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# A robust Fe-based heterogeneous photocatalyst for the visible-light-mediated selective reduction of an impure CO<sub>2</sub> stream<sup>†</sup>

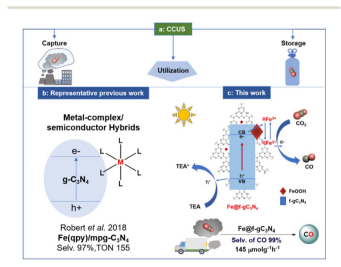
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The transformation of  $CO_2$  into value-added products from an impure  $CO_2$  stream, such as flue gas or exhaust gas, directly contributes to the principle of carbon capture and utilization (CCU). Thus, we have developed a robust iron-based heterogeneous photocatalyst that can convert the exhaust gas from the car into CO with an exceptional production rate of 145  $\mu$ mol g<sup>-1</sup> h<sup>-1</sup>. We characterized this photocatalyst by PXRD, XPS, ssNMR, EXAFS, XANES, HR-TEM, and further provided mechanistic experiments, and multi-scale/level computational studies. We have reached a clear understanding of its properties and performance that indicates that this highly robust photocatalyst could be used to design an efficient visible-light-mediated reduction strategy for the transformation of impure  $CO_2$  streams into value-added products.

#### Introduction

The development of iron-based catalysts is attractive due to the high abundance of iron in the Earth's crust and low cost compared to the other transition metals. Furthermore, iron can adopt diverse oxidation states (from -2 to +5) and can promote single electron transfer reactions. These advantages triggered scientists to develop novel iron-based homo-/heterogeneous catalysts, and many of them are comparable to

those of the 4d and 5d-based transition metal analogs. <sup>10-13</sup> Parallel to the development of iron-based catalysts, the direct transformation of CO<sub>2</sub> into value-added products has tremendous potential because CO<sub>2</sub> is non-toxic and abundant in the atmosphere. <sup>14-27</sup> However, in most cases, only pure CO<sub>2</sub> streams are used as carbon sources. If impure CO<sub>2</sub> streams such as flue gas from industries or exhaust gas from a car could be utilized, it will avoid the associated cost and energy requirement for the



Scheme 1 Reported photocatalytic approaches and our work for the reduction of carbon dioxide.

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 ${\rm CO_2}$  purification procedure. This new approach contributes to the Carbon Capture and Utilization (CCU) principle (Scheme 1a).<sup>28-39</sup> Nonetheless, the presence of impurities such as  ${\rm O_2}$ , water vapor, CO,  ${\rm NO_x}$ , and hydrocarbons in the impure  ${\rm CO_2}$  stream can be harmful to the photocatalyst and can be detrimental to the desired product formation.<sup>40-44</sup>

Recently, graphitic carbon nitrides (g-C<sub>3</sub>N<sub>4</sub>) have become highly attractive as photocatalysts due to their high chemical and thermal stability, appropriate band structures, and low cost.45-51 Thus, researchers have employed g-C3N4-based hybrid photocatalysts (g-C<sub>3</sub>N<sub>4</sub> combined with Ru,<sup>52</sup> Co,<sup>53-56</sup> and Fe<sup>57-59</sup>based metal complexes) to improve the selectivity and reactivity for the reduction of CO<sub>2</sub> into CO (Scheme 1b). Among them, Cobased transition metal complexes such as [Co(bpy)<sub>3</sub>]<sup>2+</sup> and Co(qpy) along with g-C<sub>3</sub>N<sub>4</sub>, exhibited the selectivity of 86 and 98% (TON of CO were 3.7 and 128, with the production rate of 37 and 7.98 μmol g<sup>-1</sup> h<sup>-1</sup> respectively) for the formation of CO in the presence of triethanolamine (TEOA) and 1,3-dimethyl-2phenyl-2,3-dihydro-1*H*-benzo[*d*]imidazole (BIH) as sacrificial reductant. 54,56 Additionally, Fe-based metal complexes such as Fe(qpy)<sub>3</sub> and Fe(qpy)-BA exhibited the formation of CO with a production rate of 91 and 141 μmol g<sup>-1</sup> h<sup>-1</sup> respectively (selectivity was 97 and 95% in the presence of TEOA and TEA + BIH as sacrificial reductant). 57,58 However, these hybrid systems always required expensive ligands and multistep synthetic routes (associated with the metal complexes), exhibited poor recyclability, and rarely showed reactivity toward the transformation of an impure CO<sub>2</sub> stream.

