used for CO<sub>2</sub> activation and adsorption.

In this review, we will present the advances, challenges, and prospects of carbon nitride based catalysts, photocatalysts, and electrocatalysts in CO<sub>2</sub> conversion to chemicals and fuels. After a brief introduction to CO<sub>2</sub> conversion, the requirements of CO<sub>2</sub> storage and conversion, the industrial process for CO<sub>2</sub> to chemicals, and the types of materials required for CO<sub>2</sub> activation and conversion are described in Section 1. Sections 2–6 focus on carbon nitride-based catalysts in the conventional catalytic conversion of CO<sub>2</sub> to chemicals and photocatalytic and electrocatalytic conversion of CO<sub>2</sub> to C<sub>1</sub> fuels. Section 2 presents a brief description of nitrogen-containing carbon-based materials, types of carbon nitride, and the structural elucidation of g-C<sub>3</sub>N<sub>4</sub> for catalytic applications. Section 3 deals with the synthesis strategy adopted for the preparation of g-C<sub>3</sub>N<sub>4</sub> for improved catalysis and photocatalysis involving CO<sub>2</sub> as a reactant. Sections 4 and 5 provide detailed information about photocatalytic and electrocatalytic CO<sub>2</sub> conversion to C<sub>1</sub> fuels, respectively, their basic concepts, the influence of various parameters influencing the CO<sub>2</sub> conversion and C<sub>1</sub> fuel selectivity, deep insight into the reaction mechanisms *via in situ* characterization techniques and product identification, catalytic activity assessment and result presentation, and the experimental set-up required for photocatalytic and electrocatalytic measurements. Finally, this review will be concluded with a summary, underlying challenges, and opportunities for the future development of catalysts for CO<sub>2</sub> conversion to chemicals and fuels.

This focused review aims to concisely cover almost all the important points, starting from the requirements of CO<sub>2</sub> conversion, fundamentals of CO<sub>2</sub> activation, strategies adopted for g-C<sub>3</sub>N<sub>4</sub> preparation for improved CO<sub>2</sub> conversion, structure elucidation of g-C<sub>3</sub>N<sub>4</sub>, and structure–activity relationship, followed by critical operational parameters for collective ‘thermo-photo-electro’-chemical CO<sub>2</sub> conversion focusing on the g-C<sub>3</sub>N<sub>4</sub> catalyst,





Scheme 1 CO<sub>2</sub> based commercial processes and amount of CO<sub>2</sub> utilized vs. product formed in the given process.

which is not available in a single article hitherto reported in the literature.

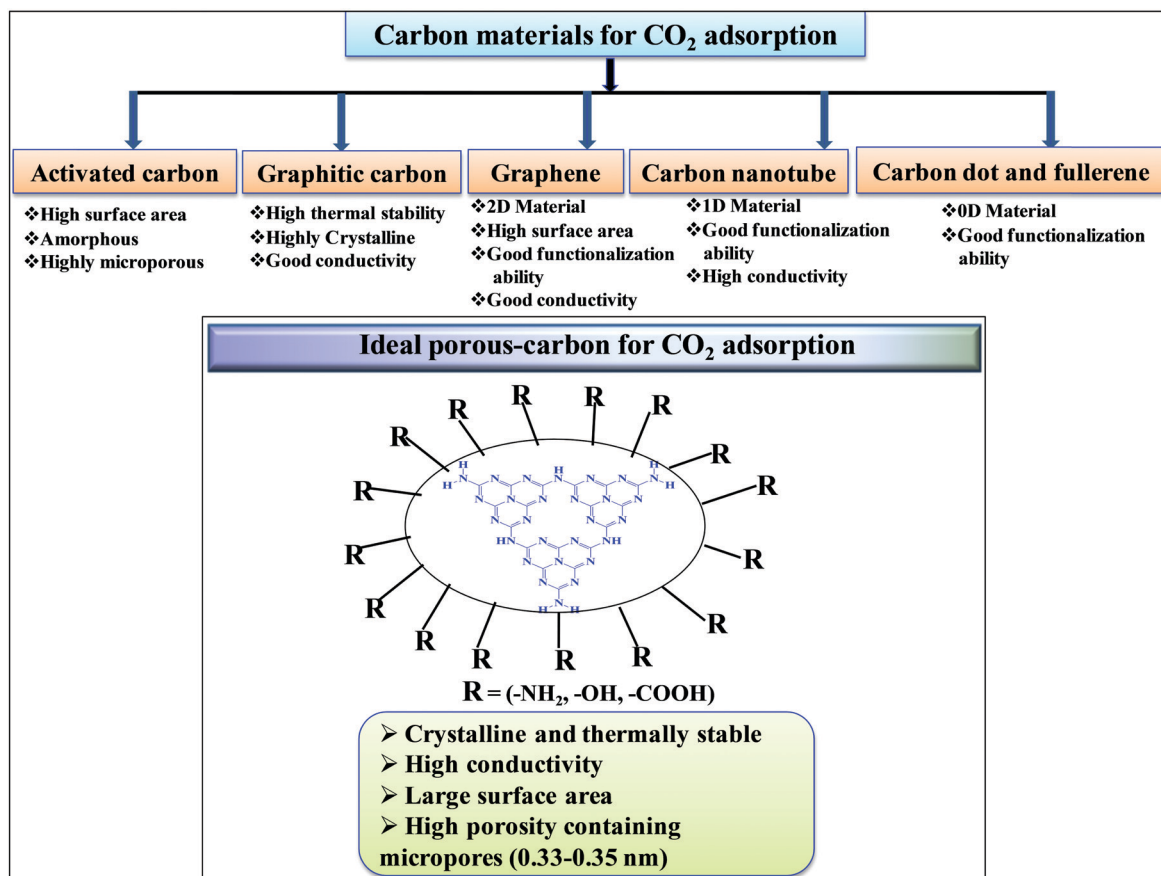
## 2. Carbon nitride, a versatile material for CO<sub>2</sub> adsorption and conversion: structural elucidation for catalysis

Among porous materials, carbon materials have a high surface area, tunable porosity, hydrophobicity for CO<sub>2</sub> adsorption under hydrated conditions, high thermal and chemical stability, and functional ability.<sup>35–37</sup> Activated carbon is the most economical and widely available carbon material having micropores,

mesopores, and macropores.<sup>38</sup> Due to the porous architecture and high surface area, it can significantly adsorb large amounts of CO<sub>2</sub> by the physisorption process but it finds difficulty in the activation and utilization of CO<sub>2</sub> in chemical synthesis due to the inability to chemisorb and activate CO<sub>2</sub>.<sup>39</sup> Biomass pyrolysis and KOH activation can produce high surface area activated carbon, suitable for CO<sub>2</sub> physisorption.<sup>40</sup> N-Doped microporous carbon prepared with coconut shell and urea has exhibited high CO<sub>2</sub> adsorption and CO<sub>2</sub>/N<sub>2</sub> selectivity at 1 bar, confirming that N-doping has improved the CO<sub>2</sub> adsorption.<sup>41</sup> Similar to inorganic porous materials, sol-gel methods have been applied for the synthesis of porous carbon materials in water and/or ethanol medium followed by acid activation



in the presence and absence of metal catalysts. Due to the high surface area and surface to volume ratio, these materials can be easily functionalized using various amines for the efficient adsorption of CO<sub>2</sub>.<sup>50,51</sup> These porous carbons have varied surface area, morphology, and porosity and, thus, the extent of CO<sub>2</sub> adsorption over these materials varies depending on their tunable textural properties. However, the adsorption and chemisorption capability improves if nitrogen-doped or nitrogen-containing carbon materials are considered. Nitrogen-containing carbon materials induce interactive forces between CO<sub>2</sub> and nitrogen groups present in the material. CO<sub>2</sub> is a weak Lewis acidic molecule because the carbon atom in CO<sub>2</sub> is electron-deficient. Nitrogen-doped carbon atoms are negatively charged and nitrogen sites act as electron donors and behave like a Lewis base; therefore, nitrogen-containing carbon materials have acid–base interactions through which CO<sub>2</sub> molecules are chemisorbed. The strong dipolar C=O bond generates an electric quadrupole moment in the CO<sub>2</sub> molecule, whereas nitrogen doping induces local polarization in the porous carbon matrix.<sup>52</sup> Notably, N<sub>2</sub> has a lower electric dipole moment than CO<sub>2</sub>. Hence, the interaction between the CO<sub>2</sub> molecular quadrupole moment and high electrostatic potential of N-doped carbons enhances the CO<sub>2</sub> adsorption energy, leading to selective and higher CO<sub>2</sub> adsorption from flue gases. Experimental and computational studies have shown that surface hydroxyl



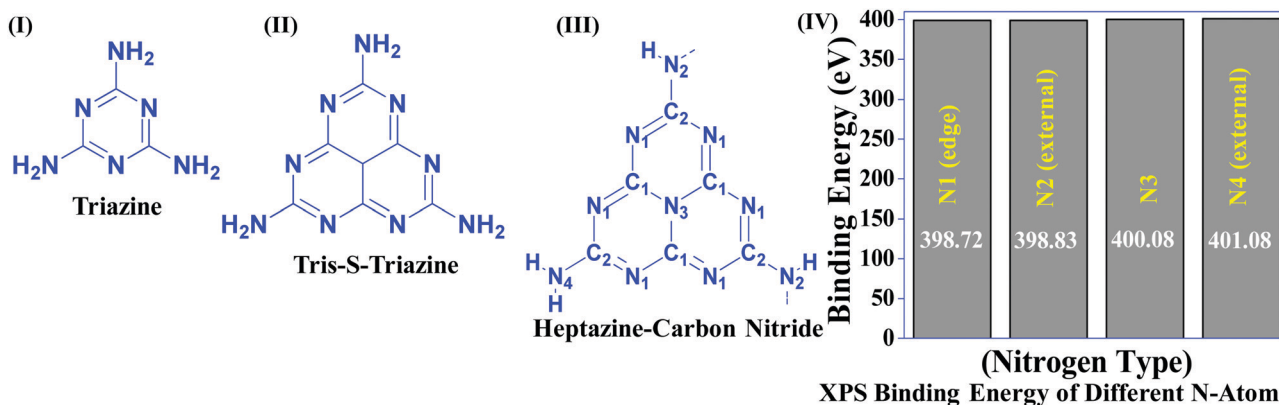
**Table 1** Nitrogen-containing carbon materials and their CO<sub>2</sub> uptake capacity<sup>44</sup>

| E. no. | Material   | $S_{\text{BET}}$<br>(m <sup>2</sup> g <sup>-1</sup> ) | Nitrogen<br>content (%) | CO <sub>2</sub> uptake<br>mmol <sup>-1</sup> g <sup>-1</sup> (298 K) |
|--------|------------|-------------------------------------------------------|-------------------------|----------------------------------------------------------------------|
| 1      | NMC-600    | 272                                                   | 6.33                    | 3.90                                                                 |
| 2      | NMC-700    | 518                                                   | 6.06                    | 3.60                                                                 |
| 3      | AC-600-0.5 | 508                                                   | 6.70                    | 3.00                                                                 |
| 4      | AC-600-2   | 966                                                   | 4.50                    | 3.64                                                                 |
| 5      | AC-800-2   | 2448                                                  | 3.90                    | 3.00                                                                 |
| 6      | Sk-0.5-700 | 1060                                                  | 0.48                    | 4.24                                                                 |
| 7      | SK-4-700   | 2792                                                  | 0.18                    | 3.05                                                                 |
| 8      | PGC-K      | 1143                                                  | 8.30                    | 4.00                                                                 |
| 9      | FCP-1-KC   | 800                                                   | 0.60                    | 5.20                                                                 |

groups present in the N-doped carbon skeleton enhance the interaction between –OH and –NH functionalities leading to high CO<sub>2</sub> adsorption in N-doped carbon materials.<sup>53</sup> Nitrogen doping and surface functionalized amines have different types of interaction. Surface functionalized amines interact with electropositive carbon of CO<sub>2</sub> and form ammonium carbamate leading to high CO<sub>2</sub> adsorption on the surface.<sup>54</sup> However, surface and pore blockage and thermal instability of functionalized amines are some of the demerits of surface amine-functionalized carbon materials; therefore, nitrogen-doped carbon is superior over amine-functionalized carbon materials. Based on the varied textural properties and chemical properties (due to N doping and amine functionalization), it can be concluded that high surface area and high nitrogen contents both are important factors contributing to high CO<sub>2</sub> adsorption (Table 1).<sup>44</sup> Moreover, when CO<sub>2</sub> activation and utilization for chemical synthesis are considered both these factors play important roles.

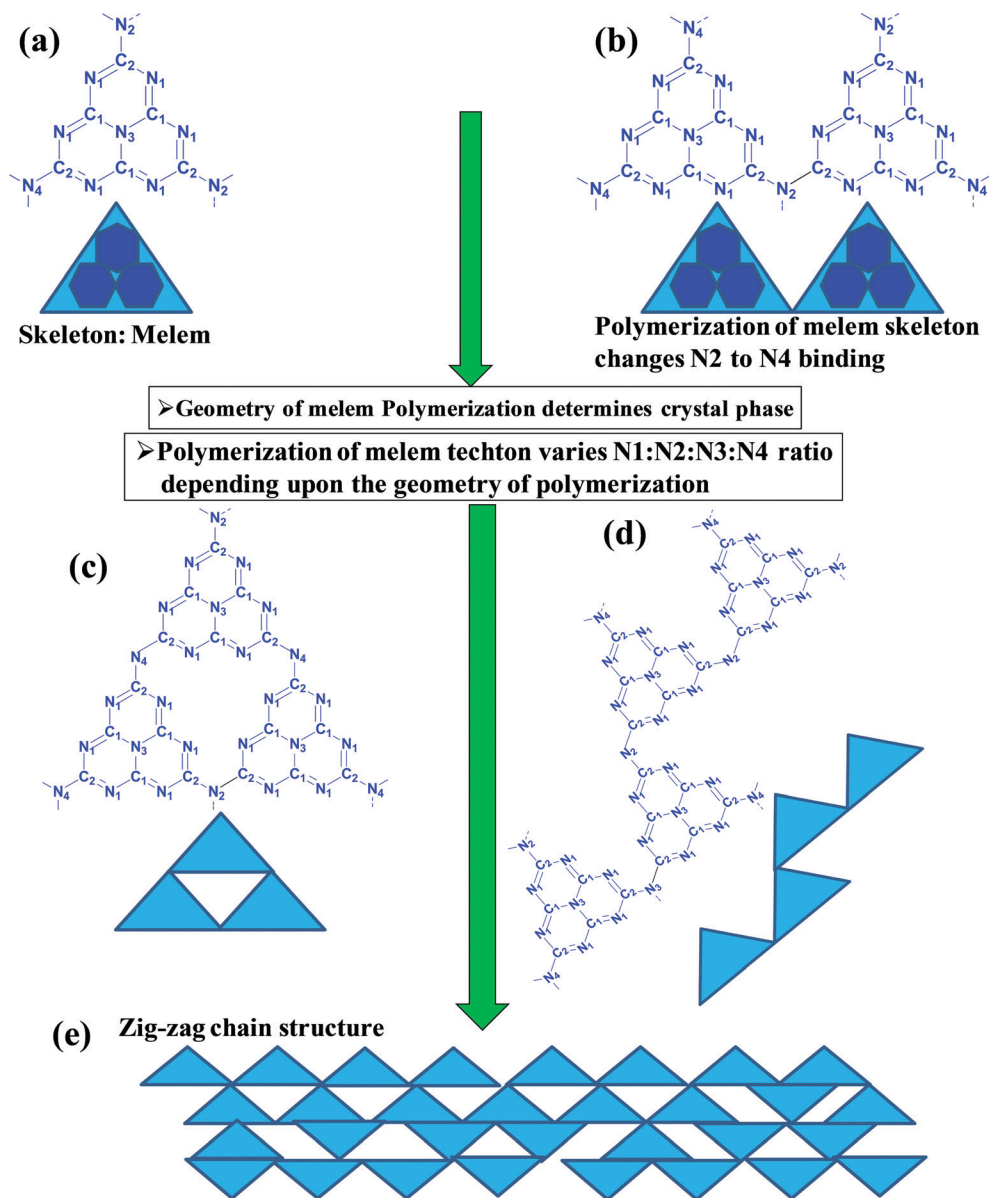
Table 1 suggests that a large number of framework substituted N atoms in a carbon matrix is required for high CO<sub>2</sub> adsorption. The graphitic carbon framework provides high thermal stability and conductivity, which would be good for photocatalysis and electrocatalysis. An ideal porous carbon material and its properties are listed in Scheme 2. Based on these characteristic features, a unique carbonaceous material known as graphitic carbon nitride (g-C<sub>3</sub>N<sub>4</sub>) can best serve this purpose. When a simple nitrogen-rich organic compound (such as urea, melamine, cyanamide, dicyandiamide, *etc.*) is thermally-

polymerized in the temperature range of 500–700 °C, it produces carbon nitride. This material was synthesized way back in 1834 by Berzelius;<sup>55</sup> however its potential as a catalyst<sup>56</sup> and photocatalyst<sup>57</sup> was realized in 2006 and 2009, respectively. This crystalline material exists in seven allotropes:  $\alpha$ -C<sub>3</sub>N<sub>4</sub>,  $\beta$ -C<sub>3</sub>N<sub>4</sub>, cubic C<sub>3</sub>N<sub>4</sub>, pseudocubic C<sub>3</sub>N<sub>4</sub>, *g-h*-triazine, *g-o*-triazine, and *g-h*-heptazine with bandgaps of 5.49, 4.85, 4.30, 4.13, 2.97, 0.93, and 2.88 eV, respectively.<sup>58</sup> These allotropes vary in their nitrogen environment and nitrogen contents. Theoretically, carbon nitride is predicted to have a C/N ratio of 0.75 without any surface functional groups and hydrogen content but the reality is far from this prediction. Carbon nitrides have been actually obtained with varied C/N ratios using different nitrogen-containing organic compounds at varied reaction temperature. It is possible to synthesize carbon nitrides such as C<sub>3</sub>N, C<sub>3</sub>N<sub>2</sub>, C<sub>10</sub>N<sub>3</sub>, C<sub>5</sub>N, C<sub>3</sub>N<sub>5</sub>, C<sub>3</sub>N<sub>6</sub>, C<sub>3</sub>N<sub>7</sub>, *etc.*, having different nitrogen contents than the theoretically predicted ideal C<sub>3</sub>N<sub>4</sub> and those are termed as nitrogen-rich (C/N < 0.75) and nitrogen lean (C/N > 0.75) C<sub>3</sub>N<sub>4</sub>.<sup>59</sup> Whenever a six-membered conducting atomic-scaffold is envisaged, structures such as those of benzene or borazine pop-up in our minds. Based on this perception, the triazine-like structure of C<sub>3</sub>N<sub>4</sub> can be predicted (Scheme 3(I)). But it has been found that triazine or tri-*s*-triazine units (Scheme 3(II)) (also known as tectonic units) propagate in three dimensions to produce allotropes of g-C<sub>3</sub>N<sub>4</sub> (Scheme 3). Computational studies based on Density Functional Theory (DFT) suggest that the tri-*s*-triazine unit is more stable than the triazine unit by 30 J mol<sup>-1</sup>.<sup>60</sup> Later, 2,5,8-triamine, tri-*s*-triazine (C<sub>6</sub>N<sub>10</sub>H<sub>6</sub>) has been isolated, which confirms the building units of g-C<sub>3</sub>N<sub>4</sub>.<sup>61</sup> Theoretical g-C<sub>3</sub>N<sub>4</sub> should have an ideal C/N ratio of 0.75 and should not possess any H or O atoms; however, XPS and elemental analysis suggest that it has H and O atoms in small quantities in addition to C and N. Notably, H and O atoms are observed due to terminal amine groups and surface adsorbed water and (C=O), (O–N), and (C–OH) groups, for which the details are provided in Scheme 3. Various types of C and N are designated in one pictorial representation of a terminal tri-*s*-triazine unit of g-C<sub>3</sub>N<sub>4</sub> (Scheme 3). The structure clearly shows that it has two different types of C represented as C1 and C2 (Scheme 3(III)). Similarly, four different types of

**Scheme 3** Chemical structure of (I) triazine, (II) tris-*s*-triazine, and (III) heptazine units of carbon nitride, and (IV) various N types and their binding energies.

nitrogen atoms are designated as N1 (pyridinic), N2, N3 (graphitic-N), and N4 (Scheme 3(III)). They can be easily distinguished from XPS analysis due to different electronic environments that provide different binding energies (Scheme 3(IV)).<sup>62</sup>

In a true heptazine building unit (also known as melem), N2 is absent and N1:N3:N4 should be present in the ratio 6:1:3 (Scheme 4a). When the melem monomer polymerizes, the N4-type of nitrogens are converted to the N2-type (Scheme 4b).

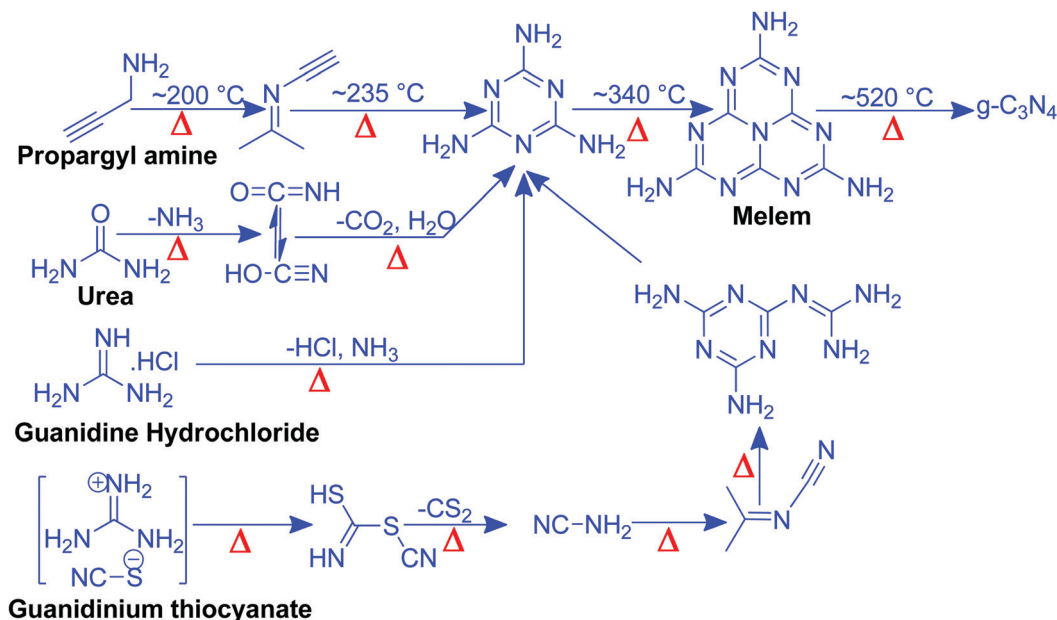


| Sl. No.        | N1 | N2 | N3 | N4  | N1 %  | N2 % | N3 %  | N4 %  |
|----------------|----|----|----|-----|-------|------|-------|-------|
| Dimer          | 12 | 1  | 2  | 4   | 63.16 | 5.26 | 10.52 | 21.05 |
| Tetramer       | 12 | 1  | 2  | 3   | 66.66 | 5.55 | 11.11 | 16.67 |
| Octamer        | 12 | 1  | 2  | 2.5 | 68.85 | 5.71 | 11.43 | 14.28 |
| Infinite chain | 12 | 1  | 2  | 2   | 70.58 | 5.88 | 11.76 | 11.76 |

Scheme 4 Evolution of different geometry and N types present in  $C_3N_4$  based on the variation of the linkages of melem units.<sup>53</sup>







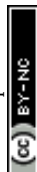
Scheme 6 Synthesis of g-C<sub>3</sub>N<sub>4</sub> via the melem unit using different precursors.