To achieve a robust and recyclable Fe-based photocatalyst, construction of composite material could be a promising route since this could address the rapid recombination of photogenerated carriers which are the common limitations for the photocatalytic reactivity of g-C<sub>3</sub>N<sub>4</sub>.60,61 Along this direction, amorphous iron-based oxyhydroxides (FeOOH), known as promising Fenton-like catalysts, should be ideal for hybridization due to their small particle size. 62-64 The coupling of amorphous FeOOH with g-C<sub>3</sub>N<sub>4</sub> could be a sound strategy for constructing an effective recyclable photocatalyst. Considering this, we first modified the g-C<sub>3</sub>N<sub>4</sub> moiety by introducing an aryl functionality (f-gC<sub>3</sub>N<sub>4</sub>) to improve the photocatalytic properties of g-C<sub>3</sub>N<sub>4</sub> (ref. 49) and later, FeOOH was introduced into the moiety to fabricate a FeOOH/f-gC<sub>3</sub>N<sub>4</sub> composite photocatalyst. This photocatalyst generated CO with a production rate of 304 μmol g<sup>-1</sup> h<sup>-1</sup> with a selectivity of 99%. Expediently, when exhaust gas from a car was applied, CO was formed with a production rate of 145 μmol g<sup>-1</sup> h<sup>-1</sup> (Scheme 1c), thanks to the enhanced interfacial electron transfer between FeOOH and f-gC<sub>3</sub>N<sub>4</sub>. To the best of our knowledge, this is the first Fe-based recyclable photocatalytic system that can be applied for the reduction of impure CO2 stream.

#### Results and discussion

At the beginning of this project, g-C<sub>3</sub>N<sub>4</sub> was synthesized by calcinating dicyandiamide (DCDA) at 550 °C for 4 h (temp. increasing rate = 2.2 °C min $^{-1}$ ) in a tube furnace under aerobic conditions (please see the detailed procedure in the ESI†).<sup>49</sup>

Followed by this method, functionalized graphitic carbon nitrides (f-gC<sub>3</sub>N<sub>4</sub>) were achieved by stirring a mixture of 9 gm of DCDA and 150 mg of 2-amino-5-trifluoromethyl benzonitrile in deionized water (45 mL) at 95 °C until it's completely dried, resulting mixture was then grinded in an algae mortar and was calcined at 550 °C for 4 h (temp. increasing rate = 2.2 °C min<sup>-1</sup>) under aerobic conditions (please see in the ESI†). Later, Fe(NO<sub>3</sub>)<sub>3</sub>·9H<sub>2</sub>O (7.23 mg for 0.2Fe@f-gC<sub>3</sub>N<sub>4</sub> 18 mg for 0.5Fe@f $gC_3N_4$ , 25.3 mg for 0.7Fe@f- $gC_3N_4$ , and 36 mg for 1Fe@f- $gC_3N_4$ ) and f-gC<sub>3</sub>N<sub>4</sub> (500 mg) were mixed in 10 mL deionized water and the reaction mixture was further stirred at 100 °C (please see in the ESI† for the detailed procedure). It should be noted that 0.2Fe@f-gC<sub>3</sub>N<sub>4</sub>, 0.5Fe@f-gC<sub>3</sub>N<sub>4</sub>, 0.7Fe@f-gC<sub>3</sub>N<sub>4</sub> and 1Fe@fgC<sub>3</sub>N<sub>4</sub> denotes 0.2, 0.5, 0.7 and 1 wt% of iron loading on f-gC<sub>3</sub>N<sub>4</sub> respectively. After the synthesis of all these photocatalysts, the Tauc plot exhibited that gC<sub>3</sub>N<sub>4</sub>, f-gC<sub>3</sub>N<sub>4</sub>, 0.5Fe@g-C<sub>3</sub>N<sub>4</sub>, 1Fe@fgC<sub>3</sub>N<sub>4</sub>, 0.7Fe@f-gC<sub>3</sub>N<sub>4</sub>, 0.5Fe@f-gC<sub>3</sub>N<sub>4</sub>, and 0.2Fe@f-gC<sub>3</sub>N<sub>4</sub> had the band gap of 2.64, 2.48, 2.46, 2.34, 2.47, 2.46 and 2.46 eV, respectively (Fig. S3†). Furthermore, Mott-Schottky plots of all these photocatalysts disclosed that the flat band (fb) potential of  $gC_3N_4$ ,  $f-gC_3N_4$ ,  $0.5Fe@g-C_3N_4$ ,  $1Fe@f-gC_3N_4$ ,  $0.7Fe@f-gC_3N_4$ , 0.5Fe@f-gC<sub>3</sub>N<sub>4</sub>, and 0.2Fe@f-gC<sub>3</sub>N<sub>4</sub> were -0.34, -0.44, -0.47, -0.52, -0.56, -0.33 and -0.38 eV vs. Normal Hydrogen Electrode (NHE). Additionally, the positive slope indicated the ntype nature of these semiconductors (Fig. S4†). The conduction band (CB) of an n-type inorganic semiconductor is commonly assumed to be  $\approx -0.2$  V negative than the flat band potentials. Thus, the CB potentials were derived by lowering the flat band potential by 0.2 V compared to the NHE (Fig. S5†). 47,65