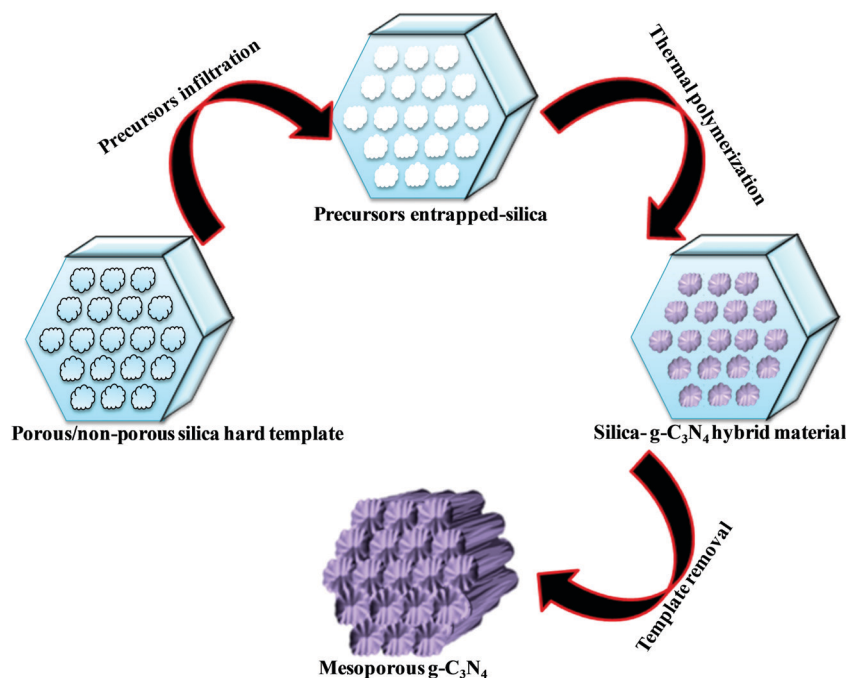
Further, the surface area of urea derived g-C<sub>3</sub>N<sub>4</sub> is improved if the pyrolysis is carried out for a longer time.<sup>74</sup> During such prolonged heating the layered structure of g-C<sub>3</sub>N<sub>4</sub> is exfoliated and splitting takes place to produce low yield g-C<sub>3</sub>N<sub>4</sub> with better surface area and photocatalytic activity.<sup>75</sup> Inspired by the synthetic paradigm of g-C<sub>3</sub>N<sub>4</sub> from urea which has resulted in higher surface area due to the liberation of CO<sub>2</sub> during urea pyrolysis, sucrose and melamine have been pyrolyzed together. The reason for selecting sucrose is that it will generate a large amount of CO<sub>2</sub> during pyrolysis and produce g-C<sub>3</sub>N<sub>4</sub> with N-deficiency and improved surface area (6 fold higher) and porosity.<sup>76</sup> The type of N species and surface area can be tailored by selecting different precursors. For example, guanidine hydrochloride or guanidine thiocyanate pyrolysis results in g-C<sub>3</sub>N<sub>4</sub> *via* a melem intermediate and produced materials with high surface area and enhanced photocatalytic activity (Scheme 6).<sup>77</sup> The bandgap of g-C<sub>3</sub>N<sub>4</sub> can be reduced and mid-gap creation is possible if a nitrogen-vacancy is created in the material. Nitrogen deficiency in g-C<sub>3</sub>N<sub>4</sub> leaves additional electrons and creates nitrogen-vacancy related C<sup>3+</sup> states leading to a reduced bandgap material. This can be easily achieved by heating the material in an H<sub>2</sub> atmosphere. H<sub>2</sub>, being a small molecule, can easily penetrate and diffuse to the interior lattice sites and reduce the lattice N sites and liberate NH<sub>3</sub> and create an N vacancy. In this process, the terminal N sites are more prone to reduction due to the lowest change in the energy of the molecule. Such an N vacancy in the system reduces the bandgap by downshifting of the CB and up-shifting of the VB position. Another situation can also arise in which new C and N sites are generated, which increases the H contents in the material and alters the bandgap of the material. Experimental analysis suggests that a three coordinated N-vacancy and H-substitution at the N sites decrease the bandgap by 0.2 and 0.55 eV.<sup>78</sup> This lowering in the bandgap is due to the formation

of additional band states below the CB by the defect C sites leading to lowering the bandgap values. The N-vacancy moves the VB energy to more negative values and H-substitution lowers the CB to more positive values and improves the visible light absorption capacity. The surface area of g-C<sub>3</sub>N<sub>4</sub> can be improved by exfoliating bulk g-C<sub>3</sub>N<sub>4</sub> in different solvent media such as water, ethanol, acetone, IPA, and NMP. Among these solvents, NMP has been found to be the best choice.<sup>79</sup> However, it is difficult to remove NMP from the exfoliated g-C<sub>3</sub>N<sub>4</sub>. Thus IPA has been explored as the next most desirable solvent to exfoliate bulk g-C<sub>3</sub>N<sub>4</sub>, so as to produce high surface area g-C<sub>3</sub>N<sub>4</sub> of the order of 300–400 m<sup>2</sup> g<sup>−1</sup>. For exfoliation of bulk g-C<sub>3</sub>N<sub>4</sub>, aqueous alkaline and acidic solutions have also been explored with the aim of obtaining higher surface area.<sup>80,81</sup> Although an improvement in the surface area is observed by varying the precursors, reaction temperatures, and pyrolysis environment, however, the materials still remain nonporous in nature. In order to introduce porosity in the material, template-assisted synthetic strategies have also been adopted as discussed below.

### 3.2 Template-mediated synthesis strategy to produce g-C<sub>3</sub>N<sub>4</sub> with high surface area and porosity for improved CO<sub>2</sub> adsorption

Various hard template-mediated synthetic strategies utilize silica-based mesoporous materials (SBA-15, SBA-16, KIT-6, MCM-41, and MCM-48) as the template (Scheme 7). Nitrogen-containing organic precursors or carbon and nitrogen precursors are infiltrated into the porous architecture of silica materials *via* the incipient wet impregnation or wet impregnation method followed by pyrolysis, which results in the formation of g-C<sub>3</sub>N<sub>4</sub>/silica hybrid materials. Finally, the silica template is removed by chemical etching using various etching agents, for example, NaOH, HF, and NH<sub>4</sub>HF<sub>2</sub>. The final material has the same architecture as that of the original silica material.<sup>82</sup>





Scheme 7 Hard template mediated synthesis of mesoporous g-C<sub>3</sub>N<sub>4</sub>.

Even silica nanoparticles have produced porous g-C<sub>3</sub>N<sub>4</sub> with an average pore diameter of 12 nm, which is equivalent to the original silica nanoparticles.<sup>83</sup> In general, the pore diameter of the resulting material is smaller than the porous silica materials due to volume shrinkage of g-C<sub>3</sub>N<sub>4</sub> filled inside the pores during the condensation.<sup>84</sup> The surface area and porosity significantly depend on the Si/precursor, hard template, and etching agent. The surface area can be further improved if the original silica material is treated with acid to enhance the surface reactivity, which allows a large number of organic precursors to in-filter, which improves the surface area of the resultant material.<sup>85</sup> This synthetic strategy produces a low yield of g-C<sub>3</sub>N<sub>4</sub> and, therefore, is difficult for application in the large scale synthesis of mesoporous g-C<sub>3</sub>N<sub>4</sub>. Further, hazardous and costly HF and NH<sub>4</sub>HF<sub>2</sub> limit its industrial-scale production.<sup>86</sup> To overcome this problem, CaCO<sub>3</sub> has been used as a hard template in which the template is removed with dilute HCl after polymerization, leading to the formation of high surface area g-C<sub>3</sub>N<sub>4</sub> with improved photocatalytic activity.<sup>87</sup>

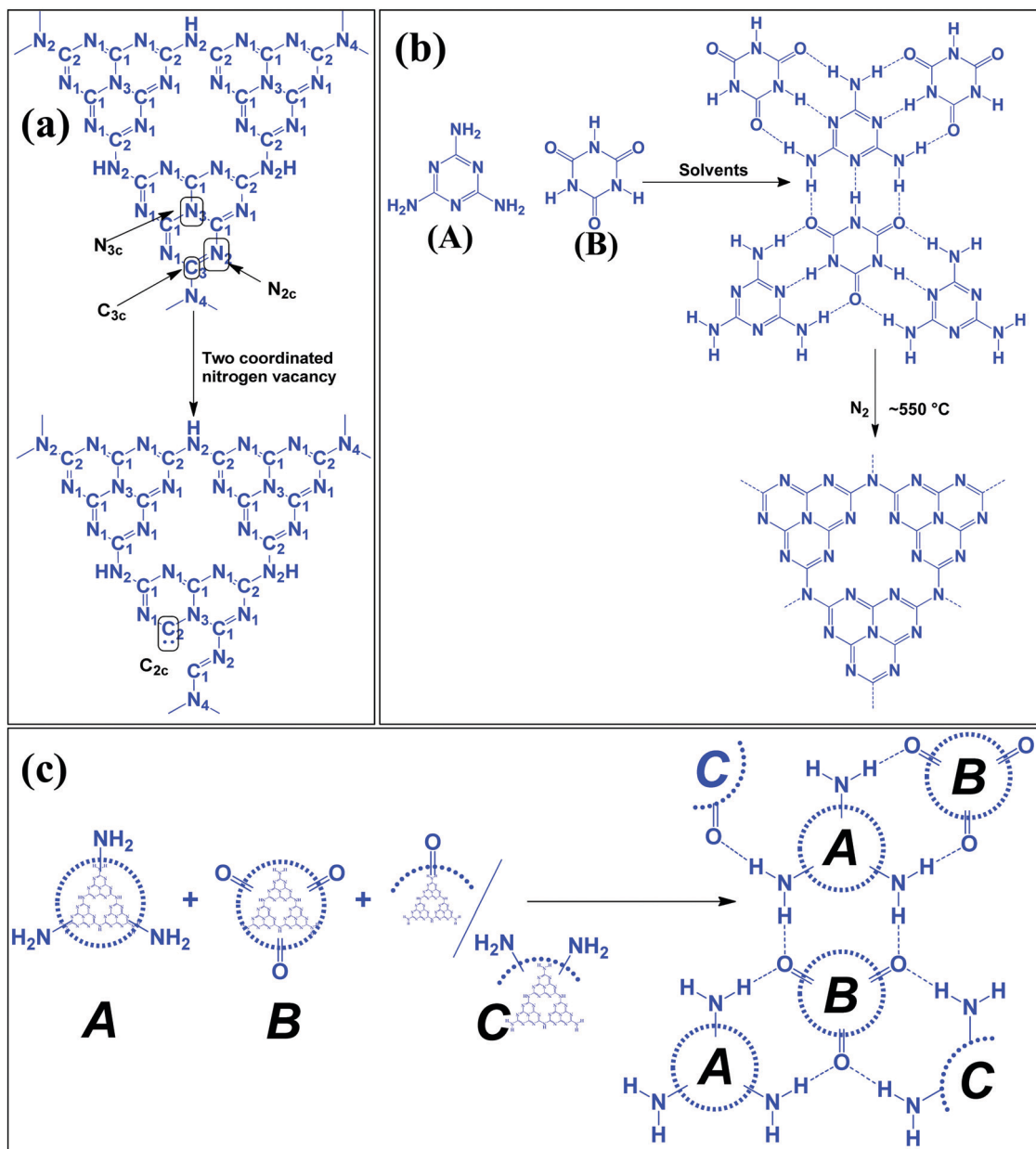
Large scale synthesis of mesoporous g-C<sub>3</sub>N<sub>4</sub> can be achieved using the soft template method. Since the soft template is a carbon-based compound, therefore g-C<sub>3</sub>N<sub>4</sub> with higher C/N is formed with diverse functionality. Ionic liquids and block copolymer-based soft templates have been widely used to prepare mesoporous g-C<sub>3</sub>N<sub>4</sub>. Among the soft templates, only a few templates such as Triton-100 and ionic liquids have produced g-C<sub>3</sub>N<sub>4</sub> with large surface area and porosity.<sup>87–89</sup> Ionic liquids act as soft modifiers for the thermal polymerization, leading to tunable textural properties, morphology, and reactivity. Anions of the ionic liquids participate in the growth process *via* C–N condensation.<sup>90</sup> Dual templating methods and cooperative assembly of two surfactants lead to formation of a

mesophase, which results in the formation of nanostructured materials. A similar strategy can be applied to synthesize mesoporous g-C<sub>3</sub>N<sub>4</sub>. Cooperative supramolecular assembly of surfactants, or a reactant and a surfactant, or reactants induced by hydrogen bonding and other weak electrostatic forces can act as a template to form mesoporous g-C<sub>3</sub>N<sub>4</sub>. Nitrogen-rich organic precursors can easily form hydrogen bonding with “O” containing organic compounds to form self-assembled porous g-C<sub>3</sub>N<sub>4</sub>. For example, melamine can form hydrogen bonding with cyanuric acid in DMSO and form stable aggregates that easily precipitate out (Scheme 8). Such precipitates can be thermally polymerized to form mesoporous g-C<sub>3</sub>N<sub>4</sub>.<sup>91</sup> In fact, supramolecular aggregates can be formed by the hydrogen bonding between three reactants such as melamine, cyanuric acid with urea/barbituric acid/caffeine, *etc.* in a different solvent medium, which results in different morphology of mesoporous g-C<sub>3</sub>N<sub>4</sub> on thermal polymerization with different textural properties and photophysical characteristics (Scheme 8).<sup>92–94</sup> Template mediated synthesis strategies can provide materials with high surface area and porosity but it is difficult to tune the N contents and type of nitrogen due to the specific use of a suitable template and nitrogen precursor. Thus more work is needed to develop soft-template mediated synthesis recipes with variable N contents and functional amine sites using a diverse range of C and N containing precursors for the appointed applications.

### 3.3 Strategies to functionalize and dope elements to improve catalytic activities and light absorption properties

Template-mediated synthetic strategies can improve the surface area and porosity. However, for tuning the catalytic activities relating to the efficient adsorption of reactants and

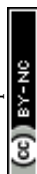




Scheme 8 Supramolecular assembly of precursors leading to formation of mesoporous g-C<sub>3</sub>N<sub>4</sub>.

light absorption, suitable doping and functionalization are required. By varying the nitrogen precursors, the C/N ratio can be tailored, but for doping other elements such as O and S, suitable precursors containing O and S should be used during the synthesis of g-C<sub>3</sub>N<sub>4</sub>. Just by H<sub>2</sub>O<sub>2</sub> treatment, O doped g-C<sub>3</sub>N<sub>4</sub> with high surface area is obtained, which exhibits efficient charge carrier separation, and a red-shift in the absorption spectra.<sup>95</sup> An oxygen group containing melamine precursor also produces O doped g-C<sub>3</sub>N<sub>4</sub>.<sup>96</sup> Organic surfactant mediated synthesis produces C doped g-C<sub>3</sub>N<sub>4</sub>.<sup>97</sup> Diammonium hydrogen phosphate, hexachloro-cyclotriphosphazene, 2-aminoethylphosphonic acid, *etc.* have been used as a P source to dope P into the g-C<sub>3</sub>N<sub>4</sub> framework.<sup>98–100</sup> H<sub>2</sub>S treatment, thiourea, and

tri-thiocyanic acid precursors are known to dope S in the g-C<sub>3</sub>N<sub>4</sub> framework.<sup>101–103</sup> B-Doped g-C<sub>3</sub>N<sub>4</sub> has been prepared by heating melamine and B<sub>2</sub>O<sub>3</sub> above 520 °C to introduce B into the g-C<sub>3</sub>N<sub>4</sub> framework.<sup>104</sup> BH<sub>3</sub>NH<sub>3</sub> and sodium-tetraphenyl boron have been used as precursors to prepare B-doped g-C<sub>3</sub>N<sub>4</sub>.<sup>105,106</sup> Recently, it has been reported that exfoliation of g-C<sub>3</sub>N<sub>4</sub> (prepared by urea and thiourea) with concentrated H<sub>2</sub>SO<sub>4</sub> aqueous solution at 80 °C has produced –SO<sub>3</sub>H functionalized g-C<sub>3</sub>N<sub>4</sub>.<sup>107</sup> Furthermore, it has been found that a small quantity of 2-methyl imidazole can be treated with urea-thiourea to produce O and C co-doped g-C<sub>3</sub>N<sub>4</sub>.<sup>108</sup> Thus, the choice of precursor and the treatment conditions can introduce desired elements and functionality in g-C<sub>3</sub>N<sub>4</sub> to impart the desired catalytic activity.



## 4. Synthesis of chemicals and fuels from CO<sub>2</sub> using g-C<sub>3</sub>N<sub>4</sub> as a catalyst in the conventional thermal catalytic process

Several recent review articles have been published, even using carbon nitrides as catalysts and photocatalysts.<sup>109–111</sup> In this section, the role of CO<sub>2</sub> as a C<sub>1</sub> source to produce chemicals and fuels is described (Scheme 9). CO<sub>2</sub> can also act as an “O” source and can be utilized as an oxidant. CO<sub>2</sub> can be efficiently reduced to provide various chemicals and fuels. All these processes are non-atom efficient processes. CO<sub>2</sub> can be inserted in organic compounds like epoxides, amines, and alkynyl acids to produce organic carbonate and carbamate in a highly atom efficient process.

### 4.1 Role of CO<sub>2</sub> as a soft oxidant or promoter in the oxidation reaction

The oxidizing ability of various gases in coke oxidation follows the order: O<sub>2</sub> (105) > H<sub>2</sub>O (3) > CO<sub>2</sub> (1), and thus CO<sub>2</sub> can act as a soft oxidant.<sup>112</sup> Such a weak oxidation ability is suitable in the oxidative dehydrogenation reaction. Oxidative dehydrogenation in the presence of oxygen leads to the generation of a large amount of CO<sub>2</sub> as a by-product due to the strong oxidizing ability of O<sub>2</sub>.<sup>113</sup> CO<sub>2</sub> is a weaker oxidant and, therefore, it requires activation energy. In general, this activation can be achieved with alkali metals such as Li, K, Na, Ba, Ca, and Cs, which activate CO<sub>2</sub> to form MCO<sub>2</sub>.<sup>114</sup> The activity of Li is higher among the above-mentioned metals due to the CO<sub>2</sub>–M valence bond angle, which is favorable for efficient CO<sub>2</sub> activation and utilization. In this dehydrogenation reaction, CO<sub>2</sub> facilitates the abstraction of hydrogen from hydrocarbons (alkanes, alkenes, alcohols, and alkyl aromatics) *via* catalytic activation to form CO and O species and suppress the total oxidation.<sup>115</sup>

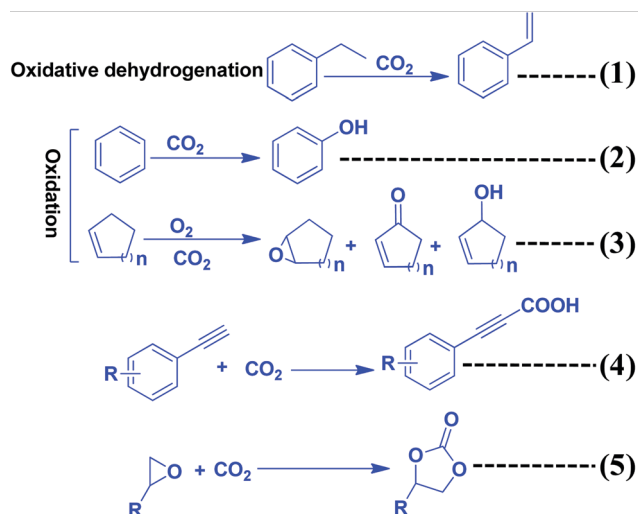
g-C<sub>3</sub>N<sub>4</sub> has been reported for the oxidation of benzene to phenol using CO<sub>2</sub> as a mild oxidant.<sup>116</sup> CO<sub>2</sub> is activated *via* the secondary amine and pyridinic N of g-C<sub>3</sub>N<sub>4</sub>. Benzene is adsorbed weakly on g-C<sub>3</sub>N<sub>4</sub>. CO<sub>2</sub> splits into an  $\ddot{O}$  diradical, followed by its reaction with weakly adsorbing benzene to form the desired phenol as a product (Scheme 10). CO is eliminated after the reaction to regenerate the active sites for further CO<sub>2</sub> adsorption.<sup>116</sup> CO<sub>2</sub> acts as a promoter in the cycloalkene oxidation using O<sub>2</sub> as an oxidant over a g-C<sub>3</sub>N<sub>4</sub> catalyst (eqn (3), Scheme 9).<sup>117</sup> CO<sub>2</sub> is efficiently adsorbed on g-C<sub>3</sub>N<sub>4</sub> to form surface O species and produce CO (confirmed from the FT-IR study). Such surface O assists in the oxidation of cyclic olefins.<sup>116</sup>

### 4.2 Carboxylation of terminal alkynes with CO<sub>2</sub>

Alkynylcarboxylic acid is an important synthetic intermediate for several important compounds. For this reaction to proceed, both the alkyne and CO<sub>2</sub> need to be activated (Scheme 9, eqn (4)). CO<sub>2</sub> can be activated with g-C<sub>3</sub>N<sub>4</sub> but for the efficient adsorption and activation of an alkyne, highly disperse active metals such as Cu or Ag are required. Ag NP decorated g-C<sub>3</sub>N<sub>4</sub> has been reported for this reaction.<sup>118</sup> This reaction follows the Langmuir–Hinshelwood mechanism in which CO<sub>2</sub> is absorbed and activated by g-C<sub>3</sub>N<sub>4</sub> and the alkyne is adsorbed and activated by Ag NPs supported on g-C<sub>3</sub>N<sub>4</sub>. Alkyne acidic hydrogen is activated by the Ag NPs and forms silver acetylide and then this reactive intermediate reacts with activated CO<sub>2</sub> to produce silver carboxylate, which can be easily converted into cesium carboxylate with the help of Cs<sub>2</sub>CO<sub>3</sub> *via* a transmetalation step. Finally, by workup with an acidified solution, it forms the desired alkynyl carboxylic acid.

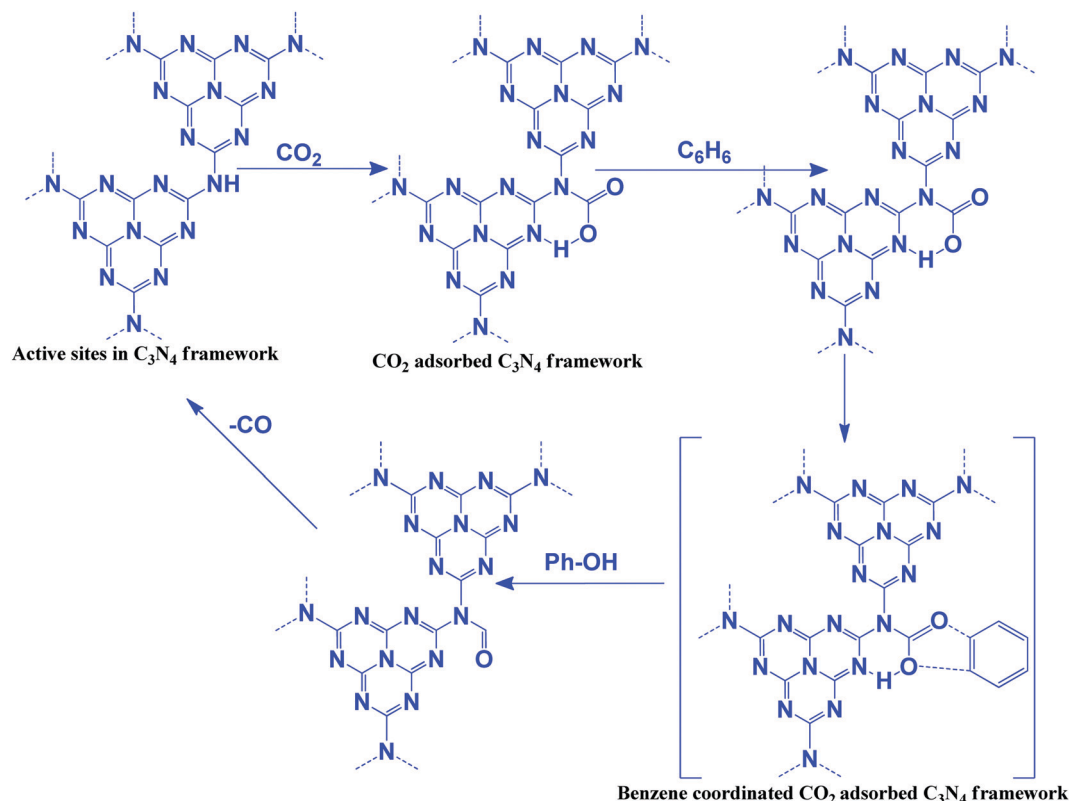
### 4.3 Insertion of CO<sub>2</sub> into epoxides to form cyclic carbonates

The synthesis of cyclic carbonates by the reaction of CO<sub>2</sub> and epoxides is one of the most widely investigated reactions for CO<sub>2</sub> utilization because it is a 100% atom economic process. The cyclic carbonate product can be used as an intermediate for the production of value-added compounds such as Bisphenol-A carbonate (a synthetic polymer used in the packaging industry) and dimethyl carbonate (a methylating agent and a green solvent), the latter being a good fuel additive and a precursor for various other industrially important chemical transformations. CO<sub>2</sub> can be inserted in the epoxide ring *via* a non-concerted process to produce the cyclic carbonate. For this reaction, generally, a catalyst should contain a Lewis basic site, which acts as a nucleophile to activate the ‘C=O’ bond of the CO<sub>2</sub> molecule, while a Lewis acidic site acts as an electrophile for activating the epoxide ring.<sup>119</sup> The Lewis acidic site polarizes the epoxide ring and forms an epoxide anion and then it acts as a nucleophile and attacks the CO<sub>2</sub> molecule, leading to the formation of a carbonate anion intermediate. The next step is either a ring-closing process for the formation of the cyclic carbonate or insertion of one or more epoxides and CO<sub>2</sub> molecules into the intermediate, which produces a polycarbonate.<sup>119</sup> g-C<sub>3</sub>N<sub>4</sub> has surface/edge located free –NH<sub>2</sub> groups which serve as Lewis basic sites and act as the



**Scheme 9** Synthesis of organic compounds using CO<sub>2</sub> as a reactant or oxidant or promoter and C<sub>3</sub>N<sub>4</sub> as a catalyst.





Scheme 10 Oxidation of benzene to phenol using CO<sub>2</sub> as an oxidant and g-C<sub>3</sub>N<sub>4</sub> as a catalyst.

nucleophile to activate CO<sub>2</sub>. However, a Lewis acid site is absent to activate the epoxide. The weak activity of g-C<sub>3</sub>N<sub>4</sub> towards this reaction is due to surface -OH sites which have the ability to weakly activate the epoxide and produce a very low yield of the product. For this reaction to proceed efficiently, a co-catalyst such as tetra butyl ammonium bromide (TBAB) is used. However, suitable metal-free bi-functional g-C<sub>3</sub>N<sub>4</sub> is reported to effectively catalyze this reaction.<sup>107</sup> Production of cyclic carbonates using different g-C<sub>3</sub>N<sub>4</sub> based catalysts is summarized in Table 2. Conventional g-C<sub>3</sub>N<sub>4</sub> exhibits very low activity (Table 2, entry 1).<sup>120</sup> Mesoporous g-C<sub>3</sub>N<sub>4</sub> (prepared with different precursors) has exhibited better activity than conventional g-C<sub>3</sub>N<sub>4</sub>. Among the prepared materials, mesoporous g-C<sub>3</sub>N<sub>4</sub> prepared with melamine exhibits 34% propylene oxide conversion, and 90% propylene carbonate selectivity with a TOF of 3.4 h<sup>-1</sup> at moderate reaction conditions in DMF (Table 2, entries 2–4).<sup>121</sup> A mixture of gas (50% O<sub>2</sub> and 50% CO<sub>2</sub>) is used during this reaction and it has been proposed that O<sub>2</sub> plays a promotional role for the activation of the epoxide by supplying additional oxygen during the course of the reaction.<sup>120</sup> Further, to improve the surface area and dispersed active sites, SBA-15 supported g-C<sub>3</sub>N<sub>4</sub> is employed for propylene carbonate synthesis under solvent-free conditions in the presence of doped metal ions, which act as Lewis acid sites. Among metal salts, Zn<sup>2+</sup> exhibits the highest conversion and selectivity of 98.3% and >99%, respectively (Table 2, entries 5–8).<sup>122</sup> The activity of doped Zn<sup>2+</sup> is better than the supported ZnCl<sub>2</sub> catalyst (Table 2, entry 9).<sup>123</sup> It is proposed that the C<sub>3</sub>N<sub>4</sub> nanosphere inside the mesopore

channel of silica supplies the activated CO<sub>2</sub> to the epoxide through a transfer mechanism and facilitates cyclic carbonate synthesis *via* Zn<sup>2+</sup> activated epoxide species. Porous and amino rich g-C<sub>3</sub>N<sub>4</sub> is prepared without using any hard template using urea as a precursor to form defective and edge enriched C<sub>3</sub>N<sub>4</sub>. The urea derived C<sub>3</sub>N<sub>4</sub> produces a wide variety of cyclic carbonates in the absence of any co-catalyst under solvent-free conditions (Table 2, entries 10–13).<sup>124</sup> It is suggested that the edge defects contain both primary amines (-NH<sub>2</sub>) and secondary amine groups (-C-NH-C) and therefore they are able to activate CO<sub>2</sub>. The epoxide is activated by the defective C<sub>3</sub>N<sub>4</sub> edge containing -OH groups. Further, it has been revealed that urea derived C<sub>3</sub>N<sub>4</sub> exhibits relatively higher surface area (41 m<sup>2</sup> g<sup>-1</sup>) than dicyandiamide derived C<sub>3</sub>N<sub>4</sub> and a porous nature, and the existence of macropores facilitates mass transfer and enhances the catalytic activity. *n*-Bromobutane grafted mesoporous g-C<sub>3</sub>N<sub>4</sub> exhibits better catalytic activity for solvent-free propylene carbonate synthesis with 88% conversion and 98% selectivity (Table 2, entry 14).<sup>125</sup> In this process, the *in situ* depleted Br<sup>-</sup> acts as the electrophile which activates the epoxide ring, while the defect rich edge positioned -NH<sub>2</sub> polarizes the CO<sub>2</sub> molecule leading to the bi-functional nature. Butane in conjugation with halides have been evaluated and it has been found that bromide exhibits the best activity because of its better leaving group ability. The better catalytic activity is attributed to the mesoporous nature (15 nm), high surface area (241 m<sup>2</sup> g<sup>-1</sup>), and coexistence of terminal -NH<sub>2</sub> groups in g-C<sub>3</sub>N<sub>4</sub> and X<sup>-</sup> groups, which is even far better than the

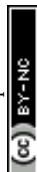
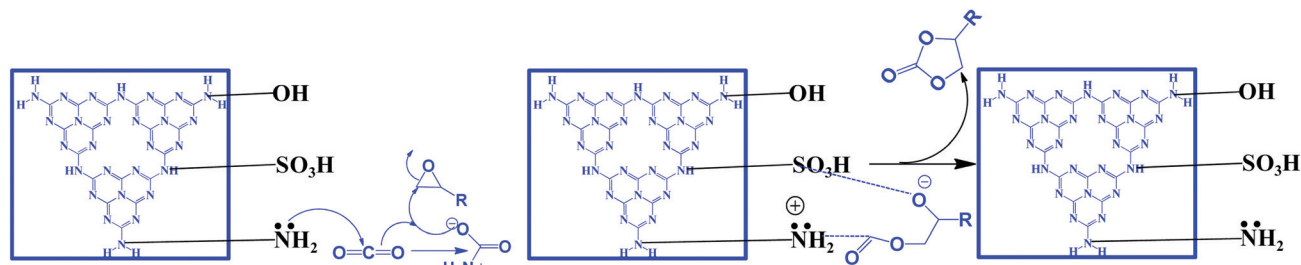


Table 2 Comparative catalytic activity of various g-C<sub>3</sub>N<sub>4</sub> based catalysts in the synthesis of cyclic carbonates

| E. no. | Catalyst                                                                                                          | $S_{\text{BET}}$<br>(m <sup>2</sup> g <sup>-1</sup> ) | Co-catalyst                | Reaction conditions                                                                                                                              | Conv. (%) | Select. (%) | TOF (h <sup>-1</sup> ) | Ref. |
|--------|-------------------------------------------------------------------------------------------------------------------|-------------------------------------------------------|----------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------|-----------|-------------|------------------------|------|
| 1.     | g-C <sub>3</sub> N <sub>4</sub> (prepared with melamine)                                                          | 64                                                    | —                          | ECH (5 mL), catalyst (50 mg), temp. (150 °C), press. (3.5 MPa), time (2 h)                                                                       | 5.6       | 96.7        | 3.3                    | 120  |
| 2.     | Mesoporous-C <sub>3</sub> N <sub>4</sub> (prepared with melamine-2D mesoporous silica)                            | —                                                     | —                          | PO (1.5 mL), catalyst (20 mg), DMF (10), temp. (100 °C), press. (0.55 MPa), time (10 h)                                                          | 34.0      | 90          | —                      | 121  |
| 3.     | Mesoporous-C <sub>3</sub> N <sub>4</sub> (prepared with urea-formaldehyde-2D mesoporous silica)                   | —                                                     | —                          | PO (1.5 mL), catalyst (20 mg), DMF (10), temp. (100 °C), press. (0.55 MPa), time (10 h)                                                          | 28.0      | 92          | —                      | 121  |
| 4.     | MS-C <sub>3</sub> N <sub>4</sub> (prepared with melamine-glyoxal-2D mesoporous silica)                            | —                                                     | —                          | PO (1.5 mL), catalyst (20 mg), DMF (10), temp. (100 °C), press. (0.55 MPa), time (10 h)                                                          | 25.0      | 98          | —                      | 121  |
| 5.     | Mesoporous-C <sub>3</sub> N <sub>4</sub> (prepared with dicyandiamide-SBA-15)                                     | 586                                                   | —                          | SO (3 mL), catalyst (100 mg), temp. (150 °C), press. (3.5 MPa), time (1.5 h)                                                                     | 28.8      | 99          | 4.8                    | 122  |
| 6.     | Mesoporous-C <sub>3</sub> N <sub>4</sub> (prepared with dicyandiamide-SBA-15)                                     | 585                                                   | Zn <sup>2+</sup> (0.8 wt%) | SO (3 mL), catalyst (100 mg), temp. (150 °C), press. (3.5 MPa), time (1.5 h)                                                                     | 94.5      | 99          | 15.8                   | 122  |
| 7.     | Mesoporous-C <sub>3</sub> N <sub>4</sub> (prepared with dicyandiamide-SBA-15)                                     | 585                                                   | Zn <sup>2+</sup> (0.8 wt%) | PO (3 mL), catalyst (100 mg), temp. (150 °C), press. (3.5 MPa), time (1.5 h)                                                                     | 97.1      | 99          | 16.2                   | 122  |
| 8.     | Mesoporous-C <sub>3</sub> N <sub>4</sub> (prepared with dicyandiamide-SBA-15)                                     | 585                                                   | Zn <sup>2+</sup> (0.8 wt%) | EO (3 mL), catalyst (100 mg), temp. (150 °C), press. (3.5 MPa), time (1.5 h)                                                                     | 98.3      | 99          | 16.4                   | 122  |
| 9.     | 10%ZnCl <sub>2</sub> /mesoporous C <sub>3</sub> N <sub>4</sub>                                                    | 206                                                   | Zn <sup>2+</sup>           | EO (7 mL), DMF (3 mL), catalyst (200 mg), temp. (140 °C), press. (2.5 MPa), time (6 h)                                                           | 73.0      | 99.4        | 5.6                    | 123  |
| 10.    | g-C <sub>3</sub> N <sub>4</sub> (prepared with urea and heated at 480 °C)                                         | 40.8                                                  | —                          | ECH (21.5 mmol), catalyst (100 mg), temp. (130 °C), press. (3.5 MPa), time (4 h)                                                                 | 99.6      | 96.4        | —                      | 124  |
| 11.    | g-C <sub>3</sub> N <sub>4</sub> (prepared with urea and heated at 480 °C)                                         | 40.8                                                  | —                          | EO (21.5 mmol), catalyst (100 mg), temp. (130 °C), press. (3.5 MPa), time (4 h)                                                                  | 91.7      | 99.3        | —                      | 124  |
| 12.    | g-C <sub>3</sub> N <sub>4</sub> (prepared with urea and heated at 480 °C)                                         | 40.8                                                  | —                          | SO (21.5 mmol), catalyst (100 mg), temp. (130 °C), press. (3.5 MPa), time (4 h)                                                                  | 98.0      | 99.8        | —                      | 124  |
| 13.    | g-C <sub>3</sub> N <sub>4</sub> (prepared with urea and heated at 480 °C)                                         | 40.8                                                  | —                          | PO (21.5 mmol), catalyst (100 mg), temp. (130 °C), press. (3.5 MPa), time (24 h)                                                                 | 99.3      | 99.6        | —                      | 124  |
| 14.    | <i>n</i> -ButBr doped mesoporous C <sub>3</sub> N <sub>4</sub> (prepared with cyanamide and 12 nm silica spheres) | 248                                                   | Br <sup>-</sup>            | PO (10 mL), catalyst (200 mg), temp. (140 °C), press. (2.5 MPa), time (6 h)                                                                      | 88.0      | 99.7        | 9.6                    | 125  |
| 15.    | H <sub>2</sub> SO <sub>4</sub> treated g-C <sub>3</sub> N <sub>4</sub> (prepared with melamine)                   | 32                                                    | —                          | ECH (5 mL), catalyst (50 mg), temp. (130 °C), press. (3.5 MPa), time (2 h)                                                                       | 99.1      | 99.5        | 58.1                   | 120  |
| 16.    | H <sub>2</sub> SO <sub>4</sub> treated g-C <sub>3</sub> N <sub>4</sub> (prepared with urea-thiourea)              | 10                                                    | —                          | ECH (63 mmol), catalyst (50 mg), Temp. (100 °C), press. (1.0 MPa), time (4 h)                                                                    | 93.5      | 99          | —                      | 107  |
| 17.    | P-Doped g-C <sub>3</sub> N <sub>4</sub> (prepared with melamine-hexachloro-triphenylphosphazene)                  | —                                                     | Bu <sub>4</sub> NBr        | PO (28.6 mmol), catalyst (150 mg), Bu <sub>4</sub> NBr (2.4 mol%), temp. (100 °C), press. (2.0 MPa), time (4 h)                                  | 99.8      | 99.9        | —                      | 126  |
| 18.    | B-Doped g-C <sub>3</sub> N <sub>4</sub> (prepared with melamine-BmimBF <sub>4</sub> )                             | 11.1                                                  | —                          | ECH (14.3 mmol), catalyst (50 mg), temp. (130 °C), press. (2.8 MPa), time (6 h)                                                                  | 89.0      | 98.0        | 5.8                    | 127  |
| 19.    | B-Doped g-C <sub>3</sub> N <sub>4</sub> (prepared with melamine-BmimBF <sub>4</sub> )                             | 11.1                                                  | —                          | PO (14.3 mmol), catalyst (50 mg), temp. (130 °C), press. (2.8 MPa), time (6 h)                                                                   | 31.0      | 99.9        | 1.5                    | 127  |
| 20.    | B-Doped mesoporous g-C <sub>3</sub> N <sub>4</sub> (prepared with dicyandiamide-B(OH) <sub>3</sub> -SBA-15)       | 123                                                   | —                          | SO (1 mL), catalyst (30 mg), temp. (130 °C), press. (3.0 MPa), time (24 h)                                                                       | 97.8      | 96.8        | —                      | 129  |
| 21.    | g-C <sub>3</sub> N <sub>4</sub> (prepared with melamine)                                                          | —                                                     | Bu <sub>4</sub> NBr        | Light assisted method, ECH/SO (13.7 mmol), catalyst (50 mg), Bu <sub>4</sub> NBr (1.8 mol%) temp. (105 °C), press. (1 atm, balloon), time (20 h) | 100       | 100         | —                      | 130  |

ECH = epichlorohydrin, PO = propylene oxide, SO = styrene oxide, TOF = turnover frequency, conv. = epoxide conversion, selec. = cyclic carbonate selectivity,  $S_{\text{BET}}$  = surface area.





**Scheme 11** Synthesis of cyclic carbonates using bi-functional g-C<sub>3</sub>N<sub>4</sub>, reproduced with permission, Copyright Royal Society of Chemistry, ref. 107.

externally induced *n*-bromobutane to carbon nitride during the course of the reaction (Table 1, entry 14).<sup>125</sup> Huang *et al.* have reported H<sub>2</sub>SO<sub>4</sub> treated, protonated acidic functionality containing carbon nitride, which has been investigated for the solvent-free cycloaddition of CO<sub>2</sub> with epichlorohydrin for the synthesis of the corresponding carbonate (Table 2, entry 15).<sup>126</sup> This protonation process has altered the electronic and textural properties and surface chemistry, which results in the incorporation of hydroxyl groups and amine groups into the bulk urea derived carbon nitride, which provides 17 fold higher catalytic activity. Surface hydroxyl groups (–OH) and amine groups (–NH<sub>2</sub>) are responsible for this improved reactivity. Inspired by this work, Samanta *et al.* have synthesized urea and thiourea derived carbon nitride CN-(UTU), which exhibits moderate catalytic activity better than the counterparts urea/thiourea under solvent-free cyclic carbonate synthesis from epichlorohydrin without any co-catalyst (Table 2, entry 16) (Scheme 11).<sup>107</sup> Further, to improve the catalytic activity, the resultant materials have been chemically exfoliated with H<sub>2</sub>SO<sub>4</sub>, which results in the simultaneous incorporation of –SO<sub>3</sub>H and –NH<sub>2</sub> groups into the melem framework (Scheme 11).

The bi-functional catalyst (which is ideal for the CO<sub>2</sub> activation reaction) has been investigated in the CO<sub>2</sub> insertion reaction by varying the epoxide under very mild conditions (100 °C and 1 MPa). Excellent conversion and selectivity are achieved over this bi-functional CN-(UTU) catalyst. The simultaneous introduction of –NH<sub>2</sub> groups (that act as basic sites for activating CO<sub>2</sub>) and –SO<sub>3</sub>H groups (that polarize the epoxide ring) has been confirmed from CO<sub>2</sub>/NH<sub>3</sub>-TPD. The synergistic participation of –NH<sub>2</sub> and –SO<sub>3</sub>H groups during the reaction is the main reason for obtaining the excellent product yield. Moreover, doping with non-metals (such as P and B) over g-C<sub>3</sub>N<sub>4</sub> is also reported for the synthesis of cyclic carbonates, which is found to be very effective for this reaction.<sup>126–129</sup> The –B(OH) and –P(OH) groups are found to be responsible for the catalytic activity enhancement (Table 2, entries 19–20). All the reactions discussed above require CO<sub>2</sub> pressure higher than one atmosphere. Only one report is available in which a cyclic carbonate is synthesized at 1 atm pressure under solvent-free conditions but in the presence of external co-catalyst (1.8%) TBAB (Table 2, entry 22).<sup>130</sup> Therefore, more work should be carried out to synthesize efficient catalysts that can catalyze this reaction under solvent-free conditions at atmospheric pressure.

## 5. Photocatalytic CO<sub>2</sub> reduction reaction (CO<sub>2</sub> photoreduction)

Photocatalytic CO<sub>2</sub> reduction is a surface catalytic reaction which employs light-induced holes and electrons generated over the surface of the catalyst and protons from the reaction medium. In ideal conditions, CO<sub>2</sub> reduction follows some fundamental steps, like other light assisted reactions such as water splitting, N<sub>2</sub> reduction, and organic transformations.<sup>131–140</sup> The basic operating steps involved in the photocatalytic CO<sub>2</sub> reduction reaction are summarized in two categories, namely optoelectronic and physicochemical steps, that are complementary to each other:

(A) Optoelectronic steps (Scheme 12A):

- (1) Creation of charge carriers (e<sup>–</sup>–h<sup>+</sup>) on the semiconductor surface by illumination with light having energy greater than the bandgap:  $E_{hv} \geq E_g$ .
- (2) Light-induced charge carrier dissociation and migration on the catalytic surface.
- (3) Charge-carrier recombination followed by the dissipation of energy in the form of heat.
- (4) The transported electrons participate in CO<sub>2</sub> reduction whereas holes participate in the oxidation of H<sub>2</sub>O to O<sub>2</sub>, and H<sup>+</sup> or any other reducing agents to their corresponding oxidized products.

(B) Physicochemical steps (Scheme 12B):

- (1) Adsorption of CO<sub>2</sub> on the catalyst surface.
- (2) CO<sub>2</sub> activation by the inherent functional groups of the catalyst.
- (3) Intermediate species adsorption and desorption of the products from the catalytically active sites during the reduction reaction.

### 5.1 Factors and parameters governing the mechanism of CO<sub>2</sub> photoreduction

Carbon dioxide is the most stable and inert oxidized product, while methane is a completely reduced form of carbon (Scheme 13a). The light-induced conversion of CO<sub>2</sub> to hydrocarbons and oxygenated hydrocarbons depends on various factors and parameters. These factors determine the fate of the reaction as well as the CO<sub>2</sub> conversion and product distribution. Various factors influencing CO<sub>2</sub> reduction, especially with graphitic carbon nitride as a photocatalyst, are discussed in this section.

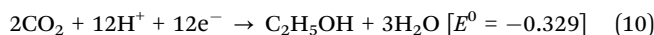
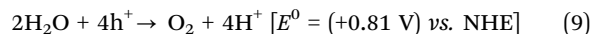
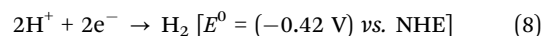
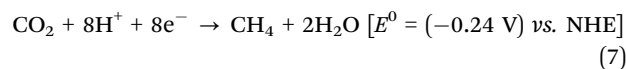
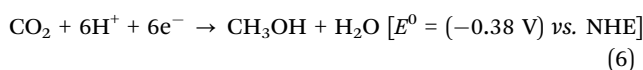
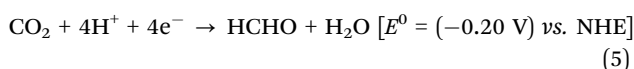
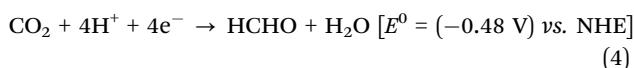
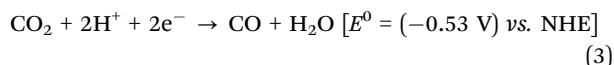
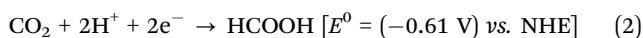
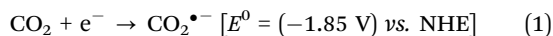
**5.1.1 Thermodynamic factors.** CO<sub>2</sub> conversion is a thermodynamically uphill energy process that requires a tremendous





**Scheme 12** Basic mechanism of the photocatalytic process: (A) optoelectronic steps and (B) physicochemical steps involved in the CO<sub>2</sub> reduction reaction.

energy input because of its stable and inert nature. An energy of 750 kJ mol<sup>-1</sup> is required for the splitting of one 'C=O' bond in the CO<sub>2</sub> molecule. CO<sub>2</sub> has a low electron affinity and a high energy gap of 13.7 eV between its lowest unoccupied molecular orbital (LUMO) and highest occupied molecular orbital (HOMO).<sup>137</sup> CO<sub>2</sub> is a linear molecule and electronic accumulation in the empty LUMO (antibonding  $\Pi^*$  orbital) facilitates the transformation of a linear to a bent geometry over the catalyst surface, which in turn expedites the kinetic barrier and enables its smooth conversion to appropriate products. The band edge position of any photocatalyst decides this electron accumulation and, accordingly, the product distributions occur. Particularly, the valence band edge must be positioned at a more negative electrode potential than the standard reduction potential of CO<sub>2</sub> to any other products otherwise the process is not thermodynamically feasible (Scheme 13b). Based on the standard reduction potential,<sup>138</sup> the reduced products are given below, which must be considered to analyze the reaction products after the reaction to establish the plausible mechanism of the reaction (Scheme 13b).



Among these reactions, eqn (1) is the most unfavorable one and, thus, the proton-coupled electron transfer process always occurs on the catalyst surface with the production of different products.