After the synthesis, photocatalytic experiments were carried out in 4 mL of CO<sub>2</sub>-saturated acetonitrile solution in the presence of a freshly distilled sacrificial electron donor (ACN: triethylamine, 4:1 V/V), under the irradiation of a Kessil lamp for 18 h ( $\lambda = 427$  nm, 100 mW cm<sup>-2</sup> light intensity, Table S1†). Indeed, CO was the primary product, with a minor quantity of CH<sub>4</sub> and H<sub>2</sub> In this photocatalytic reaction, TEA got oxidized to form TEAH<sup>+</sup>.58 To our observation, only a trace amount of CO and H<sub>2</sub> were obtained in the case of g-C<sub>3</sub>N<sub>4</sub>, while a moderate amount of CO with 91% selectivity was observed in the presence f-gC<sub>3</sub>N<sub>4</sub>. Furthermore, when 0.5 wt% of Fe was deposited onto both gC<sub>3</sub>N<sub>4</sub> and f-gC<sub>3</sub>N<sub>4</sub>, the Fe@f-gC<sub>3</sub>N<sub>4</sub> system exhibited nearly 12 times higher production rate of CO. To investigate the superior role of 0.5 wt% Fe@f-gC<sub>3</sub>N<sub>4</sub> photocatalyst, pure Fe(NO<sub>3</sub>)<sub>3</sub>·9H<sub>2</sub>O was mixed externally with f-gC<sub>3</sub>N<sub>4</sub> (iron content was the same as 0.5Fe@f-gC<sub>3</sub>N<sub>4</sub>). A lower quantity of CO clearly confirmed the importance of the deposition of iron onto fgC<sub>3</sub>N<sub>4</sub> structure. It could be due to the fact that the metal deposition enhanced the charge transfer efficiency from the conduction band of f-gC<sub>3</sub>N<sub>4</sub> to the active metal site of Fe<sup>+n</sup> and that was ideal for the effective reduction of CO2.57 After that, we were able to further increase the catalytic reactivity through different loadings of iron (0.2-1 wt%) on f-gC<sub>3</sub>N<sub>4</sub> (Fig. 1a and b). We observed that the production rate of CO was linearly increased up to 0.7 wt% and was drastically boosted for 1Fe@f $gC_3N_4$  (172  $\mu$ mol  $g^{-1}$   $h^{-1}$ ). Indeed, iron sites are prone to adsorb CO2; therefore, with the increase of iron loadings, more electrons will be transferred to the iron sites to reduce CO<sub>2</sub>. Thus,