**5.1.2 Reaction mechanism and selectivity control.** CO<sub>2</sub> reduction occurs sequentially involving multiple electrons and protons. Also, it requires the splitting of highly stable 'C=O' bonds and the creation of C-H bonds. The complete reduction occurs *via* the formation of several radical intermediates, which also tend to recombine at different stages during the course of a reaction.<sup>139</sup> The product distribution depends upon the specific set of reaction conditions. The simultaneous and competitive photo-oxidation and reduction reactions bring complexity to CO<sub>2</sub> photoreduction. The most important parameters influencing these reactions include the type of photocatalyst, band edge potentials and bandgap, reaction conditions and medium, co-catalyst, *etc.*, which are briefly described here. The first study reported for CO<sub>2</sub> photoreduction involves TiO<sub>2</sub> powder under photo-electrochemical conditions.<sup>140</sup> Based on the mechanistic investigation reported in the literature, three possible pathways have been proposed, (I) the formaldehyde pathway forming CO<sub>2</sub> → CH<sub>4</sub> (fast hydrogenation pathway), (II) the carbene pathway forming CO<sub>2</sub> → CH<sub>4</sub> (fast deoxygenation pathway), and (III) the glyoxal pathway (Scheme 14I–III).<sup>131,141</sup> It may be noted that at some stage of the reaction, all the suggested





**Scheme 13** (a) The different states of carbon with the potential difference, and (b) photocatalytic  $\text{CO}_2$  reduction with respect to the standard redox potential of different products and band straddling.

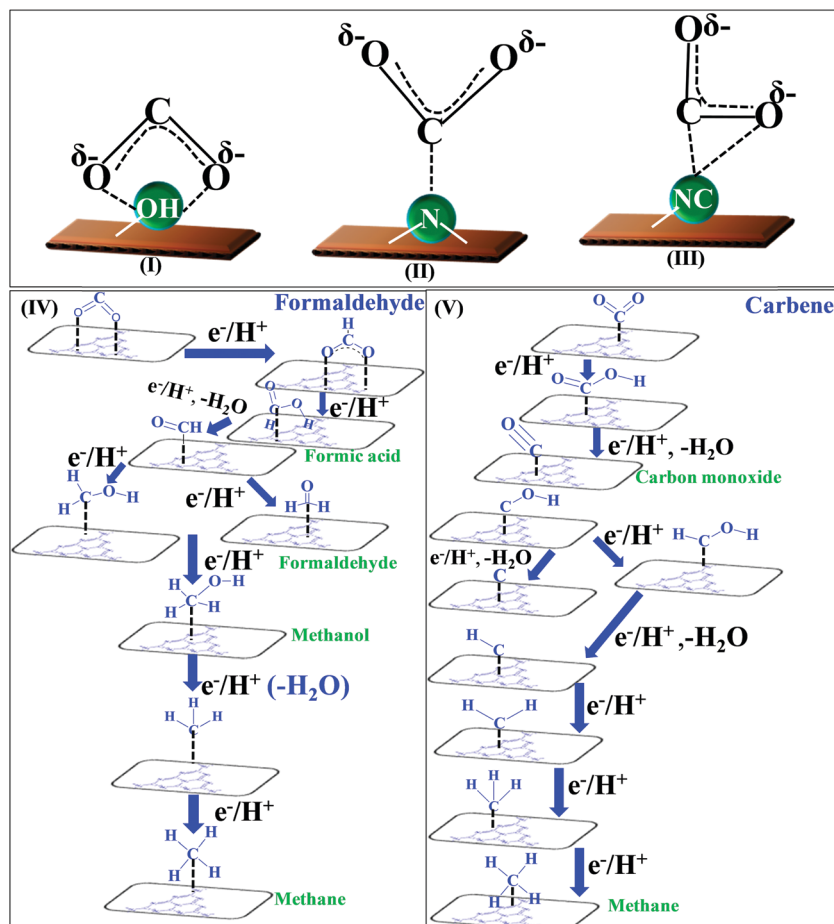
pathways can equally dominate. The elementary steps based on the specific reaction conditions need more work to establish the possible involvement of several radical species. The above-mentioned pathways are specifically discussed here.

The conversion of  $\text{CO}_2$  through the formaldehyde pathway involves multiple electrons and protons, which leads to the formation of several products such as  $\text{HCHO}$ ,  $\text{HCOOH}$  and  $\text{CH}_3\text{OH}$ , and finally the deoxygenated product  $\text{CH}_4$ . Starting from  $\text{CO}_2$  involving  $2\text{e}^-/2\text{H}^+$ , it first produces  $\text{HCOOH}$  (oxyhydrogenation), followed by involving  $2\text{e}^-/2\text{H}^+$  again in the next step, which produces  $\text{HCHO}$  (reductive dehydrogenation).<sup>142</sup> These two species formed during the reaction are intermediates and not side products. They can proceed to form unstable and reactive intermediates like  $(^*\text{CHO})/(^*\text{COOH})$ , which in the next step takes up  $2\text{e}^-/2\text{H}^+$  to produce  $\text{CH}_3\text{OH}$ .  $\text{CH}_3\text{OH}$  can split to form  $(^*\text{CH}_3)$  and  $(^*\text{OH})$  radicals.  $(^*\text{CH}_3)$  can take one  $(^*\text{H})$  produced from the cleavage of  $\text{HCHO}$  to form  $\text{CH}_4$  at the end. The formation of  $\text{CH}_3\text{OH}/\text{HCHO}$  also depends on the adsorption and desorption energy over the catalyst surface. If the

coordination is strong enough and the desorption process is not facile, then  $\text{CH}_3\text{OH}/\text{HCHO}$  will be the main product (Scheme 14(IV)). However, a weak binding energy allows this intermediate to produce the completely reduced  $\text{CH}_4$ .<sup>141</sup> Depending upon the intermediate binding energy, the yields and selectivity are regulated.

If the conversion process of  $\text{CO}_2 \rightarrow \text{CH}_4/\text{CH}_3\text{OH}$  follows the carbene pathway, then several reactive intermediates can form *via* de-oxygenation followed by hydrogenation elementary steps. It can propagate like  $\text{CO}_2 \rightarrow \text{CO} \rightarrow (^*\text{C}) \rightarrow (^*\text{CH}_3) \rightarrow \text{CH}_3\text{OH}/\text{CH}_4$ . Thus this suggests the following: (1)  $\text{CO}$  formation and radical formation of  $(^*\text{C})$  and  $(^*\text{CH}_3)$ . (2) In the second step, the competitive reduction tendency between  $(^*\text{CH}_3/^*\text{C})$  and  $(^*\text{OH}/^*\text{H})$  often determines the product selectivity. In the first step,  $\text{CO}_2$  is adsorbed on the catalyst surface through carbon co-ordination by the  $1\text{e}^-/1\text{H}^+$  attachment to form the carboxyl radical  $(^*\text{COOH})$ . After coupling with several  $\text{e}^-/\text{H}^+$ , this can undergo decarboxylation to produce the formyl radical  $(^*\text{CHO})$ . In the next step, it can break into  $(^*\text{C})$  and also can take three protons





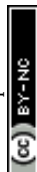
**Scheme 14** Co-ordination geometry of CO<sub>2</sub> over the catalyst surface (I–III), and plausible reaction mechanism of CO<sub>2</sub> reduction: formaldehyde pathway (IV), carbene pathway (V).

to produce the (<sup>•</sup>CH<sub>3</sub>) radical intermediate. In the final state, it couples with one H<sup>+</sup> or (<sup>•</sup>OH) to form CH<sub>3</sub>OH or CH<sub>4</sub> as the end product (Scheme 14(V)).<sup>143</sup> The adsorption strength of CO on the catalyst surface also influences the product selectivity. If the CO absorption is very strong then it could lead to the formation of several interim products and, consequently, desorption as different products. But if the absorption is weaker, then the easy desorption produces CO as an exclusive reduced product. In this process, the formation of radical species (<sup>•</sup>CH<sub>3</sub>) and (<sup>•</sup>C) has been postulated and therefore electron paramagnetic resonance (EPR) measurements are very crucial to obtain a clear understanding of the reaction mechanism.

In the glyoxal mechanism, the first step is the formation of the one-electron reduction of (CO<sub>2</sub><sup>•-</sup>), which takes 1e<sup>-</sup>/1H<sup>+</sup> to form the formyl radical (<sup>•</sup>CHO). This further undergoes dimerization to form the glyoxal product. The complete sequence of the elementary steps involves oxidation and reduction reactions as follows: (CO<sub>2</sub> → CO<sub>2</sub><sup>•-</sup>) → (<sup>•</sup>CHO) → CHO-CHO → CH<sub>2</sub>OH-CHO → (<sup>•</sup>CH<sub>2</sub>-CHO) → CHO-CH<sub>3</sub> → (CH<sub>3</sub>OC<sup>•</sup>) → (<sup>•</sup>CH<sub>3</sub>) → CH<sub>4</sub>/CH<sub>3</sub>OH. However, this process is less explored and not completely understood yet.<sup>144</sup> Various C<sub>2</sub> products such as C<sub>2</sub>H<sub>5</sub>OH or CH<sub>3</sub>COOH *etc.* are believed to proceed through this pathway.

Based on the experimental observation and radical trapping/detection experiments, mechanistic routes are postulated, which requires further exploration. However, recent theoretical calculations suggest that defects and preferable coordination such as oxygen or carbon binding of the reactant, CO<sub>2</sub>, over the catalyst frequently determine the selectivity of the obtained product. A defect-less catalyst surface produces CH<sub>4</sub> only, while oxygen defect containing g-C<sub>3</sub>N<sub>4</sub> catalysts simultaneously form CH<sub>3</sub>OH/CH<sub>4</sub>.<sup>145,146</sup> This is an interesting observation and more detailed work is required.

**5.1.3 Light-harvesting capacity and band straddling.** Apparently, 40% is visible radiation within the spectral range of 0.4 to 0.7 μm, 51% is infrared radiation in the spectral region of 0.7 to 4 μm and 4% is UV radiation of the solar radiation received on the Earth's surface on a sunny day.<sup>147</sup> The total radiation energy emitted by the Sun in unit time remains practically constant. A blue shift of the light-harvesting domain also changes the thermal energy associated with the phonons. However, in a lab-scale reaction system, the most widely used 1.5 AM G light source provides energy equivalent to the energy of 1 Sun, which is 100 mW cm<sup>-2</sup>.<sup>148</sup> The focus should be on developing materials that absorb in a wide area of the light spectrum and have a bandgap of 1.23 eV or higher so that the kinetic



**5.1.5 CO<sub>2</sub>-Philicity.** CO<sub>2</sub> adsorption on the photocatalyst surface is the very first and crucial step. Higher CO<sub>2</sub> philicity

**5.1.6 Effect of the reaction medium.** CO<sub>2</sub> photoreduction can be carried out in two different reaction conditions, liquid phase suspension and solid-gas interaction.<sup>153</sup> In the suspension medium, the photocatalyst is dispersed in water medium. In the second one the photocatalyst is immobilized on a solid substrate and then water vapor is introduced through the reactor. In the first case, sometimes a CO<sub>2</sub> miscibility agent like acetonitrile or DMF is used, which is not an ideal reaction condition. The catalyst at the periphery of the reactor would get less light illumination and thus uniform light absorption may not be achieved. Moreover, the symmetrical photocatalyst particle distribution also depends upon mechanical stirring and those particles in the close vicinity of water molecules would contribute more to the resultant catalytic activity. The mixing of acetonitrile or DMF into water raises a question about the actual source of carbon during the CO<sub>2</sub> photoreduction reaction. So in ideal reaction conditions, a solid photocatalyst dispersed in CO<sub>2</sub> gas purge-pure water must be considered when performing this reaction to avoid any unwanted products. Besides, in the gas-solid medium, the gaseous CO<sub>2</sub> is allowed to react with the immobilized solid catalyst and H<sub>2</sub>O vapor, which itself is in the gaseous phase. Considering the two states of H<sub>2</sub>O and its ease of oxidation either in the liquid state or in the vapor phase, the activity and selectivity are greatly affected. Moreover, in this case, uniform light illumination is reached over the photocatalyst surface. Therefore, the total photons available in the solid-gas interaction are always more than in the case of the liquid-suspension medium. Additionally, the vapor phase of

H<sub>2</sub>O is associated with extra thermal kinetic energy, which is eventually beneficial for overcoming the thermodynamic barriers. While in the first case oxygenated hydrocarbons like CH<sub>3</sub>OH, HCOOH, *etc.* are preferably formed, in the second case complete hydrocarbons (CH<sub>4</sub>) or oxy-carbons (CO) are achieved as products. Thus one should select different reaction media to obtain the desired product.

**5.1.7 Impact of the CO<sub>2</sub> source.** CO<sub>2</sub> can be directly inserted in the reaction medium from a gas cylinder or it can be *in situ* generated. The gaseous CO<sub>2</sub> is purged into water (pH = 7), which is a typical reaction condition of liquid-suspension medium as discussed in the previous section. It remains in a comparably less activated state than the *in situ* generated CO<sub>2</sub> from different sources.<sup>139</sup> Moreover, CO<sub>2</sub> can be *in situ* generated by the reaction between NaHCO<sub>3</sub> and H<sub>2</sub>SO<sub>4</sub> or can be activated by employing NaOH. Sometimes elevated temperature and pressure are employed to make the interaction between CO<sub>2</sub> and the photocatalyst more kinetically favorable. This, in turn, favors HCO<sub>3</sub><sup>−</sup> formation and also lowers the pH of the medium (pH < 7). Therefore, the CO<sub>2</sub> reduction follows the proton-coupled-electron-transfer (PCET) mechanism, leading to the formation of a wide range of products.<sup>154</sup> Ideally, employing additional thermal energy or *in situ* CO<sub>2</sub> generation inside the reactor is not a true photocatalytic reaction, but it is preferably used in the electrocatalytic process since NaHCO<sub>3</sub> itself acts as a CO<sub>2</sub> source as well as an electrolyte.

**5.1.8 Influence of the reactor design.** The reactor design and fabrication also significantly influence the CO<sub>2</sub> photo-conversion efficiency. The major aim of designing a suitable photoreactor is to enhance the light-harvesting capacity uniformly, prohibiting the loss of photon energy, facilitating the ease of product separation, and allowing an appropriate interaction between the reactants and light energy with even distribution.<sup>155</sup> Basically, on the lab-scale, photoreactor systems can be classified into two categories: (1) a quartz glass reactor that allows light irradiation from the horizontal direction on the reaction system and (2) a moderately pressurized reactor containing a quartz glass window which can sustain a pressure up to 10 bar CO<sub>2</sub> and simultaneously allow light from the vertical direction. In the first case, both the liquid-suspension and the gas–solid system can be operated at atmospheric pressure where the CO<sub>2</sub> gas can be purged through a mass flow controller while H<sub>2</sub>O vapor is inserted simultaneously. However, in the second case, only the liquid-suspension system can be operated where an external pressure and elevated temperature can be applied to activate the inert CO<sub>2</sub> gas. Since both the reactors involve a difference in their operating process, it is quite obvious that different efficiency and selectivity will be obtained in the two different reactors.<sup>156</sup> In the first case, light is passed through the reactor and therefore the catalyst encountering the light illumination on the front side will exhibit better activity than the backside one. However, in the gas–solid system, it will face the same light illumination with even intensity and so the obtained activity will be uniform. Therefore, the apparent quantum yield would be different depending upon the photoreactor since photocatalysis depends

on the area of illumination and not on the amount of catalyst. Similarly, in the second case, the catalyst at the bottom will get less light exposure and so its activity will be different from the catalyst placed at the top of the reactor. It is not possible to exactly propose the selectivity of the CO<sub>2</sub> photo-reduction products exactly based on the applied reactor. For example, the pressure sustaining reactor allows CO to stay inside the reactor, which may further undergo hydrogenation, leading to the formation of multiple products. On the contrary, in the continuous flow process, lower numbers of reduced products are achieved as it allows the reactant to spend less time inside the reactor.

**5.1.9 Effect of sacrificial agents.** Ideal CO<sub>2</sub> photoreduction uses water as the medium, where it acts as a sacrificial hole scavenging agent. But this fact totally depends upon the valence band edge position of the catalyst. If the valence band edge potential is more positive than the standard  $E^0(\text{H}_2\text{O}/\text{O}_2)$ , then it can exhibit sacrificial hole scavenging behavior. For the oxidation of one H<sub>2</sub>O → O<sub>2</sub>, four holes are required and the produced O<sub>2</sub> can act as an oxidizing agent, which has an adverse impact on the CO<sub>2</sub> reduction reaction.<sup>141,153</sup> But in several cases, additional sacrificial electron donors like triethanolamine (TEOA), *etc.* are used to facilitate the hole scavenging process, which makes the electrons free to reduce the CO<sub>2</sub> molecule. The hole induced oxidation of TEOA can also produce CO<sub>2</sub> and thus creates a disturbance in the operating reaction conditions. Therefore, the actual source of carbon in the reduced products it inevitably in doubt. Thus, a <sup>13</sup>CO<sub>2</sub> isotope tracing experiment is also important in those cases where TEOA is used as a sacrificial electron donor. In some cases, H<sub>2</sub> gas is used as proton and electron sources respectively. But it is imperative to state that a true CO<sub>2</sub> photoreduction reaction involves H<sub>2</sub>O only as a sacrificial electron donor.

**5.1.10 Effect of a co-catalyst.** Noble metal-based co-catalysts such as Pt, Ag, and Au are widely used over a solid catalyst to add extra momentum to the overall thermodynamics of the CO<sub>2</sub> photoreduction reaction.<sup>157,158</sup> This has a severe influence on both the performance and product selectivity of the CO<sub>2</sub> photoreduction process. It is well documented that when noble metals like Au, Pt, and Pd are decorated over a photocatalyst, a Schottky barrier is created at the junction. At this interface, the metal nanoparticles act as electron traps and prohibit the recombination of the photo-generated charge carriers. Additionally, plasmonic nanoparticles, specifically Au and Ag, facilitate photocatalysts to harness an extended range of visible light and enhance their photocatalytic performance. Also, the localized surface plasmon resonance induces the injection of hot electrons into the conduction band of the appointed photocatalyst and therefore electron accumulation enhances the CO<sub>2</sub> reduction tendency. These exciting features of noble metal co-catalysts are applicable to provide additional benefits in the CO<sub>2</sub> photocatalytic reaction. At the same time, they also enhance the competitive H<sub>2</sub> reduction tendency, which prohibits the CO<sub>2</sub> reduction reaction. But the overall bandgap, as well as the band edge positions, is altered, which provides opportunities to produce different CO<sub>2</sub> photo-reduction products



if it is the case of *ex situ* decoration of noble metals. Another distinctive point to be noted here is that the different affinity towards different interim species over different noble metal nanoparticles truly regulates the product distribution. Since noble metals are costly, their application as a co-catalyst is rather discouraging in ideal reaction conditions. But to obtain better activity and desired selectivity, one can employ such a co-catalyst.

**5.1.11 Stoichiometric ratio of reaction consumed electrons and holes against the yield of the products.** The stoichiometric involvement of electrons and holes in CO<sub>2</sub> reduction and H<sub>2</sub>O oxidation is another important parameter because this will definitely influence the rate of formation of different products after the reaction. For example, the rate of total reactant conversion can be correlated with the rate of the different products formed and the numbers of electrons and holes consumed. For the reduction of 2 CO<sub>2</sub> molecules to produce 2CH<sub>3</sub>OH molecules, 12 electrons are required and, at the same instance, 12 holes must be consumed by the water molecules to produce 6O<sub>2</sub> molecules. This is also a feasible thermodynamic concern in the CO<sub>2</sub> photo-reduction reaction, which can provide a meaningful idea about the photo-reduction and light-induced charge consumption during the reaction.

## 5.2 Graphitic carbon nitride as a CO<sub>2</sub> reduction photocatalyst

The basic phenomenon of CO<sub>2</sub> photo-reduction, especially with respect to heterogeneous catalysis, have already been addressed through several critical, tutorial, perspective, and edge articles by eminent researchers across the globe.<sup>159–165</sup> Most importantly, CO<sub>2</sub> photoreduction over g-C<sub>3</sub>N<sub>4</sub> has also been reviewed recently.<sup>153–165</sup> Still, various aspects of this material and its further manipulation for CO<sub>2</sub> photoreduction catalytic efficiency improvement remain uncovered.

It is pertinent to mention that g-C<sub>3</sub>N<sub>4</sub> has been continuously investigated in different photocatalytic applications since its application as a semiconductor HER photocatalyst by Wang *et al.* in 2009. However, among the seven different types of polymorphic phases (all are semiconductors), only the *g-h*-heptazine and *g-h*-triazine phases possess appropriate bandgaps and band straddling to catalyze various photocatalytic reactions. But, a close look at the valence band edge potential and theoretical calculations confirm that only *g-h*-heptazine fulfills the thermodynamic requirements for water oxidation.<sup>62</sup> Notably, the *g-h*-heptazine based polymorphic g-C<sub>3</sub>N<sub>4</sub> material has been most widely investigated as a photocatalyst with a bandgap of 2.7 eV where the conduction band is situated at (−1.2 V vs. NHE) and the valence band is positioned at (+1.5 V vs. NHE) at pH = 7. The CB and VB indicate that it is capable of reducing the CO<sub>2</sub> molecule and also can oxidize H<sub>2</sub>O to produce O<sub>2</sub> and H<sup>+</sup> required for the production of hydrocarbons/oxygenated hydrocarbons in CO<sub>2</sub> photoreduction. Thus various nanostructured carbon nitrides with diverse morphologies such as nanosheets, microspheres, nanotubes, *etc.* are proven to be efficient catalysts for CO<sub>2</sub> photoreduction. Thermodynamic constraints and optoelectronic limitations are the main bottlenecks of this catalyst for CO<sub>2</sub> photoreduction as simultaneous CO<sub>2</sub> and H<sub>2</sub> reduction tendencies are observed for this particular catalyst.

Also, the rapid charge recombination and low active surface area are other shortcomings that significantly hamper the efficiency of this catalyst. Strategic improvement of the physico-chemical and optoelectronic properties is a pragmatic approach to overcome the poor CO<sub>2</sub> conversion efficiency of pristine g-C<sub>3</sub>N<sub>4</sub>. In the forthcoming sections, a concise summary of the recent literature reports dealing with CO<sub>2</sub> photoreduction efficiency improvement is compiled to give a clear understanding to researchers actively working in this area.

Dong *et al.* demonstrated, for the first time, CO<sub>2</sub> photo-reduction involving porous carbon nitride (PCN) as a catalyst in 2012.<sup>166</sup> In this study, the authors propose that the porosity is introduced in bulk g-C<sub>3</sub>N<sub>4</sub> by changing the precursors from melamine to melamine hydrochloride, which exhibits selective CO production of 4.5 mM in 11 h during CO<sub>2</sub> photoreduction. However, on the contrary, the activity of bulk g-C<sub>3</sub>N<sub>4</sub> is shown to be better (20.6 mM in 11 h) than porous g-C<sub>3</sub>N<sub>4</sub> and this is attributed to the presence of predominant inherent defect sites in PCN compared to bulk g-C<sub>3</sub>N<sub>4</sub> that act as charge carrier recombination sites (Table 3, entry 1).<sup>166</sup> However, this adverse trend in the catalytic activity is unfortunately not fully understandable due to inadequate experimental evidence. Micro-structure dependent CO<sub>2</sub> reduction activity has been reported by Mao *et al.* The authors have observed that urea derived, mesoporous, flake-like g-C<sub>3</sub>N<sub>4</sub> has produced a mixture of oxygenated hydrocarbons (C<sub>2</sub>H<sub>5</sub>OH and CH<sub>3</sub>OH), while melamine derived, non-porous, flaky C<sub>3</sub>N<sub>4</sub> has produced CO and CH<sub>4</sub> as products. The varied product selectivity is correlated to the reaction medium NaHCO<sub>3</sub> and CO<sub>2</sub> gas, where CO<sub>2</sub> is available in the activated form (HCO<sub>3</sub><sup>−</sup>), which is more accessible to the catalyst. In this process, O<sub>2</sub> is obtained as a side product and it is established that the holes and electrons are consumed in a close ratio of ~1 during the reaction with an apparent quantum efficiency (A.Q.E.) of 0.18% at 420 nm (Table 3, entry 2).<sup>167</sup> The fine-tuning of the product selectivity is also reported in different studies, in which g-C<sub>3</sub>N<sub>4</sub> nanosheets produce CH<sub>4</sub> as a selective product, while the bulk g-C<sub>3</sub>N<sub>4</sub> produces CH<sub>3</sub>CHO. This change in the product selectivity is proposed due to the switching of band edge potentials and also due to the different bandgap values. g-C<sub>3</sub>N<sub>4</sub> nanosheets exhibit bandgap enlargement by 0.2 eV, which leads to a more positive valence band edge by 80 meV and a negative shift of the conduction band potential by 120 meV, which are the key reasons behind the different activity and selectivity as compared to bulk g-C<sub>3</sub>N<sub>4</sub> (Table 3, entry 3).<sup>168</sup> Thermal polymerization of pre-synthesized urea derived C<sub>3</sub>N<sub>4</sub> and red phosphorous (P) has formed a heterojunction and, with the aid of Pt as a co-catalyst, the optimized catalyst (PCN-30) has converted CO<sub>2</sub> to CH<sub>4</sub> with a rate of 295 μmol h<sup>−1</sup> g<sup>−1</sup> in the suspension medium. The authors have proposed that the interfacial heterojunction has transferred the electrons from the high position g-C<sub>3</sub>N<sub>4</sub> conduction band to a relatively lower position red P conduction band and the holes have been transferred in the reverse direction, which effectively reduces CO<sub>2</sub> to CH<sub>4</sub> and oxidizes OH to the (OH•) radical, respectively. However, the authors have not emphasized the exact role of Pt involved in the reaction as a co-catalyst



Table 3 Photocatalytic CO<sub>2</sub> reduction reaction over g-C<sub>3</sub>N<sub>4</sub> based photocatalysts

| E. Catalyst/<br>no. amount                                                          | S <sub>BET</sub> (m <sup>2</sup> g <sup>-1</sup> )/<br>precursors and<br>process                 | Co-catalyst                                                                   | Reaction conditions<br>and (CO <sub>2</sub> source)                                               | Light source                              | Products/rate                                                                                              | Time (h) | <sup>13</sup> C<br>isotope CO <sub>2</sub><br>tracing pressure Q.E. | Bandgap and<br>band straddling<br>CO <sub>2</sub> adsorption                                                                                                   | Ref. |
|-------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------|-------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------|-------------------------------------------|------------------------------------------------------------------------------------------------------------|----------|---------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------|------|
| 1. g-C <sub>3</sub> N <sub>4</sub> (100 mg)                                         | —                                                                                                | None                                                                          | CO <sub>2</sub> gas + H <sub>2</sub> O<br>Supercritical fluid                                     | 300 W Xe lamp                             | 20.6 mM CO in 11 h                                                                                         | 11       | No                                                                  | —                                                                                                                                                              | 166  |
| 2. g-C <sub>3</sub> N <sub>4</sub> (200 mg)                                         | 39.5                                                                                             | —                                                                             | grade-CO <sub>2</sub> + NaOH<br>solution                                                          | 300 W Xe lamp                             | 15.1 μmol CH <sub>3</sub> OH +<br>10.8 μmol C <sub>2</sub> H <sub>5</sub> OH +<br>51.2 μmol O <sub>2</sub> | 12       | —                                                                   | 2.78 eV<br>[E <sub>cb</sub> = -1.23 eV,<br>E <sub>vb</sub> not defined]                                                                                        | 167  |
| 3. g-C <sub>3</sub> N <sub>4</sub> nano-<br>sheet (100 mg)                          | (306) dicyandia-<br>mide in thermal<br>oxidation in air                                          | —                                                                             | CO <sub>2</sub> + H <sub>2</sub> O                                                                | 300 W Xe lamp                             | ~0.12 μmol CH <sub>4</sub><br>and ~0.25 μmol<br>CH <sub>3</sub> CHO                                        | 5        | No                                                                  | E <sub>g</sub> = 2.97 eV;<br>E <sub>cb</sub> = (+1.85 eV),<br>E <sub>vb</sub> not defined                                                                      | 168  |
| 4. Red P doped<br>g-C <sub>3</sub> N <sub>4</sub> PCN-30<br>(20 mg)                 | (Not provided)<br>urea derived C <sub>3</sub> N <sub>4</sub><br>combined pyrolysis<br>of red P   | 0.5 wt% Pt                                                                    | CO <sub>2</sub> gas + H <sub>2</sub> O                                                            | 500 W Xe lamp                             | 295 μmol h <sup>-1</sup> g <sup>-1</sup>                                                                   | —        | No                                                                  | Not mentioned                                                                                                                                                  | 169  |
| 5. CN-12-1.5<br>(8 mg)                                                              | (240) (silica hard<br>template/<br>cyanamide)                                                    | Ru co-catalyst<br>(3.9 μmol g <sup>-1</sup> )                                 | CO <sub>2</sub> gas + (H <sub>2</sub> O +<br>TEOA) solution                                       | 400 W high-<br>pressure Hg<br>lamp        | 1854 nmol HCOOH                                                                                            | 5        | Yes                                                                 | E <sub>g</sub> and E <sub>vb</sub> not<br>mentioned; E <sub>cb</sub> =<br>(-1.29 V) vs. Ag/<br>AgCl at pH 6.6                                                  | 170  |
| 6. g-C <sub>3</sub> N <sub>4</sub> nano-<br>sheet CNU-BA <sub>0.03</sub><br>(30 mg) | (54) thermal<br>polymerization<br>of (urea and<br>barbituric acid)                               | (CoCl <sub>2</sub> + 2,2'-<br>bipyridine) as<br>electron donor<br>co-catalyst | CO <sub>2</sub> + H <sub>2</sub> O + (TEOA<br>as electron donor)                                  | 300 W Xe lamp                             | 56.3 μmol CO                                                                                               | 4        | Yes                                                                 | No data available                                                                                                                                              | 171  |
| 7. S-Doped g-C <sub>3</sub> N <sub>4</sub><br>(TCN) (100 mg)                        | (4,4) thermal poly-<br>merization of<br>thiourea at 520 °C<br>in an ambient<br>environment       | 1 wt% Pt                                                                      | (120 mg NaHCO <sub>3</sub> +<br>0.25 mL 4 M HCl) →<br>(CO <sub>2</sub> + H <sub>2</sub> O) vapour | Simulated<br>xenon lamp                   | Yield 1.12 CH <sub>3</sub> OH<br>μmole g <sup>-1</sup>                                                     | 3        | No                                                                  | E <sub>g</sub> = 2.63 eV, no<br>data available for<br>band edge<br>potentials                                                                                  | 172  |
| 8. O-Doped g-C <sub>3</sub> N <sub>4</sub><br>tube (50 mg)                          | (36) melamine in<br>two-step heating<br>in air and N <sub>2</sub>                                | —                                                                             | NaHCO <sub>3</sub> + H <sub>2</sub> SO <sub>4</sub>                                               | Not mentioned<br>(420 nm)                 | 2.6 μmol CH <sub>3</sub> OH                                                                                | 3        | No                                                                  | E <sub>g</sub> = 2.97 eV;<br>E <sub>vb</sub> = not<br>mentioned, E <sub>cb</sub> =<br>-0.87 eV vs. Ag/<br>AgCl at pH = 7                                       | 173  |
| 9. P-Doped g-C <sub>3</sub> N <sub>4</sub><br>nanotube (50<br>mg)                   | (13.4) melamine<br>and sodium hypo-<br>phosphite mono-<br>hydrate in 550 °C<br>4 h <sup>-1</sup> | —                                                                             | NaHCO <sub>3</sub> + H <sub>2</sub> SO <sub>4</sub>                                               | A high-<br>pressure xenon<br>lamp (300 W) | 9.48 μmol CO and<br>7.24 μmol CH <sub>4</sub>                                                              | 4        | No                                                                  | (E <sub>g</sub> = 2.97 eV,<br>E <sub>cb</sub> = -0.90 eV<br>vs. E <sub>NHE</sub> at<br>pH = 7), E <sub>vb</sub> = not<br>mentioned,                            | 174  |
| 10. Phenyl grafted<br>g-C <sub>3</sub> N <sub>4</sub> (100 mg)                      | (94.7) thermal<br>polymerization of<br>urea and phenyl<br>urea at 550 °C for<br>2 h              | —                                                                             | DMF + TEOA +<br>H <sub>2</sub> O + CO <sub>2</sub>                                                | No informa-<br>tion available             | 0.907 μmol CH <sub>4</sub> +<br>450 micromole<br>CH <sub>3</sub> OH + 5 μmol<br>HCOOH                      | 24       | No                                                                  | 0.96% using E <sub>g</sub> = 2.97 eV,<br>20 W LED<br>bulb<br>E <sub>cb</sub> = -1.01 eV,<br>E <sub>vb</sub> = +1.65<br>calculated from<br>Kulbeka monk<br>plot | 175  |
| 11. C <sub>2</sub> O co-doped<br>g-C <sub>3</sub> N <sub>4</sub> (100 mg)           | (50) thermal co-<br>polymerization of<br>(urea + thouria +<br>2-methyl<br>imidazole)             | —                                                                             | (CO <sub>2</sub> + H <sub>2</sub> O) + (TEOA<br>acts as a sacrificial<br>electron donor)          | Xenon lamp<br>(300 W)                     | 4.18 mmol g <sup>-1</sup><br>CH <sub>3</sub> OH                                                            | 6        | No                                                                  | 0.69<br>mmol g <sup>-1</sup>                                                                                                                                   | 110  |



Table 3 (continued)

| E. Catalyst/<br>no. amount                                                                                                               | $S_{\text{BET}}$ ( $\text{m}^2 \text{g}^{-1}$ )/<br>precursors and<br>process                      | Co-catalyst                                                                                                                                  | Reaction conditions<br>and (CO <sub>2</sub> source)                                                                                                                                                           | Light source                                                                               | Products/rate                                                                                                                                           | Time (h) | <sup>13</sup> C<br>isotope CO <sub>2</sub><br>tracing pressure | Q.E.                                                                                | Bandgap and<br>band straddling                                                                                                      | CO <sub>2</sub><br>adsorption                                 | Ref. |
|------------------------------------------------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------|----------|----------------------------------------------------------------|-------------------------------------------------------------------------------------|-------------------------------------------------------------------------------------------------------------------------------------|---------------------------------------------------------------|------|
| 12. mp-g-CN <sub>x</sub> (2 mg)                                                                                                          | (104) cyandiamide<br>and <i>in situ</i><br>synthesized cobalt<br>phthalocyanine                    | CoPPc                                                                                                                                        | CO <sub>2</sub> -Saturated (4 : 1)<br>MeCN/TEOA                                                                                                                                                               | Xenon lamp,<br>100 mW cm <sup>-2</sup> ,<br>AM 1.5G,<br>$\lambda > 400 \text{ nm}$         | 455 $\mu\text{mol g}^{-1} \text{CO}$                                                                                                                    | 24       | Yes                                                            | (EQE) for<br>CO: 0.11%<br>and 0.03%<br>at $\lambda_{\text{ex}} = 360$<br>and 400 nm | $E_g = 2.82 \text{ eV}$ ,<br>$E_{\text{cb}} = -1.10 \text{ eV}$<br>vs. $E_{\text{NHE}}^0$ , $E_{\text{vb}} =$<br>$+1.60 \text{ eV}$ | —                                                             | 176  |
| 13. DCN-0.05 (not<br>mentioned)                                                                                                          | (29.5) one step<br>thermal<br>polymerization of<br>cyandiamide and<br>tartaric acid                | Bi-pyridine +<br>CoCl <sub>2</sub>                                                                                                           | CO <sub>2</sub> gas saturated in<br>5 mL solution (4 mL lamp<br>ACN + 1 mL of<br>TEOA) containing<br>aq. bi-pyridine<br>(10 mmol L <sup>-1</sup> + 25<br>mL of 20 mmol L <sup>-1</sup><br>CoCl <sub>2</sub> ) | 300 W xenon<br>lamp (420 nm<br>cut off filter)                                             | 284.7 $\mu\text{mol g}^{-1} \text{CO}$<br>and<br>51.6 $\mu\text{mol g}^{-1} \text{H}_2$                                                                 | 15       | No                                                             | —                                                                                   | $E_g = 2.49 \text{ eV}$ ,<br>$E_{\text{cb}} = -0.83 \text{ eV}$<br>vs. $E_{\text{RHE}}^0$ , $E_{\text{vb}} =$<br>$+1.66 \text{ eV}$ | —                                                             | 177  |
| 14. BA20-CNS-PDA15<br>nanosphere (not<br>mentioned)                                                                                      | (95) silica template<br>(barbituric acid +<br>polydopamine)                                        | Co(bpy) <sub>3</sub> <sup>2+</sup><br>(150 mg 2,2-<br>bipyridine,<br>+10 $\mu\text{mol}$<br>CoCl <sub>2</sub> )                              | CO <sub>2</sub> gas purged in<br>50 mL suspension<br>(30 mL ACN + 10 mL<br>TEOA + 10 mL H <sub>2</sub> O)                                                                                                     | 300 W xenon<br>lamp (420 nm<br>cut off filter)                                             | 158 $\mu\text{mol CO} + 26$<br>$\mu\text{mol H}_2$ (86%<br>selectivity)                                                                                 | 4        | No                                                             | —                                                                                   | $E_g = 2.63 \text{ eV}$                                                                                                             | Isotherm<br>provided,<br>binding<br>energy did<br>not mention | 178  |
| 15. CN-ATZ-NaK<br>(50 mg)                                                                                                                | (15) (5-amino-<br>tetrazole in a or<br>NaCl/KCl eutectic<br>mixture)                               | —                                                                                                                                            | (0.084 g NaHCO <sub>3</sub> ) + 350 W Xe lamp<br>(0.3 mL 2 M H <sub>2</sub> SO <sub>4</sub> )<br>with an AM1.5<br>filter                                                                                      | 350 W Xe lamp<br>with an AM1.5<br>filter                                                   | 14 $\mu\text{mol g}^{-1} \text{h}^{-1} \text{CO}$ Not<br>mentioned<br>CH <sub>4</sub> + <1 $\mu\text{mol g}^{-1}$<br>h <sup>-1</sup> CH <sub>3</sub> OH | 12       | No                                                             | —                                                                                   | $E_g = 2.53 \text{ eV}$ ,<br>$E_{\text{cb}} = (+1.87 \text{ eV})$ ,<br>$E_{\text{cb}} = (-0.66 \text{ eV})$                         | No data<br>available                                          | 179  |
| 16. O,C co-doped<br>C <sub>3</sub> N <sub>4</sub> (MG <sub>3</sub> )<br>(30 mg)                                                          | (10.8) (melamine +<br>glucose) water<br>dispersion flowed<br>by heating                            | —                                                                                                                                            | CO <sub>2</sub> gas purged in<br>(ACN+ TEOA + H <sub>2</sub> O)                                                                                                                                               | 300 W Xe lamp<br>with a UV cut-<br>off filter<br>>420 nm                                   | 55.2 $\mu\text{mol CO g}^{-1}$                                                                                                                          | 12       | No                                                             | —                                                                                   | $E_g = 2.66 \text{ eV}$ ,<br>$E_{\text{cb}} = (+1.38)$ ,<br>$E_{\text{cb}} = (-1.28 \text{ eV})$                                    | —                                                             | 180  |
| 17. Nitrogen defect-<br>modified g-C <sub>3</sub> N <sub>4</sub> and<br>atomic layers:<br>(CN <sub>550</sub> ) (amount<br>not mentioned) | (103.1) (melamine<br>and NH <sub>4</sub> Cl)<br>heating at different<br>temperature                | —                                                                                                                                            | Catalyst coated on a<br>graphite rod + H <sub>2</sub> O + arc lamp<br>CO <sub>2</sub> gas                                                                                                                     | 500 W xenon<br>assembled<br>with a UV<br>cut-off filter<br>( $\lambda > 400 \text{ nm}$ )  | 9.28 $\mu\text{mol g}^{-1} \text{CH}_4$                                                                                                                 | 8        | No                                                             | 0.089%                                                                              | $E_g = 2.72 \text{ eV}$ ,<br>$E_{\text{cb}} = (+2.17)$ ,<br>$E_{\text{cb}} = (-0.55 \text{ eV})$                                    | —                                                             | 181  |
| 18. 1%B/g-C <sub>3</sub> N <sub>4</sub><br>(amount not<br>mentioned)                                                                     | (62.5) one-step<br>calcination boric<br>acid + urea                                                | —                                                                                                                                            | CO <sub>2</sub> and H <sub>2</sub> O<br>vapor = (H <sub>2</sub> SO <sub>4</sub><br>solution + NaHCO <sub>3</sub><br>powder)                                                                                   | 300 W Xe lamp<br>with a UV/<br>visible cut-off<br>filter<br>( $\lambda > 400 \text{ nm}$ ) | 0.32 $\mu\text{mol g}^{-1} \text{CH}_4$ + 3<br>0.45 $\mu\text{mol g}^{-1} \text{CO}$                                                                    | 3        | Yes                                                            | —                                                                                   | Not provided                                                                                                                        | —                                                             | 182  |
| 19. CNU-TM-13.0<br>(30 mg)                                                                                                               | (109.1) (urea +<br>trimesic acid)                                                                  | Co(bpy) <sub>3</sub> <sup>2+</sup> ;<br>1 $\mu\text{mol}$ of<br>cobalt chlor-<br>ide (CoCl <sub>2</sub> ) +<br>(15 mg of 2,2-<br>bipyridine) | CO <sub>2</sub> gas purged in 5<br>mL solution (1 mL<br>TEOA + H <sub>2</sub> O)                                                                                                                              | Visible light<br>( $\lambda > 420 \text{ nm}$ )<br>filter                                  | 70.63 $\mu\text{mol of CO}$<br>and 14.99 $\mu\text{mol of H}_2$                                                                                         | 4        | Yes                                                            | 1 bar                                                                               | Not provided                                                                                                                        | Isotherm<br>provided,<br>binding<br>energy did<br>not mention | 183  |
| 20. B,K-Doped C <sub>3</sub> N <sub>4</sub><br>(100 mg)                                                                                  | 5.3 (di-cyandia-<br>mide + KBH <sub>4</sub> ) in<br>H <sub>2</sub> thermal etching                 | —                                                                                                                                            | H <sub>2</sub> O + CO <sub>2</sub> gas                                                                                                                                                                        | 300 W Xe lamp                                                                              | 5.93 $\mu\text{mol g}^{-1} \text{CH}_4$ + 5<br>3.16 $\mu\text{mol g}^{-1} \text{CO}$                                                                    | 5        | No                                                             | 2 bar                                                                               | $E_g = 2.35 \text{ eV}$ ,<br>$E_{\text{cb}} = (+1.45)$ ,<br>$E_{\text{cb}} = (-0.90 \text{ eV})$                                    | —                                                             | 184  |
| 21. Tubular yolk-<br>shell g-C <sub>3</sub> N <sub>4</sub><br>(100 mg)                                                                   | (75) (urea +<br>melamine + DI<br>H <sub>2</sub> O) hydrothermal<br>treatment at 180 °C<br>for 24 h | —                                                                                                                                            | supercritical fluid-<br>grade CO <sub>2</sub> gas + H <sub>2</sub> O                                                                                                                                          | 300 W Xe lamp                                                                              | 40.3 $\mu\text{mol g}^{-1} \text{CO}$                                                                                                                   | 4        | Yes                                                            | —                                                                                   | $E_g = 2.72 \text{ eV}$ ,<br>edge potential<br>not mentioned                                                                        | Isotherm<br>provided,<br>binding<br>energy did<br>not mention | 185  |



Table 3 (continued)

| E. Catalyst/<br>no. amount                                                                                                          | $S_{\text{BET}}$ ( $\text{m}^2 \text{g}^{-1}$ )/<br>precursors and<br>process                                                                                                                                         | Co-catalyst                                               | Reaction conditions<br>and ( $\text{CO}_2$ source)                                                                                                                  | Light source                                                                               | Products/rate                                                                                                                                                                                                      | Time (h)          | $^{13}\text{C}$<br>isotope $\text{CO}_2$<br>tracing pressure | Bandgap and<br>band straddling<br>$\text{CO}_2$<br>adsorption                                                                                           | Ref.                                    |
|-------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------------|--------------------------------------------------------------|---------------------------------------------------------------------------------------------------------------------------------------------------------|-----------------------------------------|
| 22. Nitrogen<br>vacancy<br>containing $\text{C}_3\text{N}_4$<br>(CN-550)                                                            | (89.5) thermal<br>combustion of<br>melamine in air<br>followed by $\text{H}_2$<br>atmosphere                                                                                                                          | —                                                         | (High-purity $\text{CO}_2$<br>gas + $\text{H}_2\text{O}$ )                                                                                                          | 300 W xenon<br>lamp equipped<br>with a 420 nm<br>cut-off filter                            | 6.21 $\mu\text{mol h}^{-1} \text{CO}$                                                                                                                                                                              | Not<br>mentioned  | No                                                           | $E_g = 2.77 \text{ eV}$ ,<br>$E_{\text{vb}} = (+1.67)$ ,<br>$E_{\text{cb}} = (-1.07 \text{ eV})$<br>vs. NHE                                             | 146                                     |
| 23. Cyano and<br>P-doped $\text{g-C}_3\text{N}_4$<br>( $\text{g-C}_3\text{N}_4$ -425)<br>(100 mg)                                   | (31.1) urea derived<br>$\text{C}_3\text{N}_4$ mixed with<br>$\text{Na}_2\text{HPO}_4$ followed<br>by heat treatment                                                                                                   | 1 wt% Au                                                  | 1 M 100 mL<br>$\text{NaHCO}_3$ + $\text{CO}_2$ gas                                                                                                                  | 300 W xenon<br>lamp equipped<br>with a<br>( $\lambda = 420 \text{ nm}$ )<br>cut-off filter | 1.135 $\mu\text{mol g}^{-1} \text{CO}$                                                                                                                                                                             | 3                 | Yes                                                          | $E_g = 2.77 \text{ eV}$ ,<br>$E_{\text{vb}} = (+1.68)$ ,<br>$E_{\text{cb}} = (-0.54 \text{ eV})$<br>vs. NHE                                             | 186                                     |
| 24. $\text{CNX}_i$ (30 mg)                                                                                                          | (29.4) (melamine +<br>HX) (X = Cl, Br, I)<br>hydrothermal<br>followed by<br>carbonization in $\text{N}_2$                                                                                                             | —                                                         | $\text{CO}_2$ gas purged in<br>(ACN + TEOA + $\text{H}_2\text{O}$ )                                                                                                 | 300 W xenon<br>lamp with a<br>UV-cut filter<br>( $\lambda = 420 \text{ nm}$ )              | 34.14 $\text{CO} \mu\text{mol g}^{-1}$ ,<br>10.42 $\mu\text{mol g}^{-1} \text{H}_2$ ,<br>1.99 $\mu\text{mol g}^{-1} \text{CH}_4$                                                                                   | 12                | No                                                           | $E_g = 2.83 \text{ eV}$ ,<br>$E_{\text{cb}} = (-1.13 \text{ eV})$<br>vs. NHE, $E_{\text{vb}}$ not<br>defined                                            | 187                                     |
| 25. Pyrene-<br>functionalized<br>polymeric<br>carbon nitride<br>(10% Py-PCN)10<br>mg                                                | (2.11) cyanuric<br>chloride (CC) +<br>cyanamide (CA) +<br>acetonitrile (ACN)<br>hydrothermal<br>treatment followed<br>by rehydrothermal<br>treatment with<br>1-(4,6-dichloro-<br>1,3,5-triazin-2-<br>yl)pyrene (PyCC) | 3 wt% Pt                                                  | Biphasic (1 mL 3 M<br>aqueous solution of<br>$\text{NaHCO}_3$ ) (3 M) and<br>9 mL cyclohexene<br>(or 9 mL toluene<br>containing 2 mmol<br>of 2-methyl-2-<br>butene) | 300 W xenon<br>lamp                                                                        | $\sim 0.7 \mu\text{mol CO}$<br>(using $\text{NaHCO}_3$ +<br>cyclohexane); $\sim 0.15$<br>$\mu\text{mol CO}$ and $\sim 0.23$<br>$\mu\text{mol CH}_4$ (using<br>$\text{NaHCO}_3$ + 2-methyl-<br>2-butene in toluene) | 4                 | No                                                           | $E_g = 2.31 \text{ eV}$ ; $E_{\text{vb}}$<br>$= (+1.15)$ , $E_{\text{cb}} =$<br>$(-1.16 \text{ eV})$ vs.<br>NHE                                         | 188                                     |
| 26. $\text{g-C}_3\text{N}_4$ (catalyst<br>amount not<br>mentioned)                                                                  | (SA analysis not<br>measured) thermal<br>polymerization of<br>melamine                                                                                                                                                | —                                                         | $\text{H}_2\text{O}$ vapour + $\text{CO}_2$<br>gas                                                                                                                  | 100 W xenon<br>lamp                                                                        | $\sim 20 \text{ ppm CO}$                                                                                                                                                                                           | 5                 | No                                                           | $E_g = 2.8 \text{ eV}$ , band<br>edge potential<br>not mentioned                                                                                        | 62                                      |
| Metal modified/doped $\text{g-C}_3\text{N}_4$ for $\text{CO}_2$ reduction                                                           |                                                                                                                                                                                                                       |                                                           |                                                                                                                                                                     |                                                                                            |                                                                                                                                                                                                                    |                   |                                                              |                                                                                                                                                         |                                         |
| 27. Se-Modified poly-<br>meric carbon<br>nitride nano-<br>sheets (m-CNNSS<br>0.08) 50 mg                                            | (133) urea derived<br>CN mixed in an<br>ethanolic solution<br>of diphenyl<br>diselenide (DDS)<br>followed by<br>calcination                                                                                           | (15 mg; 2,2-<br>bipyridine, 1<br>$\mu\text{mol CoCl}_2$ ) | 1 mL TEOA in 5 mL<br>solvent (ACN; 3 mL;<br>$\text{H}_2\text{O}$ , 2 mL), 40 °C                                                                                     | 300 W xenon<br>lamp<br>( $\lambda > 420 \text{ nm}$ )                                      | $\sim 10 \mu\text{mol CO}$ ,<br>$\sim 1.42 \mu\text{mol H}_2$                                                                                                                                                      | 1                 | No                                                           | No information<br>available                                                                                                                             | 189                                     |
| 28. Mg-Doped<br>$\text{g-C}_3\text{N}_4$ Mg/<br>PCN-20%<br>(50 mg)                                                                  | (59.9) urea derived<br>$\text{g-C}_3\text{N}_4$ mixed in an<br>ethanolic solution<br>of Mg ( $\text{NO}_3$ ) $_2$ ·6 $\text{H}_2\text{O}$<br>followed by hydro-<br>thermal treatment                                  | —                                                         | (100 mL $\text{H}_2\text{O}$ + $\text{CO}_2$<br>gas)                                                                                                                | 300 W xenon<br>lamp                                                                        | $\text{CH}_4$ 17.09 $\mu\text{mol g}^{-1}$ ,<br>$\text{CO}$ 4.13 $\mu\text{mol g}^{-1}$                                                                                                                            | 6                 | No                                                           | $E_g = 2.54 \text{ eV}$ , no<br>information<br>available about<br>$\text{g-C}_3\text{N}_4$                                                              | No informa-<br>tion<br>available<br>190 |
| 29. CN1/Pd nano-<br>tetrahedron (NT)<br>(50 mg of the<br>catalyst was<br>suspended in<br>6 mL of DI water<br>in a glass<br>reactor) | Urea derived<br>$\text{g-C}_3\text{N}_4$ followed<br>by seed-mediated<br>polyol-assisted<br>reduction of<br>$\text{Na}_2\text{PdCl}_4$                                                                                | Pd                                                        | ( $\text{CO}_2$ and $\text{H}_2\text{O}$<br>vapor) 0.084 g of<br>$\text{NaHCO}_3$ + 0.3 mL of<br>2 M $\text{H}_2\text{SO}_4$ solution                               | 300 W Xe arc<br>lamp                                                                       | 2.13 $\mu\text{mol h}^{-1} \text{g}^{-1}$<br>$\text{CH}_4$ -production rate<br>of 0.58 $\mu\text{mol h}^{-1} \text{g}^{-1}$ available                                                                              | No<br>information | No                                                           | $E_g = 2.75 \text{ eV}$<br>$\text{CO}_2$ adsorp-<br>tion energy<br>( $E_{\text{ads}} =$<br>$-0.0083 \text{ eV}$ )<br>at the 100<br>surfaces of<br>Pd NT | Theoretical<br>191                      |