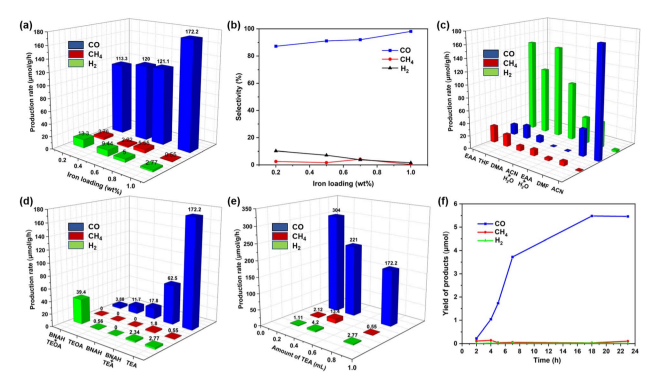


Fig. 1 (a) Reduction of  $CO_2$  by using different loading of iron catalysts; (b) selectivity in  $CO_2$  reduced products by using different loading of iron catalysts; (a and b) reaction conditions: iron catalyst (1 mg), TEA (0.8 mL), ACN (3.2 mL), time = 18 h,  $\lambda$  = 427 nm, reaction temperature = 30 °C. (c) Photocatalytic  $CO_2$  reduction by using different solvents. Reaction conditions: 1Fe@f-gC<sub>3</sub>N<sub>4</sub> (1 mg), TEA (0.8 mL), different solvents (3.2 mL), time = 18 h,  $\lambda$  = 427 nm, reaction temperature = 30 °C. (d) Photocatalytic reduction of  $CO_2$  by using different reductants. Reaction conditions: 1Fe@f-gC<sub>3</sub>N<sub>4</sub> (1 mg), different reductant for single reductant system TEA or TEOA (0.8 mL), 0.1 M BNAH, for binary reductant system BNAH (32 mg) + TEOA (0.4 mL), BNAH (32 mg) + TEA (0.4 mL), ACN (3.2 mL), time = 18 h,  $\lambda$  = 427 nm, reaction temperature = 30° (e) production rate of photocatalytic  $CO_2$  reduction by using different amount of TEA. Reaction conditions: catalyst 1Fe@f-gC<sub>3</sub>N<sub>4</sub> (1 mg), TEA (0.2, 0.4, 0.8 mL), solvent ACN (3.2-3.8 mL), time = 18 h,  $\lambda$  = 427 nm, reaction temperature = 30 °C. (f) Kinetic studies for the photocatalytic reduction of pure  $CO_2$ . Reaction conditions: 1Fe@f-gC<sub>3</sub>N<sub>4</sub> (1 mg), TEA (0.2 mL), ACN (3.8 mL), time = 2-23 h,  $\lambda$  = 427 nm, reaction temperature = 30 °C.

the increased Fe-loading prolonged the lifetime of the charge carriers and enhanced the transfer of the photogenerated electrons from f-gC<sub>3</sub>N<sub>4</sub> to the Fe<sup>+n</sup> center and then to CO<sub>2</sub>. Instead, a smaller number of electrons were transferred toward proton reduction, which in turn suppressed H<sub>2</sub> evolution,<sup>67</sup> and the production rate and selectivity to CO were linearly increased with the increase of iron loading.

Then, we evaluated the importance of different solvents (CH<sub>3</sub>CN, DMF, DMA, EtOAc, THF and H<sub>2</sub>O) in the presence of 1Fe@f-gC<sub>3</sub>N<sub>4</sub> (Fig. 1c). While all of them were favorable for this transformation, no reaction took place in pure water, and further addition of water to organic solvents such as CH<sub>3</sub>CN and EtOAc resulted in a lower evolution rate of CO. Surprisingly, in the aqueous binary solvent system, CH<sub>4</sub> and H<sub>2</sub> were the major products, and CO was the minor product (Fig. 1c). In fact, by adding 37% of water into CH<sub>3</sub>CN and EtOAc, the selectivity of the CO<sub>2</sub> reduction product was changed entirely from CO to CH<sub>4</sub> (8e<sup>-</sup>/8H<sup>+</sup> reduction process) with a production rate of 9.30 and 3.53  $\mu$ mol g<sup>-1</sup> h<sup>-1</sup> respectively. This could be attributed to the fact that the addition of water increased the number of available protons in the solution, which in turn took part in the CO<sub>2</sub> reduction process to form CH<sub>4</sub>.68 Nevertheless, among all the solvents, CH<sub>3</sub>CN was the best for the photochemical reduction of CO<sub>2</sub> to CO, with a high production rate of 172.2 μmol g<sup>-1</sup> h<sup>-1</sup> and an excellent selectivity of 98%. Additionally, sacrificial reductants such as triethylamine (TEA), triethanolamine (TEOA) and 1-benzyl-1,4-dihydronicotinamide (BNAH) were also investigated and a production rate of 172, 11.7, and 17.8 μmol g<sup>-1</sup> h<sup>-1</sup> with 98, 95, and 100% selectivity were obtained (Fig. 1d). Further investigations by using different amounts of TEA and reducing the amount of TEA to 0.2 mL resulted in an excellent production rate of 304 μmol g<sup>-1</sup> h<sup>-1</sup> (Fig. 1e). We argued that a higher concentration of TEA probably quenched the excited state of the photocatalyst and decreased the photocatalytic efficiency.<sup>69</sup>