Table 3 (continued)

| E. Catalyst/<br>no. amount                                                                                             | $S_{\text{BET}}$ ( $\text{m}^2 \text{g}^{-1}$ )/<br>precursors and<br>process                                                                                                                                                  | Co-catalyst | Reaction conditions<br>and ( $\text{CO}_2$ source) | Light source                                                                        | Products/rate                | Time (h) | $^{13}\text{C}$<br>isotope $\text{CO}_2$<br>tracing pressure | Q.E.                                                                     | Bandgap and<br>band straddling                                                                                             | $\text{CO}_2$<br>adsorption | Ref. |
|------------------------------------------------------------------------------------------------------------------------|--------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|-------------|----------------------------------------------------|-------------------------------------------------------------------------------------|------------------------------|----------|--------------------------------------------------------------|--------------------------------------------------------------------------|----------------------------------------------------------------------------------------------------------------------------|-----------------------------|------|
| 30. Au/P-Doped<br>$\text{g-C}_3\text{N}_4$ (200 mg<br>of the sample<br>was dispersed<br>in 3 mL $\text{H}_2\text{O}$ ) | [Ionic liquid<br>1-butyl-3-methyl<br>imidazoliumhexa-<br>fluorophosphate<br>(BmimPF <sub>6</sub> ) +<br>dicyandiamide<br>(DCDA)] thermal<br>annealing followed<br>by <i>in situ</i> photo-<br>reduction of<br>$\text{HAuCl}_4$ | Au          | ( $\text{H}_2\text{O}$ + $\text{CO}_2$ ) gas       | 300 W xe-lamp                                                                       | 24 $\mu\text{mol CH}_4$      | 5        | No                                                           | $\text{CH}_4 \sim$<br>2.83%,<br>respectively<br>at $\lambda =$<br>420 nm | $E_g = 2.38 \text{ eV}$ ,<br>$E_{\text{vb}} = (+1.4 \text{ eV})$ ,<br>respectively<br>$E_{\text{cb}} = (-0.98 \text{ eV})$ | —                           | 192  |
| 31. Pt/ $\text{g-C}_3\text{N}_4$                                                                                       | (The surface area<br>was not men-<br>tioned) urea<br>derived $\text{g-C}_3\text{N}_4$<br>followed by polyol-<br>assisted reduction<br>of $\text{H}_2\text{PtCl}_6$                                                             | 2 wt% Pt    | ( $\text{H}_2\text{O}$ + $\text{CO}_2$ ) gas       | 15 W bulb,<br>the average<br>intensity at<br>the reactor<br>8.5 $\text{mW cm}^{-2}$ | 13.02 $\mu\text{mol g}^{-1}$ | 10       | No                                                           | —                                                                        | $E_g = 2.8 \text{ eV}$ , No<br>information<br>available about<br>$E_{\text{cb}}$ and $E_{\text{bv}}$                       | —                           | 193  |

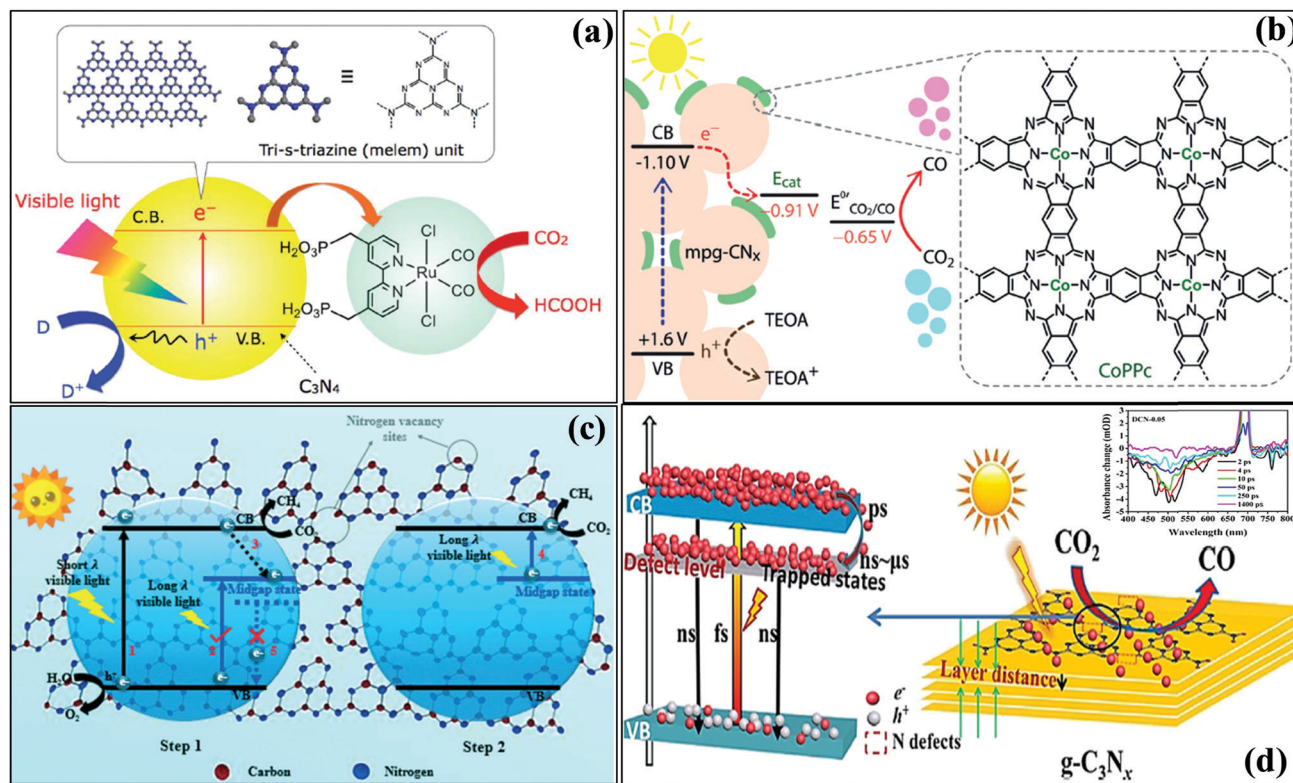
(Table 3, entry 4).<sup>169</sup> K. Maeda *et al.* have synthesized high surface area mesoporous  $\text{g-C}_3\text{N}_4$  using  $\text{SiO}_2$  as a hard template and dicyanamide as a precursor by varying the silica/precursor ratio.<sup>170</sup> The dependency of the pore wall geometry on the  $\text{CO}_2$  photoreduction activity has been uncovered in this study. Furthermore, the resulting mesoporous  $\text{C}_3\text{N}_4$  has been grafted with a Ru complex *trans* (Cl)-[Ru{4,4'-( $\text{CH}_2\text{PO}_3\text{H}_2$ )<sub>2</sub>-2,2'-bipyridine}(CO)<sub>2</sub>Cl<sub>2</sub>], where the electrons from the conduction band of  $\text{g-C}_3\text{N}_4$  are transferred to the Ru and the bare holes in the valence band of  $\text{g-C}_3\text{N}_4$  are engaged in the oxidation of water/triethanolamine, thus completing the total redox cycle (Scheme 15a).<sup>170</sup> The best catalyst of this study exhibits the selective formation of 1854 nmol  $\text{HCOOH}$  in 5 h by involving ( $\text{H}_2\text{O}$  + TEOA +  $\text{CO}_2$ ) in the catalytic system under visible light illumination. The most fascinating outcome of this work is that high surface area is not always beneficial, rather the average surface area ( $240 \text{ m}^2 \text{g}^{-1}$ ) with higher pore volume ( $0.74 \text{ cm}^3 \text{g}^{-1}$ ) has minimized the charge carrier recombination and has shortened the electron-hole diffusion to the reactive sites (Table 3, entry 5).<sup>170</sup> J. Qin *et al.* have investigated the co-polymerization effect of monomer burbutyric acid with urea to synthesize  $\text{g-C}_3\text{N}_4$ , which eventually extends the pi-conjugated network, engineers the molecular hetero-interface and tailors the optical, surface area and nanostructural properties of  $\text{g-C}_3\text{N}_4$  semiconductors for  $\text{CO}_2$  reduction with the involvement of  $\text{Co}(\text{bpy})_3^{2+}$  as a co-catalyst and acetonitrile containing triethanolamine solution as a sacrificial electron donor, and  $\text{CO}_2$  gas.<sup>171</sup> In this study, 56.3  $\mu\text{mol}$  selective CO gas production is achieved in 4 h.  $^{13}\text{C}$  isotope tracing measurements have confirmed that the  $\text{CO}_2$  gas is the sole source of carbon, and, in fact, the control experiment has proved that all the components are essential for triggering  $\text{CO}_2$  photoreduction (Table 3, entry 6).<sup>171</sup> K. Wang *et al.* have reported S-doped  $\text{g-C}_3\text{N}_4$  (TCN) which has been prepared *via* a simple heat combustion method and first-principles calculations suggest that some of the bi-dentate N atoms of the tris-triazine melem framework are replaced by sulfur during this process, having a narrow gap energy ( $E_g = 2.63 \text{ eV}$ ) as compared to the melamine derived bulk  $\text{g-C}_3\text{N}_4$  (2.7 eV). The  $\text{CO}_2$  photoreduction result demonstrates the selective production of methanol with a yield of  $1.25 \mu\text{mol g}^{-1}$  in 3 h from the ( $\text{NaHCO}_3$  +  $\text{HCl}$ ) induced ( $\text{CO}_2$  +  $\text{H}_2\text{O}$ ) vapor medium and Pt co-catalyst (Table 3, entry 7).<sup>172</sup> J. Fu *et al.* have reported O-doped  $\text{C}_3\text{N}_4$  nanotubes from controlled thermal oxidation etching which exhibit selective  $\text{CH}_3\text{OH}$  production with a yield of 2.6  $\mu\text{mol}$  in 3 h from a ( $\text{NaHCO}_3$  +  $\text{H}_2\text{SO}_4$ ) mixture as  $\text{CO}_2$  and proton sources, respectively. The oxygen doping causes a lowering of the conduction band by 0.16 eV by substituting the nitrogen atom of the melem unit, as evidenced by theoretical calculations, which is the prime reason for this selectivity control. Moreover, the higher  $\text{CO}_2$  affinity and  $\text{CO}_2$  uptake are also supported by  $\text{CO}_2$ -TPD and  $\text{CO}_2$ -adsorption analysis. The mechanism of methanol formation has also been confirmed by *in situ* FTIR and GC analysis (Table 3, entry 8).<sup>173</sup> B. Liu *et al.* have synthesized phosphorus-doped  $\text{g-C}_3\text{N}_4$  nanotubes through a one-step thermal condensation between melamine and sodium hypophosphite monohydrate, which simultaneously dopes P and introduces surface amino groups. These modifications have led to higher



CO<sub>2</sub> uptake and favorable CO<sub>2</sub> adsorption, which have been experimentally confirmed by CO<sub>2</sub>-uptake and zeta potential measurements. The resultant material has converted CO<sub>2</sub> to 9.48 μmol CO and 7.24 μmol CH<sub>4</sub> in 3 h. The experimental evidence suggests that phosphorus is interstitially doped into g-C<sub>3</sub>N<sub>4</sub> during the synthesis, narrows the bandgap, exhibits a 1D nanotube morphology with high surface area, enhances the visible light absorption, and promotes the separation of charge carriers, which are the inclusive factors for this enhanced catalytic activity (Table 3, entry 9).<sup>174</sup> D. Vidyasagar *et al.* have synthesized a phenyl grafted g-C<sub>3</sub>N<sub>4</sub> scaffold by reacting phenyl urea, melamine, and urea with enhanced pi-pi stacking, which lowers the bandgap and provides a favorable band edge potential for CO<sub>2</sub> photoconversion from (DMF + TEOA + H<sub>2</sub>O + CO<sub>2</sub> gas) to produce CH<sub>4</sub>, CH<sub>3</sub>OH, and HCOOH with a yield of 0.907 μmol CH<sub>4</sub>, 450 μmol CH<sub>3</sub>OH, and 5 μmol HCOOH in 24 h using a 20 W LED bulb. The achieved apparent quantum yield is calculated to be 0.96% (Table 3, entry 10).<sup>175</sup> S. Samanta *et al.* have synthesized C, O co-doped C<sub>3</sub>N<sub>4</sub> through a thermal poly-condensation of urea, thiourea, and 2-methyl imidazole as starting materials in an air atmosphere. The co-doping has increased the CO<sub>2</sub> uptake, gives a porous cotton-like morphology, lowers the conduction band edge potentials, and also red-shifts the light-harvesting domain. The material converts CO<sub>2</sub> gas saturated (TEOA + H<sub>2</sub>O) solution to methanol with a yield of 4.18 mmol g<sup>-1</sup> in 6 h under visible light illumination. More importantly, theoretical calculations have provided favorable evidence about the high CO<sub>2</sub> binding energy after C, O co-doping, which is also supported by CO<sub>2</sub>-TPD and CO<sub>2</sub>-adsorption studies (Table 3, entry 11).<sup>108</sup> In interesting work, A. Roy *et al.* have synthesized a cobalt phthalocyanine (CoPPC) grafted polymeric graphitic carbon nitride photocatalyst in which the favorable CO<sub>2</sub> electro-reduction capability, as well as the light sensitization properties of cobalt phthalocyanine, has been emphasized. Combining the characteristic features of C<sub>3</sub>N<sub>4</sub> and CoPPC enables the material to reduce CO<sub>2</sub> to CO in organic solvents (ACN + TEOA) under UV/vis light (AM 1.5 G, 100 mW cm<sup>-2</sup>, λ > 300 nm) with a cobalt-based turnover number of 90 for CO after 60 h. More interestingly, in complete aqueous phase reaction conditions, the catalyst exhibits 60 μmol CO corresponding to a TON of 5.1 per CO. When the reaction condition is changed by keeping acetonitrile fixed and replacing TEOA with 10% H<sub>2</sub>O the activity is reduced to 41%, indicating phase separation in aqueous medium (Table 3, entry 12).<sup>178</sup> CoPPC acts as an electron acceptor and lowers the conduction band edge potential of the overall system (Scheme 15b). A <sup>13</sup>C isotope experiment fully agrees that CO<sub>2</sub> gas is the only source of CO in the reaction. The external quantum efficiency (E.Q.E) of CO is 0.11% and 0.03% at λ<sub>ex</sub> = 400 and 360 nm, respectively (Table 3, entry 12).<sup>178</sup> Very recently, a 2D g-C<sub>3</sub>N<sub>4</sub> nanosheet has been prepared *via* one-step tartaric acid-assisted thermal polymerization of dicyandiamide having nitrogen vacancies (g-C<sub>3</sub>N<sub>x</sub>) with preferential location at both three-coordinate N atoms and uncondensed terminal NH<sub>x</sub> species. The material has produced 284.7 μmol g<sup>-1</sup> CO and 51.6 μmol g<sup>-1</sup> H<sub>2</sub> from the CO<sub>2</sub> gas dissolved in (TEOA + ACN) solution and Co(bpy)<sub>3</sub><sup>2+</sup> as a co-catalyst in 15 h. The charge carrier migration and utilization

are demonstrated with transient absorption spectroscopy (Table 3, entry 13) (Scheme 15d).<sup>179</sup> M. Li *et al.* have synthesized polydopamine and barbituric acid co-modified carbon nitride nanospheres through thermal silica template-polymerization followed by a post-modification method, which has produced 158 μmol g<sup>-1</sup> CO and 28 μmol H<sub>2</sub> from CO<sub>2</sub> gas dissolved in aqueous (TEOA + ACN) solution and Co(bpy)<sub>3</sub><sup>2+</sup> as a co-catalyst in 4 h using a 300 W xenon lamp (visible light). The co-modification tunes the band edge potentials and favors CO<sub>2</sub>-adsorption, which are the key factors for the activity improvement (Table 3, entry 14).<sup>180</sup> J. Zhang *et al.* have synthesized grain boundary controlled carbon nitride using 5-amino tetrazole as a high nitrogen-containing organic compound and NaCl/KCl. The synthesized catalyst (CN-ATZ-NaK) exhibits 14 μmol g<sup>-1</sup> h<sup>-1</sup> CO, 1.2 μmol g<sup>-1</sup> h<sup>-1</sup> CH<sub>4</sub>, and <1 μmol g<sup>-1</sup> h<sup>-1</sup> CH<sub>3</sub>OH production rates from an activated CO<sub>2</sub> source (NaHCO<sub>3</sub>) and H<sub>2</sub>SO<sub>4</sub> as a proton donor. The grain boundary tailoring is the main reason for this impressive catalytic activity (Table 3, entry 15).<sup>181</sup> Another carbon and oxygen co-doped g-C<sub>3</sub>N<sub>4</sub> has been achieved by aqueous dispersion of glucose with melamine followed by thermal treatment. The resultant optimized material has produced 55.2 μmol CO g<sup>-1</sup> with nominal amounts of CH<sub>4</sub> and CH<sub>3</sub>OH under visible light from CO<sub>2</sub> dissolved in a (TEOA + ACN + TEOA) reaction mixture in 12 h. Simultaneous co-doping has improved the visible light absorption and given fast electron extraction, which are responsible for this activity enhancement (Table 3, entry 16).<sup>182</sup> Very recently J.-Y. Tiang *et al.* have synthesized nitrogen-vacancy enriched carbon nitride by two-step thermal copolymerization between NH<sub>4</sub>Cl and melamine in a controlled environment, which induces mid-gap band state dangling between the conduction and valence band edge positions. The material has produced 9.28 μmol g<sup>-1</sup> CO in 8 h in the presence of a catalyst-coated graphite rod and (CO<sub>2</sub> + H<sub>2</sub>O) vapors under visible light illumination. The main reasons underlying this enhanced CO<sub>2</sub> reduction activity are the introduction of dense nitrogen vacancies, extended light absorptivity of visible light leading to a wide range of light wavelength due to the strategic mid-gap arising, and prolonged lifetime of the charge carriers trapped inside the midgap. The obtained apparent quantum efficiency is calculated to be 0.089% (Table 3, entry 17) (Scheme 15c).<sup>183</sup> J. Fu *et al.* have reported boron as a dopant that induces charge localization inside the cavity of tris tri-azine by the interaction with the two co-ordinated -(N)<sub>2</sub> at the interstitial position. This claim is also justified by DFT calculations. The resulting 1%B/C<sub>3</sub>N<sub>4</sub> has produced 0.32 μmol g<sup>-1</sup> CH<sub>4</sub> + 0.45 μmol g<sup>-1</sup> CO from (CO<sub>2</sub> + H<sub>2</sub>O) vapor under visible light irradiation in 3 h. A <sup>13</sup>C isotope tracing experiment has supported the CO intermediate end product CH<sub>4</sub> formation (Table 3, entry 18) (J. Fu *et al.*, 2019).<sup>184</sup> Trimesic acid-modified g-C<sub>3</sub>N<sub>4</sub> (CNU-TMU-13) has been synthesized *via* a thermal combustion method which produces 70.63 μmol of CO and 14.99 μmol of H<sub>2</sub> in 5 h from aqueous TEOA saturated with CO<sub>2</sub> gas and Co(bpy)<sub>3</sub><sup>2+</sup> as a co-catalyst. The origin of the carbon in the produced carbon is analyzed by a <sup>13</sup>C isotope experiment and high CO<sub>2</sub> uptake is also in favor of the surface modification

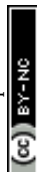




**Scheme 15** (a) CO<sub>2</sub> photoreduction over (a) Ru-complex grafted g-C<sub>3</sub>N<sub>4</sub>, reprinted with permission, Copyright Royal Society of Chemistry, ref. 172; (b) Co(bpy)<sub>3</sub><sup>2+</sup>/g-C<sub>3</sub>N<sub>4</sub>, reprinted with permission, Copyright John Wiley and Sons, ref. 178; (c) N-vacancy containing g-C<sub>3</sub>N<sub>4</sub> with a midgap state, reprinted with permission, Copyright Royal Society of Chemistry, ref. 183; and (d) defect and trap state containing g-C<sub>3</sub>N<sub>4</sub>, reprinted with permission, Copyright John Wiley and Sons, ref. 179.