Furthermore, control experiments suggested that in the absence of  $CO_2$ , photocatalyst, and light, no formation of CO was observed (Fig. S8†). On the other hand, in the absence of TEA, the formation of CO was observed, but with a lower production rate of 76.6  $\mu$ mol g<sup>-1</sup> h<sup>-1</sup>. In these conditions, the kinetics of  $CO_2$  reduction exhibited a linear increase in CO production up to 18 h, and after this, the yield of CO remained constant but the production of  $CH_4$  was slightly increased. Furthermore, kinetic studies demonstrated that the evolution of CO was stable up to 23 h, which was comparable with the recently reported photocatalysts (Fig. 1f).<sup>57,70</sup>

To demonstrate the application of this chemistry, the exhaust gas (containing impurities besides CO<sub>2</sub>, are shown in

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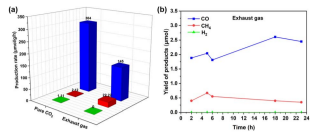


Fig. 2 (a) Comparison of photocatalytic CO $_2$  reduction by using pure CO $_2$  and by using exhaust gas. Reaction conditions: 1Fe@f-gC $_3$ N $_4$  (1 mg), TEA (0.2 mL), ACN (3.8 mL), time = 18 h,  $\lambda$  = 427 nm, reaction temp. = 30 °C. (b) Kinetic studies for the photocatalytic reduction of exhaust gas. Reaction conditions: 1Fe@f-gC $_3$ N $_4$  (1 mg), TEA (0.2 mL), ACN (3.8 mL),  $\lambda$  = 427 nm, reaction temp. = 30 °C.

Fig. S7†) was directly collected from a vehicle by using gas sampling bags (Fig. S6†) and was applied directly under these reaction conditions, showing a CO production rate of 145  $\mu$ mol g<sup>-1</sup> h<sup>-1</sup> (Fig. 2a). This decreased catalytic reactivity was due to the combined result of lower CO<sub>2</sub> concentration in the exhaust gas as well as the presence of NO<sub>x</sub> or SO<sub>x</sub> in the exhaust gas, which can also be adsorbed on the active catalytic sites.<sup>71</sup> In addition, the kinetic studies demonstrated the high stability of this photocatalyst in the presence of impurities such as H<sub>2</sub>, CO, CH<sub>4</sub>, O<sub>2</sub>, N<sub>2</sub>, and others, which are typically present in car exhaust gas (Fig. 2b). Furthermore, to estimate the reusability of this photocatalyst, 10 mg of the material was successfully used (under the same reaction conditions) and recycled for up to three cycles (Fig. S10†).