by trimesic acid, leading to high catalytic activity (Table 3, entry 19).<sup>185</sup> Simultaneous B, K co-doped g-C<sub>3</sub>N<sub>4</sub> has been synthesized using dicyandiamide and KBH<sub>4</sub> under an H<sub>2</sub> thermal etching process, which has accelerated the CO<sub>2</sub> reduction activity with the formation of 5.93 μmol g<sup>-1</sup> CH<sub>4</sub> and 3.16 μmol g<sup>-1</sup> CO from CO<sub>2</sub> dissolved in H<sub>2</sub>O in the absence of any co-catalyst, electron donor, or CO<sub>2</sub> miscibility agent like DMF or acetonitrile in 5 h under visible light. This is also ascribed to the creation of nitrogen-vacancies and the incorporation of B and C together according to the reported theoretical simulation (Table 3, entry 20).<sup>186</sup> The morphology also has a great impact on photocatalytic CO<sub>2</sub> reduction. Tubular yolk-shell g-C<sub>3</sub>N<sub>4</sub> has been hydrothermally synthesized from melamine and urea at 200 °C. The material has produced 40.3 μmol g<sup>-1</sup> CO in 4 h from super-critical grade CO<sub>2</sub> and H<sub>2</sub>O. The formation of CO is also analyzed by a <sup>13</sup>C isotopic experiment. This catalytic improvement is correlated to the higher CO<sub>2</sub> uptake as evident by the CO<sub>2</sub>-uptake profile and also due to the obtained tubular morphology (Table 3, entry 21).<sup>187</sup> Another study reports the synthesis of nitrogen-vacancy containing C<sub>3</sub>N<sub>4</sub> through the controlled polymerization of melamine in different atmospheres, which selectively produces 6.21 μmol h<sup>-1</sup> CO from the catalyst suspension medium of CO<sub>2</sub> and H<sub>2</sub>O under visible light illumination (Table 3, entry 22).<sup>146</sup> The replacement of nitrogen has profoundly controlled selective CO production, which is established *via* DFT calculations

(Table 3, entry 22).<sup>146</sup> In a very strategic approach, X. Liu *et al.* have synthesized cyano-groups and phosphorus-doped C<sub>3</sub>N<sub>4</sub> by treating pre-synthesized carbon nitride with NaH<sub>2</sub>PO<sub>2</sub> under an argon atmosphere at different temperatures. The authors have proposed and proved that the thermal decomposition of NaH<sub>2</sub>PO<sub>2</sub> produced PH<sub>3</sub> gas, which itself was doped as phosphorus and was also accountable for the conversion of the terminal, uncondensed =C-NH<sub>2</sub> to -CN groups during the synthesis process. In fact, the attachment of -CN groups served as electron-withdrawing groups over the carbon nitride surface, which benefited CO<sub>2</sub> activation. The phosphorus doping has enlarged the light absorption and has inhibited photogenerated charge recombination. The active material (g-C<sub>3</sub>N<sub>4</sub>-425) has converted NaHCO<sub>3</sub> saturated CO<sub>2</sub> gas to CO with a yield of ~1.14 μmol g<sup>-1</sup> in 3 h using 1 wt% Au as a co-catalyst under a 300 W xenon lamp (Table 3, entry 23).<sup>188</sup> S. Wan *et al.* have prepared hexagonal tubular carbon nitride by the heat treatment of the supra-molecular organization of the *in situ* hydrogen-bonded melamine cyanuric acid assembly under hydrothermal conditions. Hydrogen fluoride is found to be the best suitable haloic acid (H-X; X = F, Cl, Br, I) that promotes the dissolution of melamine, which eventually facilitated the pi-pi stacking as evidenced by the intense EPR spectra. The light absorption has been blue-shifted and charge separation is maximized with HF. The best active catalyst has produced 34.14 CO μmol g<sup>-1</sup>, 10.42 μmol g<sup>-1</sup> H<sub>2</sub>, and 1.99 μmol g<sup>-1</sup> CH<sub>4</sub> in 12 h under



visible light from 0.8 bar CO<sub>2</sub> gas pressure dissolved in a mixture solution of (acetonitrile + TEOA + H<sub>2</sub>) (Table 3, entry 24).<sup>189</sup> The authors have also calculated the average electron consumption rate ( $R_{\text{electron}}$ ) during the CO<sub>2</sub> reduction reaction as equal to 8.96 mol g<sup>-1</sup> h<sup>-1</sup> on the basis of the equation  $R_E = (2R_{\text{H}_2}) + (2R_{\text{CO}}) + (8R_{\text{CH}_4})$ , where  $R$  is the rate, expressed in μmol g<sup>-1</sup> h<sup>-1</sup> (Table 3, entry 24).<sup>189</sup> For the first time ever, the stability aspect of g-C<sub>3</sub>N<sub>4</sub> in the CO<sub>2</sub> photoreduction reaction has been investigated in-depth by F. R. Poimilla *et al.*, in which the authors provide insights about the stability of the melem framework before and after catalytic activity evaluation (Table 3, entry 25).<sup>62</sup> They have synthesized g-C<sub>3</sub>N<sub>4</sub> by following a two-step thermal polymerization of melamine. For comparison purposes, they have also synthesized O-doped g-C<sub>3</sub>N<sub>4</sub> by treating carbon nitride with H<sub>2</sub>O<sub>2</sub>. They observed a marginal difference in the catalytic activity after repetitive cycles. In gas phase CO<sub>2</sub> reduction, the catalyst exhibits 20 ppm selective CO in 5 h under 100 W xenon light (Table 3, entry 25).<sup>62</sup> But, surprisingly, CO is obtained even in the absence of CO<sub>2</sub> gas under light illumination, which raised the question about the origin of carbon from the graphitic carbon nitride photocatalyst. In association with detailed optoelectronic and physicochemical measurements and also with controlled analysis, the authors concluded that the photo-corrosion rather than the thermal decomposition of melamine is responsible for this unexpected behavior of the catalyst under their experimental conditions. In another breakthrough, X. Gong *et al.* have synthesized pyrene functionalized graphitic carbon nitride, which endows biphasic CO<sub>2</sub> reduction combining CO<sub>2</sub> reduction by the electrons in the aqueous phase and hole generated (OH) radical-induced olefin oxidation in the organic phase. The process of accessibility of the organic olefin into the aqueous system is facilitated by the increased lipophilicity of the grafted pyrene-functionality over g-C<sub>3</sub>N<sub>4</sub>. The authors have achieved ~0.7 μmol CO from (NaHCO<sub>3</sub> + cyclohexane) and (~0.15 μmol CO + ~0.23 μmol CH<sub>4</sub>) from (NaHCO<sub>3</sub> + 2-methyl-2-butene in toluene) under 300 W xenon light illumination in 4 h (Table 3, entry 26).<sup>190</sup>

Various noble and non-noble metal modified carbon nitrides also exhibited profound CO<sub>2</sub> reduction catalytic activity. H. Ou *et al.* have prepared selenium modified thin g-C<sub>3</sub>N<sub>4</sub> porous nanosheets (m-CNNS<sub>s</sub>) using pre-synthesized g-C<sub>3</sub>N<sub>4</sub> in an ethanolic solution of diphenyl diselenide (DDS) followed by thermal polymerization. The evolution of ~10 μmol CO and ~1.42 μmol H<sub>2</sub> is obtained over this catalyst from a reaction mixture containing (TEOA + acetonitrile + H<sub>2</sub>O) at 40 °C in 1 h using 420 nm light irradiation using Co(bpy)<sub>3</sub><sup>2+</sup> as a co-catalyst (Table 3, entry 27).<sup>191</sup> The role of selenium is investigated with density of state calculations, which evidences that selenium has narrowed the conduction band by creating an electronic distribution with the conduction band of CN. But it has not hampered the conduction and valence band composition as its Fermi level exists in the middle of the CB and VB level of g-C<sub>3</sub>N<sub>4</sub>. J. Tang *et al.* have synthesized Mg-doped g-C<sub>3</sub>N<sub>4</sub> considering the high CO<sub>2</sub> philicity of MgO and the light-harnessing capacity of the Mg<sup>2+</sup> ions in natural chlorophyll. The synthesized material has produced 17.09 μmol g<sup>-1</sup> CH<sub>4</sub> and 4.13 μmol g<sup>-1</sup>

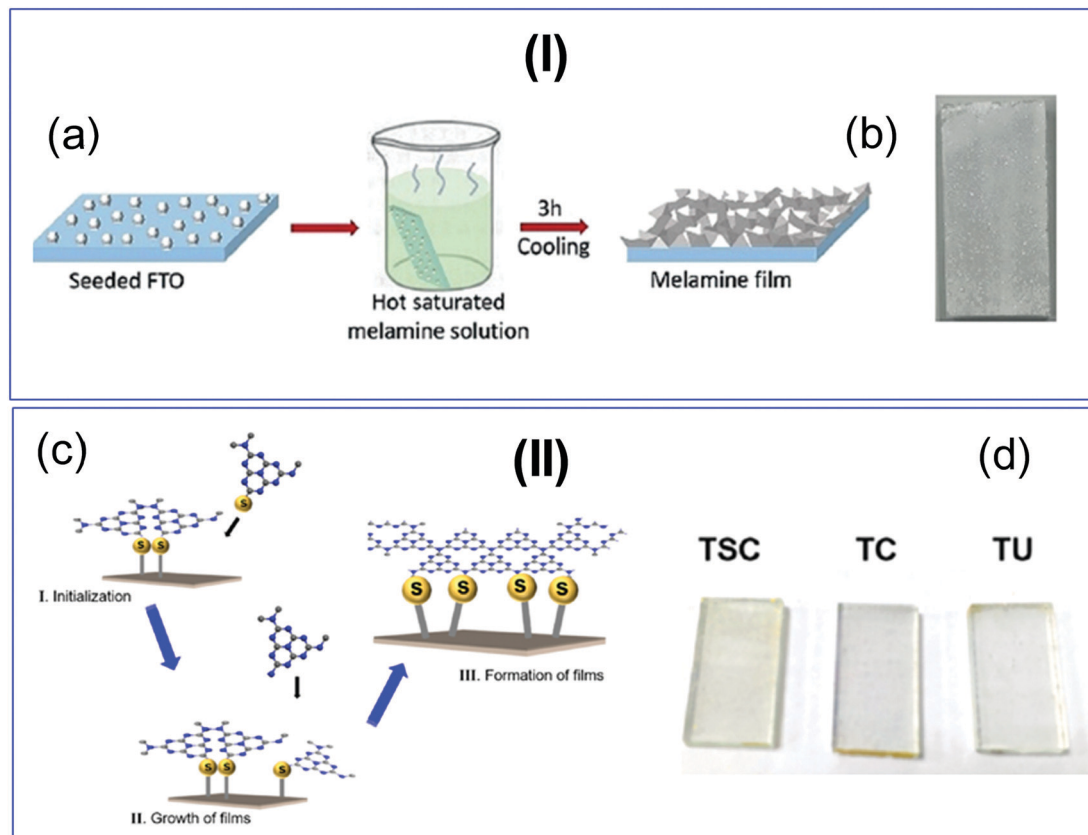
CO from CO<sub>2</sub> gas saturated H<sub>2</sub>O under a 300 W xenon lamp in 6 h (Table 3, entry 28).<sup>192</sup> Though the light-harvesting capacity is enhanced, they did not emphasize the CO<sub>2</sub> uptake phenomenon with any experimental evidence. S. Cao *et al.* have prepared Pd nano-tetrahedron doped g-C<sub>3</sub>N<sub>4</sub> from urea derived g-C<sub>3</sub>N<sub>4</sub> followed by seed-mediated polyol-assisted reduction of Na<sub>2</sub>PdCl<sub>4</sub>. They focused on the facet dependent CO<sub>2</sub> reduction catalytic activity of Pd NT/g-C<sub>3</sub>N<sub>4</sub>, which exhibits a CH<sub>3</sub>OH and CH<sub>4</sub> production rate of 2.13 μmol h<sup>-1</sup> g<sup>-1</sup> and 0.58 μmol h<sup>-1</sup> g<sup>-1</sup>, respectively, from activated (CO<sub>2</sub> + H<sub>2</sub>O) vapor under 300 W visible light illumination (Table 3, entry 29).<sup>193</sup> Furthermore, DFT calculations confirm that the Pd(100) facet over g-C<sub>3</sub>N<sub>4</sub> exhibits high CO<sub>2</sub> uptake and high CO<sub>2</sub> binding energy and thus facilitates CH<sub>3</sub>OH desorption. M. Humayun *et al.* have synthesized Au NP loaded P-doped g-C<sub>3</sub>N<sub>4</sub> in ionic liquid followed by thermal annealing of a mixture of [1-butyl-3-methyl imidazoliumhexafluorophosphate (BmimPF<sub>6</sub>) + dicyandiamide (DCDA)] and then finally the *in situ* photoreduction of HAuCl<sub>4</sub> for the decoration of Au NPs. The catalyst exhibits selective CH<sub>4</sub> evolution of 24 μmol in 5 h from CO<sub>2</sub> gas saturated H<sub>2</sub>O under 300 W visible light (Table 3, entry 30).<sup>194</sup> The calculated quantum efficiency for CH<sub>4</sub> formation at 420 nm is ~2.83%. The impressive catalytic activity is ascribed to the extended visible-light absorption along with the improved charge carrier separation *via* the phosphorus induced surface states and the Au nanoparticle decoration. The impact of Pt on the CO<sub>2</sub> reduction over g-C<sub>3</sub>N<sub>4</sub> is revealed by W.-J. Ong *et al.*, using 2 wt% Pt loaded g-C<sub>3</sub>N<sub>4</sub> nanocomposites. The catalyst exhibits the highest CH<sub>4</sub> yield of 13.02 μmol g<sup>-1</sup> catalyst<sup>-1</sup> after 10 h of light irradiation, which is 5.1-fold higher in comparison with pure g-C<sub>3</sub>N<sub>4</sub> (2.55 μmol g<sup>-1</sup> catalyst<sup>-1</sup>), from CO<sub>2</sub> gas dissolved in H<sub>2</sub>O under a 300 W xenon lamp (Table 3, entry 31).<sup>195</sup> The enhancement in the photocatalytic activity of the Pt/g-C<sub>3</sub>N<sub>4</sub> nanocomposite for CO<sub>2</sub> photo-reduction CH<sub>4</sub> production can be attributed to the enhanced visible light absorption and smooth interfacial photogenerated electron transfer from g-C<sub>3</sub>N<sub>4</sub> to Pt by lowering the Fermi level of Pt in the Pt/g-C<sub>3</sub>N<sub>4</sub> hybrid as evidenced by the UV-vis and photoluminescence studies. The accumulated electron density on Pt facilitates the reduction of CO<sub>2</sub> to CH<sub>4</sub> *via* a multi-electron transfer process.

### 5.3 Critical factors regarding catalytic activity assessment and result presentation

Amidst the CO<sub>2</sub> photoreduction reports published in several journals, there is great concern regarding the precise catalytic activity evaluation, supportive characterization, and data presentation. Despite addressing this matter in several perspectives, edge articles, and editorials, this problem still persists.<sup>196–200</sup> Although the intention is not to indicate or draw attention to any inappropriateness about the results reported on g-C<sub>3</sub>N<sub>4</sub> for CO<sub>2</sub> photoreduction, the aim is to emphasize and regulate the standard requirements for a particular report that can be followed precisely by the readers. The following points may be considered during an experiment and result presentation. If g-C<sub>3</sub>N<sub>4</sub> is the appointed photocatalyst, the abundance or source of carbon in the resultant product may be ensured with







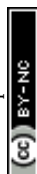
**Scheme 16** (I) (a) Fabrication of a crystalline carbon nitride based-PEC device, and (b) the resultant FTO based device, reprinted with permission, Copyright John Wiley and Sons, ref. 201; and (II) (c) a polymeric carbon nitride-based adhesive connected-PEC device, and (d) corresponding images of the FTO coated PEC devices synthesized from different precursors: TSC-thiosemicarbazide, TC-tri-thiocyanuric acid, TU-thiourea and urea, respectively, reprinted with permission, Copyright American Chemical Society, ref. 204.

on an electrode material with the loss of its absolute linearity; (ii) cleavage of the C–O bonds to form C–H bonds by electron transfer and/or proton migration processes; and (iii) desorption of the product preceded by configurational rearrangement and

followed by diffusion into the electrolyte.<sup>207</sup> Further, it is found that both the e-CO<sub>2</sub>RR and pe-CO<sub>2</sub>RR involve multiple steps with overall two-, four-, six-, or eight-electron transfer processes (Scheme 17).<sup>208</sup> Naturally, the range of products that form



**Scheme 17** Various products formed due to multi-step processes during electrochemical CO<sub>2</sub> reduction.



during such processes is diverse. They include carbon monoxide, formic acid or formate, formaldehyde, methanol, ethanol, methane, ethylene *etc.* The half-cell reactions involved in these processes along with their standard electrode potentials have already been listed in Section 5.1.1.

### 6.1 The various rudiments of an e-CO<sub>2</sub>RR experiment

Based on the mechanistic pathways of the abovementioned processes, electrocatalysts can be tailored for efficient performance and high selectivity. The performance of various electrocatalysts can be judged by electrochemical techniques such as linear sweep voltammetry (LSV), cyclic voltammetry (CV) and electrochemical impedance spectroscopy (EIS). While the overpotentials can be calculated for different products at various current densities directly from the LSV and CV profiles, the kinetics can be understood using the Tafel slopes that can be derived from the relevant voltammograms and/or the impedance spectra. All experiments should be performed using a two-compartment H-shaped electrochemical cell separating the cathode and the anode. This separation can be achieved using a thin frit glass membrane or a semi-permeable conducting membrane such as that of Nafion. A detailed gas-leak-proof setup design has been proposed by Pander *et al.*<sup>209</sup> Since e-CO<sub>2</sub>RR involves the reduction of CO<sub>2</sub>, the working electrode is always a cathode which is capable of reducing other gaseous contaminants such as O<sub>2</sub> and N<sub>2</sub>. Therefore, the e-CO<sub>2</sub>RR process should be preceded by the removal of air from the cathodic chamber. Subsequently, to accurately determine the products formed during the e-CO<sub>2</sub>RR, it is inevitable to saturate the cathodic chamber with high purity CO<sub>2</sub>. The electrolyte is generally KHCO<sub>3</sub> and the counter-electrode (anode in this case) is graphite or a carbon electrode. In several cases, researchers have demonstrated the e-CO<sub>2</sub>RR in KOH medium in order to elevate the pH, and, thereby, suppress the reduction of protons/water (hydrogen evolution). However, saturation of the KOH electrolyte with CO<sub>2</sub> reduces its pH due to the spontaneous formation of K<sub>2</sub>CO<sub>3</sub> and KHCO<sub>3</sub>. Although the pH is maintained between 6.8 and 7.5 under normal conditions, the exact pH of the medium should be monitored to correctly convert the potential *vs.* reference electrode (commonly, a pre-calibrated Ag/AgCl or SCE) to the potential *vs.* reversible hydrogen electrode (RHE). The reference electrode is placed in the cathode chamber of the H-cell. The working electrode should be kept protected from metal impurities present in the electrolyte, such as that of Fe and Zn, in order to avoid poisoning and severe deterioration of the catalytic performance. For this, the aqueous electrolyte can be pre-electrolyzed so as to eradicate the contaminants by depositing them on a sacrificial cathode.<sup>210</sup> Chelex is another option to prevent metal ion deposition due to its chelating action. The current densities that are reported for electrochemical experiments such as CV, LSV *etc.* should be normalized with respect to the electrochemical surface area (ECSA), which can be calculated as per an earlier report.<sup>211</sup> To evaluate the faradaic efficiencies (FE) of various electrocatalysts, the aforesaid electrochemical measurements are generally correlated with the

chromatograms obtained from GC (gas chromatography) and/or HPLC (high-performance liquid chromatography) of the product mixtures. A bulk-electrolysis experiment should be carried out for about an hour to obtain products that are enough for further analysis.<sup>212</sup> It is wise to carry out experiments under potentiostatic conditions to avoid deviations from the product selectivity. Galvanostatic experiments are likely to generate potentials that are not suitable for the e-CO<sub>2</sub>RR, or can generate a variety of products. However, the latter may be preferred only under certain special conditions where the working electrode is studied well in advance and the applied current is precise for scaling up e-CO<sub>2</sub>RR processes.<sup>209</sup> The complexity of e-CO<sub>2</sub>RR experiments, therefore, lies in the varied physical states of the products that are generated in the reaction medium. In the separate evaluation of the FE for various products, the faradaic selectivity or faradaic yield, defined as the fraction of the FE for a given product with respect to the total FE for all products, should also be calculated and reported. For the analysis of the H<sub>2</sub> gas which is invariably evolved during e-CO<sub>2</sub>RR experiments due to very close potentials, the evolving gases should be directly delivered to the sampling loop of an online gas chromatograph equipped with a thermal conductivity detector (TCD) using argon as the inert carrier gas. Besides, a flame ionization detector (FID) should be used to analyze the possible carbon-containing gaseous products such as CO, CH<sub>4</sub>, C<sub>2</sub>H<sub>4</sub>, and C<sub>2</sub>H<sub>6</sub> as described in the mechanistic pathway (Scheme 17). On identification of the constituents present in the liquid (*e.g.*, HCOOH, CH<sub>3</sub>OH, and C<sub>2</sub>H<sub>5</sub>OH) and gaseous mixtures, the selectivity of the product can be evaluated. Relevantly, nuclear magnetic resonance (NMR) spectroscopy and mass spectroscopy (MS) are often used to confirm the associated products. However, to minimize interfering proton signals arising from the aqueous electrolyte, a solvent-suppression method (suppresses the solvent peak, such as water) and deuterium locking (usually done to every NMR sample before acquisition of the spectra) should be necessarily adopted during NMR measurements.<sup>209</sup> As the mechanistic pathway involved in the catalysis over a given heterogeneous surface may be completely changed with the change in the nanoarchitecture of various carbon-based electrodes, structural characterizations of the electrode materials are also important. These include Fourier transform-infrared (FT-IR) spectroscopy, Raman spectroscopy, X-ray diffraction measurements (XRD), X-ray photoelectron spectroscopy (XPS), X-ray absorption near-edge spectroscopy (XANES), X-ray absorption fine structure (XAFS), scanning electron microscopy (SEM), transmission electron microscopy (TEM) and aberration-corrected-scanning transmission electron microscopy (AC-STEM). Even when coupled with active metallic catalyst nanoparticles, various carbon materials may often be seen generating very different products than what can be expected from the purely metallic catalysts. Hence, it is important to understand the structure-activity relationships of the active catalysts with the reactant and product molecules. For detailed analyses of these processes, *in situ* techniques are often used. To understand the source of CO<sub>2</sub> especially with carbon-based electrocatalysts or



Table 4 Electro catalysts and electrocatalytic parameters of the CO<sub>2</sub> reduction reaction

| E. no. | Electrocatalyst                                                     | Electrolyte              | Major product (F.E.%, optimal potential) | Onset potential      | Ref. |
|--------|---------------------------------------------------------------------|--------------------------|------------------------------------------|----------------------|------|
| 1.     | 2D-pg-C <sub>3</sub> N <sub>4</sub>                                 | 2 M NaHCO <sub>3</sub>   | CO (~80%, -0.35 V vs. RHE)               | -0.13 V (vs. RHE)    | 214  |
| 2.     | Au-BCN <sub>3.0</sub> (gold decorated boron-doped carbon nitride)   | 0.5 M NaHCO <sub>3</sub> | Ethanol (78%)                            | 0.15 V (vs. Ag/AgCl) | 215  |
| 3.     | BCN <sub>3.0</sub> (boron-doped carbon nitride)                     | 0.5 M NaHCO <sub>3</sub> | Ethanol (47%)                            | 0.15 V (vs. Ag/AgCl) | 217  |
| 4.     | g-C <sub>3</sub> N <sub>4</sub> /MWCNT                              | 0.5 M NaHCO <sub>3</sub> | CO (~60%, -0.75 V vs. RHE)               | —                    | 217  |
| 5.     | Cu-C <sub>3</sub> N <sub>4</sub>                                    | 0.1 M KHCO <sub>3</sub>  | —                                        | —                    | 219  |
| 6.     | Au/C <sub>3</sub> N <sub>4</sub>                                    | 0.5 M KHCO <sub>3</sub>  | CO (90%, -0.45 V vs. RHE)                | —                    | 220  |
| 7.     | Au-CDot-C <sub>3</sub> N <sub>4</sub>                               | 0.5 M KHCO <sub>3</sub>  | CO (79.8%)                               | -0.3 V               | 221  |
| 8.     | OA-PCN (hydroxyl and amino functionalized polymeric carbon nitride) | 0.5 M NaHCO <sub>3</sub> | CO (40.1%)                               | -0.56 V              | 222  |

electrode materials/substrates, it is often important to carry out isotopic <sup>13</sup>C-labelling experiments where <sup>13</sup>CO<sub>2</sub> is used as the feed gas instead of CO<sub>2</sub>. Since the focus of this article is on C<sub>3</sub>N<sub>4</sub>-based electrodes that have been studied for CO<sub>2</sub> utilization to date, a relevant discussion in the said direction is imperative. Studies related to other carbon-based materials or carbon-free catalysts have however been regarded to exist beyond the scope of the present article.

## 6.2 Impact of g-C<sub>3</sub>N<sub>4</sub> as an interesting e-CO<sub>2</sub>RR catalyst

In recent times, heteroatom doping in carbon materials has evolved as an interesting domain of research to fine-tune their catalytic performances. For example, Sagara *et al.* have reported a boron-doped g-C<sub>3</sub>N<sub>4</sub> photocathode catalyst that produces ethanol as the major product from CO<sub>2</sub> in an aqueous phase with or without the loading of metal nanoparticle co-catalysts Au, Ag or Rh for activity enhancement. It has been generally understood through this study that the enhanced production of ethanol is an outcome of restrained recombination of C1 moieties and the increased selectivity owing to metal nanoparticle loading could be due to the multi-electron reduction by electron capture on metal particles.<sup>213</sup> However, CO is generally the major product when Au or Ag is used as the catalyst, whereas alcohols are formed over Cu electrocatalysts.<sup>214</sup> Zhang *et al.* have reported a polarized few-layer metal-free g-C<sub>3</sub>N<sub>4</sub> electrocatalyst that can selectively form CO (with trace HCOOH) from CO<sub>2</sub> with a high faradaic efficiency (F.E.) of ~80% (Table 4, entry 1).<sup>214</sup> This electrocatalyst has been prepared using hydrothermal pre-treatment of dicyandiamide followed by thermal exfoliation to reduce the number of stray basic sites. This catalyst is able to suppress the hydrogen evolution reaction and, thereby, efficiently catalyze CO<sub>2</sub> reduction. The authors have argued that the large surface area and mesoporous structure of the benchmark photocatalyst mpg-C<sub>3</sub>N<sub>4</sub> (mesoporous graphitic carbon nitride) are not capable of electroreduction of CO<sub>2</sub> whereas the hierarchical structure of the foam-like monolith composed of 2D-pg-C<sub>3</sub>N<sub>4</sub> is proven to facilitate the diffusion of carbon dioxide molecules to the active surface of the catalyst. Electrochemical impedance spectroscopy, further, has justified the improved mass transfer efficiency of 2D-pg-C<sub>3</sub>N<sub>4</sub> exhibiting a larger slope in the low-frequency range (Table 4, entry 2).<sup>215</sup> This result is better than an earlier report on a metal-free g-C<sub>3</sub>N<sub>4</sub>/MWCNT electrocatalyst that converts CO<sub>2</sub> to CO. Of the total 98% faradaic efficiency obtained for H<sub>2</sub> and CO,

H<sub>2</sub> accounts for 58% of the total FE, while that of CO is only 41% (Table 4, entry 3).<sup>216</sup> Zhao *et al.* on the other hand have reported a ternary Au-CDot-C<sub>3</sub>N<sub>4</sub> electrocatalyst with 4 wt% Au that can reduce CO<sub>2</sub> to CO selectively with an F.E. of ~79.8% at a potential of -0.5 V (vs. RHE). Employing theoretical calculations, the authors have proven the importance of the interaction between the CDots and C<sub>3</sub>N<sub>4</sub>. The authors have suggested the enhancement of the catalytic activity to be a direct outcome of the superior adsorption capability of CDots for both H<sup>+</sup> and CO<sub>2</sub>. A narrower bandgap of 1.35 eV due to the newly introduced valence band maximum has been observed in CDot-C<sub>3</sub>N<sub>4</sub> in contrast to pure g-C<sub>3</sub>N<sub>4</sub> (1.92 eV). Again, with the introduction of AuNPs, the electrocatalyst Au-CDot-C<sub>3</sub>N<sub>4</sub> has shown a further decrease in the bandgap to 0.32 eV (Table 4, entry 3).<sup>216</sup> Also, adsorption studies have shown better capability of the 4 wt% Au-CDot-C<sub>3</sub>N<sub>4</sub> electrocatalyst to adsorb both CO<sub>2</sub> (~0.34 mmol g<sup>-1</sup> at 1.2 atm) and H<sup>+</sup> (0.37 g g<sup>-1</sup>, on the basis of HCl) in comparison to 4 wt% Au-C<sub>3</sub>N<sub>4</sub> (~0.22 mmol g<sup>-1</sup> at 1.2 atm of CO<sub>2</sub> and 0.18 mmol g<sup>-1</sup> of H<sup>+</sup> on the basis of HCl) affirming the necessity of CDot incorporation (Table 4, entry 3).<sup>217</sup> Employing Cu as the most widely investigated probe for CO<sub>2</sub> reduction, Jiao *et al.* have studied g-C<sub>3</sub>N<sub>4</sub> as a molecular scaffold that appropriately modifies the electronic structure of Cu in the resultant Cu-C<sub>3</sub>N<sub>4</sub> complex. Through a combination of DFT computation and experimental justification, their work contributes to understanding the high selectivity of the e-CO<sub>2</sub>RR toward hydrocarbons/alcohols. The existence of 'mock chemisorption' on the nitrogen atom of the C<sub>3</sub>N<sub>4</sub> scaffold is suggested in the study, where CO<sub>2</sub> is adsorbed through a carbon-nitrogen bond on Cu-C<sub>3</sub>N<sub>4</sub>. This is believed to afford Cu-C<sub>3</sub>N<sub>4</sub> to activate CO<sub>2</sub> with the application of a suitable electrode potential. Further, the C-centres of the C<sub>3</sub>N<sub>4</sub> framework are shown to have a higher ability to bind with species that ligate using oxygen atoms such as \*OCH<sub>2</sub>, \*OCH<sub>3</sub>, \*O, and \*OH. This is due to the electrophilicity in the C-atoms of C<sub>3</sub>N<sub>4</sub>, which are connected to nitrogen atoms with high electronegativity. Moreover, the Cu active site is shown to have a higher preference to bind with intermediates, \*COOH, \*CO and \*CHO, that use the C-centre for coordination. This kind of dual active center model has been demonstrated to uplift copper's d-orbital position toward the Fermi level, which is critical in adsorbing the intermediate for CO<sub>2</sub> activation. Their work successfully validates the formation of C2 products with the presence of two catalytically active centers (Table 4, entry 4).<sup>218</sup> In contrast to this mechanism for CO<sub>2</sub> activation,



Xu *et al.* have unraveled the intriguing role of nitrogen species in the carbon framework during the electroreduction of CO<sub>2</sub> (Table 4, entry 5).<sup>219</sup> They have enumerated the activity of four types of N-species present in N-doped carbon nanotubes: pyrrolic, pyridinic, and quaternary nitrogen. A conjugated outcome of systematic DFT calculations and experimental observations suggested that the improved activity in N-doped CNTs, in contrast to undoped CNTs, is due to the stabilization of the key intermediate CO<sub>2</sub><sup>•−</sup>. The best correlation is found to be the presence of quaternary N-species (Table 4, entry 5).<sup>219</sup> Recently, Zhang *et al.* have corroborated that carbon nitride supported Au nanoparticles (Au/C<sub>3</sub>N<sub>4</sub>) are better for CO<sub>2</sub> electroreduction than carbon-supported Au nanoparticles (Au/C). This is consistent with an earlier report<sup>220</sup> and the study has been extended to the interaction of Ag with C<sub>3</sub>N<sub>4</sub>. Experimental and theoretical data prove that C<sub>3</sub>N<sub>4</sub> intensifies the electron-density over the Au surface, strengthening the adsorption of key reaction intermediate \*COOH. The F.E. for CO is found to be maintained above 90% over a wide potential window of −0.45 V to −0.85 V *vs.* RHE for Au/C<sub>3</sub>N<sub>4</sub>, while that for Au/C is always lesser than 80% in this range (Table 4, entry 6).<sup>220</sup> This result is comparable to a recently reported electrocatalyst Au-Py-CNT-O which reportedly reduces CO<sub>2</sub> to CO with 93% F.E. over a wide potential range (−0.58 to −0.98 V) (Table 4, entry 7).<sup>221</sup> Meng *et al.*, in a recent report, have shown that a surface-modification strategy to introduce hydroxyl and amino groups on polymeric carbon nitride (PCN) can successfully be used to develop a metal-free electrocatalyst which is able to enhance the CO yield rate 17.1-fold as compared to its unmodified counterpart. The modification process has been carried out by treating PCN at 520 °C for 6 h in a controlled airflow to obtain O-PCN (PCN containing only hydroxyl species). Further, for 1 h O-PCN has been annealed in an NH<sub>3</sub> atmosphere to obtain OA-PCN (containing amino groups along with hydroxyl groups). The catalytic efficiency of OA-PCN towards CO<sub>2</sub> electroreduction is compared with that of A-PCN (containing only amino groups), O-PCN, and PCN. It is observed that the introduction of surface hydroxyl or amino groups alone cannot boost the e-CO<sub>2</sub>RR rate but the one with both functional groups has been found to suppress hydrogen evolution, thereby enhancing the rate of CO production. The CO/H<sub>2</sub> ratio for the OA-PCN electrocatalyst is determined to be 0.67, which is in the range for the Fischer-Tropsch reaction (0.25–3.34). To investigate the ostensibly magical role of dual functionalization of PCN in the e-CO<sub>2</sub>RR, density functional theory (DFT) calculations have been carried out. It is revealed that the conversion of CO<sub>2</sub> to COOH\* possessed the highest energy barrier of 0.62 eV at the NH<sub>2</sub> terminal. Furthermore, a higher barrier of 0.83 eV is required, at the −OH terminal, to desorb the CO product. Therefore, mono-functionalization does not boost the overall performance. However, according to the theoretical calculations, it is possible to infer that a proton–electron transfer process initiates the generation of the \*COOH intermediate at the −OH terminal, which is transferred to the −NH<sub>2</sub> terminal and, then, converted to \*CO. The more facile desorption of the CO product is believed to occur at the −NH<sub>2</sub> terminal, thereby exhibiting a fascinatingly high performance of the dual functionalized OA-PCN electrocatalyst

(Table 4, entry 8).<sup>222</sup> From the perspective of consistency, reproducibility and ease of synthesis, C<sub>3</sub>N<sub>4</sub>-based materials should, therefore, be advocated as efficient potential electrocatalysts for CO<sub>2</sub> reduction.

## 7. Overall summary, conclusions, and future perspectives

In summary, this article covered three important CO<sub>2</sub> conversion processes, thermal CO<sub>2</sub> activation, photocatalytic CO<sub>2</sub> conversion, and electrochemical CO<sub>2</sub> reduction, focused on the rising star carbonaceous material ‘g-C<sub>3</sub>N<sub>4</sub>’. A summary indicating the general characteristics, advantages, and disadvantages of the thermocatalytic/photocatalytic/electrocatalytic CO<sub>2</sub> conversion process is provided in Table 5.

The merits of g-C<sub>3</sub>N<sub>4</sub> for the above mentioned processes are: (a) It is a thermally stable, non-toxic, easy to synthesize material at moderate temperature (550 °C), whereas other carbon-based catalysts need high temperature (1200–1400 °C) and also require chemical activation. (b) In contrast to other carbon materials, N-doped carbon materials, especially g-C<sub>3</sub>N<sub>4</sub>, are more effective for acidic CO<sub>2</sub> activation and conversion. Tailored synthesis parameters, varied carbon and nitrogen precursors, and different synthesis strategies allowed us to synthesize g-C<sub>3</sub>N<sub>4</sub> with different types of N species having Lewis and Brønsted basic sites (uncondensed −NH<sub>2</sub>) suitable for CO<sub>2</sub> activation and utilization. Moreover, it is also possible to incorporate other functional sites in g-C<sub>3</sub>N<sub>4</sub> so that a multi-functional catalyst for the appointed CO<sub>2</sub> activation reactions can be designed to deliver a designated product. (c) g-C<sub>3</sub>N<sub>4</sub> is the only carbon catalyst having a bandgap energy with appropriate valence and conduction band edge potentials for the utilization of visible light of the solar spectrum. It can reduce CO<sub>2</sub> and oxidize H<sub>2</sub>O simultaneously and thus effectively serve in the CO<sub>2</sub> photoreduction reaction. Except for carbon dots, all other carbons do not have any bandgap and, therefore, are photocatalytically inactive for the CO<sub>2</sub> reduction reaction. However, carbon dots cannot themselves trigger CO<sub>2</sub> reduction; instead, they need another support material eventually to exhibit their up-conversion photoluminescence property. (d) It is easy to functionalize g-C<sub>3</sub>N<sub>4</sub> to improve the visible-light harnessing capability, which is not possible with other carbonaceous materials.

Some of the demerits of g-C<sub>3</sub>N<sub>4</sub> are: (a) It is thermally stable, but it is prone to light-induced framework oxidation/decomposition. If CO<sub>2</sub> is a reactant, then it requires deep investigation because the framework oxidation/decomposition product and CO<sub>2</sub> derived product could give misleading information. Therefore, integrating g-C<sub>3</sub>N<sub>4</sub> with another semiconductor is necessary to stop its framework degradation caused by light absorption. (b) It contains less oxygen in the form of hydroxyl groups, which is not enough to make it a steam reforming chemical looping catalyst. (c) It has low electrochemical conductivity and thus exhibits inferior activity for electrocatalytic CO<sub>2</sub> reduction. Moreover, in the absence of an

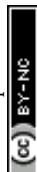


Table 5 Summary of the thermocatalytic/photocatalytic/electrocatalytic CO<sub>2</sub> conversion process

| Process          | General characteristics                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  | Advantages                                                                                                                                                                                                                                                                                                                                                                                                                                                 | Disadvantages                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                            |
|------------------|----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|
| Thermocatalytic  | <p>(1) Thermal energy is the driving force to disrupt the stable atomic and molecular bonds of CO<sub>2</sub> and other reactants over a bi-functional catalyst to produce various platform chemicals and fuels.</p> <p>(2) Industrially available high-pressure liquid phase and gas phase reactors can be employed to convert CO<sub>2</sub> to important feedstock chemicals and fuels under the influence of mild to high CO<sub>2</sub> pressure (1–500 bar or so).</p> <p>(3) Catalysts can be easily designed for the appointed applications.</p> | <p>(1) Easy to implement in large scale production.</p> <p>(2) A variety of chemicals and building blocks can be produced.</p> <p>(3) Catalyst recovery is easy.</p> <p>(4) More than 90% of CO<sub>2</sub> is converted <i>via</i> this process.</p> <p>(5) Can be applied in a broad range of pressure (1–500 bar or so).</p> <p>(6) Thermodynamic and kinetic parameters can be easily tuned.</p>                                                       | <p>(1) Not sustainable due to the involvement of a non-renewable energy source.</p> <p>(2) The trade-off between conversion and energy efficiency is very low.</p> <p>(3) The requirement of high pressure and temperature.</p> <p>(4) Pure CO<sub>2</sub> gas is required.</p> <p>(5) Atmospheric CO<sub>2</sub> gas cannot be converted.</p>                                                                                                                                                                                                                                                                                                                                                                                                                                           |
| Photocatalytic   | <p>(1) The driving force is light-energy under ambient conditions to bring down the thermodynamic constraint of stable CO<sub>2</sub> molecules over a semiconductor photocatalyst having efficient photon capturing ability.</p> <p>(2) It depends on the chemisorption magnitude of CO<sub>2</sub> at <math>[P/P_0 = 1]</math>.</p> <p>(3) The design of a photocatalyst is more-challenging than the thermocatalytic process.</p>                                                                                                                     | <p>(1) Light energy is available abundantly.</p> <p>(2) Eco-friendly and sustainable process.</p> <p>(3) A selective product can be achieved since one-pot CO<sub>2</sub> reduction and H<sub>2</sub>O oxidation takes place.</p> <p>(4) The source of CO<sub>2</sub> can be varied and, accordingly, different products can be achieved.</p> <p>(5) Recyclability is high if a powder suspension system is employed.</p>                                  | <p>(1) A custom-made catalytic reactor coupled with a suitable light source makes the process less user-friendly.</p> <p>(2) Low light-harnessing and quick charge carrier recombination of the photocatalyst are a major drawback.</p> <p>(3) Product isolation and separation are comparatively tough.</p> <p>(4) Mechanistic investigation is somewhat difficult.</p> <p>(5) The competitive reduction tendency between CO<sub>2</sub> and H<sup>+</sup> by the light-induced electrons suppresses the overall activity.</p> <p>(6) Large-scale implementation and night time operation are not possible due to the absence of sunlight.</p> <p>(7) Only low CO<sub>2</sub> pressure can be employed (&lt;10 bar).</p> <p>(8) Atmospheric CO<sub>2</sub> gas cannot be converted.</p> |
| Electrocatalytic | <p>(1) A suitable external potential is a driving force (under ambient conditions) to bring down the thermodynamic constraint of a stable CO<sub>2</sub> molecule over a highly electrochemically conductive electrocatalyst.</p> <p>(2) Mainly carried out under alkaline solution as a CO<sub>2</sub> source.</p> <p>(3) Electrolyte and potential bias both play a key role.</p> <p>(4) The design of an electrocatalyst is highly challenging.</p>                                                                                                   | <p>(1) Easy to operate.</p> <p>(2) Eco-friendly and sustainable process.</p> <p>(3) Highly economic.</p> <p>(4) The trade-off between conversion and energy efficiency is very high.</p> <p>(5) Product selectivity can be finely tuned.</p> <p>(6) Less dependent on the CO<sub>2</sub> chemisorption process.</p> <p>(7) The liquid phase reaction medium is primarily followed.</p> <p>(8) High turnover number (TON) and turnover frequency (TOF).</p> | <p>(1) It is extremely difficult to design the electrocatalytic reactor.</p> <p>(2) An appropriate membrane is difficult to synthesize.</p> <p>(3) A redox-active metal is required owing to its inherent redox properties.</p> <p>(4) Catalyst recovery is very difficult if a thin-film is used.</p> <p>(5) Product quantification from a basic medium by GC, GC-MS, and HPLC leads to low yield.</p> <p>(6) Only low CO<sub>2</sub> pressure can be employed (&lt;10 bar).</p> <p>(7) Atmospheric CO<sub>2</sub> gas cannot be converted.</p> <p>(8) Mechanistic investigation is difficult.</p>                                                                                                                                                                                      |



active metal, it exhibits limited redox activity in electrochemical CO<sub>2</sub> reduction. Other carbonaceous materials, primarily N-doped carbons, are more suitable for electrochemical CO<sub>2</sub> reduction owing to their superior electronic conductivity. (d) Porous carbon can be derived from any natural resources, but specific carbon and nitrogen-containing precursors are required for g-C<sub>3</sub>N<sub>4</sub> synthesis. Moreover, it is a visible light active catalyst and cannot cover the UV and NIR regions of the solar spectrum, which is also a drawback for its limited photocatalytic CO<sub>2</sub> reduction activity.

The most crucial feature of g-C<sub>3</sub>N<sub>4</sub> is its facile synthesis, which can be performed without any complex synthesis setup. But plenty of research is required to improve its catalytic CO<sub>2</sub> conversion efficiency. g-C<sub>3</sub>N<sub>4</sub> exhibited activity in all these three processes but with limited efficiency. For example, thermal CO<sub>2</sub> activation needs high temperature and pressure, which is not sustainable. Moreover, the involvement of an added co-catalyst raises the question of its effectiveness. Integrating an alkali metal oxide with the g-C<sub>3</sub>N<sub>4</sub> framework can enhance CO<sub>2</sub> adsorption and activation. Therefore, further improvement of g-C<sub>3</sub>N<sub>4</sub> based catalysts is required for CO<sub>2</sub> ambient condition activation and utilization. Since g-C<sub>3</sub>N<sub>4</sub> does not have enough oxygen content, integrating it with transition metal oxides like CeO<sub>2</sub> and MnO<sub>2</sub>, or spinel based oxides like NiCo<sub>2</sub>O<sub>4</sub> with high oxygen exchange capacity can make it an active catalyst for steam reforming of methane coupled with the CO<sub>2</sub> splitting reaction. In the case of photocatalytic CO<sub>2</sub> reduction, the target of mole scale product yield is not yet achieved. Moreover, selectivity control and inclusion of additional organic additives like acetonitrile and dimethylformamide, and sacrificial agents like triethanolamine along with a co-catalyst are the serious drawbacks of this photocatalytic process. It is noteworthy that an ideal CO<sub>2</sub> photoreduction system involves a light-harvesting material, CO<sub>2</sub> gas, and H<sub>2</sub>O only. But very rarely has this target been served by this catalyst to date. Another controversial aspect is the source of carbon in the converted product during CO<sub>2</sub> photo-reduction, especially when the catalyst itself is a carbon-based material. Utilization of suitable precursors during thermal polymerization (or other synthesis strategies) can incorporate required C, N, O, or S functionalities that have the capability to alter its solar light harnessing capability and can improve the charge carrier separation suitable for high photocatalytic CO<sub>2</sub> reduction reaction activity. Another vital area is the configuration of thin-film based-device fabrication for practical CO<sub>2</sub> photoreduction. g-C<sub>3</sub>N<sub>4</sub> has a triazine skeleton, and therefore single-metal-atoms can be entrapped in the framework *via* wet impregnation followed by thermal annealing in a reducing environment. It will minimize attrition and deactivation, which are advantages for catalyst long term stability. High overpotential, low current densities, and limited redox-tunability are the shortcomings in its electrochemical CO<sub>2</sub> reduction application. Therefore these concerns should be carefully considered for its further efficiency improvement. Carbon nitride-based single-atom-catalysts can be a viable solution to improve photocatalytic, electrocatalytic, or photoelectrocatalytic CO<sub>2</sub> reduction.

We have concisely covered almost all the parameters starting from the synthesis of materials to fundamentals of CO<sub>2</sub>

activation, followed by critical operational parameters for collective 'thermo-photo-electro'-chemical CO<sub>2</sub> conversion focusing on the g-C<sub>3</sub>N<sub>4</sub> catalyst. The conclusion section has provided the details of the merits and demerits of g-C<sub>3</sub>N<sub>4</sub> along with a concise perspective on the application of carbon nitride for CO<sub>2</sub> conversion. We envisage that this review will be helpful for beginners in their early stage of research on heterogeneous CO<sub>2</sub> conversion reactions for further technological advancement in the 21st century.

## Author contributions

The theme of the review article was proposed by SS and RS jointly. The manuscript was co-written by S. S. and R. S. The authors thank Mr Aniruddha Mukherjee for the preparation of some figures and helpful discussion and drafting the electrochemical section. All the authors proofread and approved the final version of the manuscript.

## Conflicts of interest

The authors state that no competitive and non-competitive conflicts of interest exist to declare. The authors also declare that some of the figures included in this manuscript have been prepared by themselves with the appropriate citations wherever it is applicable. Copyright permission has been obtained for the required figures of this manuscript.

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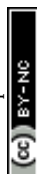
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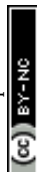
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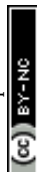
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