To achieve a deep characterization of the synthesized materials, we investigated the morphology and the structure of 1Fe@f-gC<sub>3</sub>N<sub>4</sub> by High-Angle Annular Dark Field Scanning Transmission Electron Microscopy (HAADF-STEM). As shown in Fig. 3a, nanoparticles (NPs) were non-uniformly distributed over f-gC<sub>3</sub>N<sub>4</sub>, and EDX-mapping demonstrated that the NPs contained Fe (Fig. 3b-e). Atomic resolution HAADF-STEM images were acquired at lower (Fig. 3f) and higher (Fig. 3g) magnification. Fast Fourier transform (FFT) of the NPs marked by the red dashed rectangle in Fig. 3g provided lattice spacings of 2.690 Å, 2.580 Å, and 2.446 Å (Fig. 3h), which corresponded to FeO(OH) (130), (0-21), and (111), respectively. These results demonstrated that the supported NPs correspond to FeO(OH) species, and further investigating the edge of the f-gC3N4 support, bright dots marked by yellow dashed circles were observed, corresponding to small clusters containing Fe (Fig. 3i).

The crystal structure of prepared  $gC_3N_4$ , f- $gC_3N_4$ , and 1Fe@f- $gC_3N_4$  were further investigated via XRD. Two distinct diffraction peaks were observed at 12.9° and 27.3°, corresponding to the (100) and (002) crystal planes of g- $C_3N_4$ , related to the inplane repeating s-triazine structural moieties and interlayer stacking of the conjugated aromatic ring (Fig. S11†). T2-74 While for f- $gC_3N_4$ , the peak position and the peak intensity were almost similar with  $gC_3N_4$ , in the presence of Fe, the intensity of diffraction peak (002 in case of 1Fe@f- $gC_3N_4$ ) was slightly decreased as well as a marginal shift to higher diffraction angle

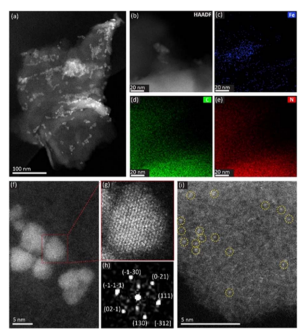


Fig. 3 (a) HAADF-STEM image showing the morphology of  $1Fe@f-gC_3N_4$ . (b–e) HAADF-STEM image (b) and EDX mapping results (c–e) showing the support  $f-gC_3N_4$  and the Fe nanoparticles. (f) Atomic resolution HAADF-STEM image showing the  $Fe_2O_3$  nanoparticles. (g) Image of magnified area in f marked by red dashed rectangles. (h) Fast Fourier transform (FFT) pattern of the FeO(OH) nanoparticles in  $1Fe@f-gC_3N_4$  (g). The direction is FeO(OH) [–312]. (i) Atomic resolution image showing the Fe clusters.

was observed.  $^{73,75}$  However, no obvious diffraction peak of  $Fe_2O_3$  or FeO(OH) phase were detected in the pattern of  $1Fe@f-gC_3N_4$ , due to relatively low amount of the Fe species.  $^{66,76,77}$  Additionally, the weakening of the intensity of (002) peak suggested a decrease in crystallinity, with a consequent increase in the number of defects. These could trap a large number of carriers, with a resulting increase in charge separation and an improvement of the photocatalytic performance.  $^{75}$ 

To verify the atomic environment of Fe in 1Fe@f-gC<sub>3</sub>N<sub>4</sub>, synchrotron X-ray absorption near-edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) at Fe K-edge were performed and compared to the X-ray absorption spectra of the Fe<sub>2</sub>O<sub>3</sub> and FeOOH reference materials. From the position of the rising edge in XANES spectra (Fig. 4a), it can be concluded that Fe in 1Fe@f-gC<sub>3</sub>N<sub>4</sub> was predominantly in a +3 oxidation state, as confirmed by the close resemblance with Fe<sub>2</sub>O<sub>3</sub> and FeOOH pre-edge peaks. EXAFS and Fourier transformed (FT)  $k^3$ weighted EXAFS curves were further extracted to probe the atomic iron-based local structures (Fig. 4b and c). When compared with the reference systems, 1Fe@f-gC3N4 nearly matched the peaks of FeOOH, which was especially noticeable for the shell scattering peaks of the longer-range order (2-3.5 Å), indicating Fe-Fe bonding. Concurrently, only FeOOH species could not fully describe XANES region and first shell, and a low Fe<sub>2</sub>O<sub>3</sub> fraction below 5% cannot be excluded.

Solid-state MAS NMR spectra of all NMR-active nuclei present in the material (<sup>1</sup>H, <sup>13</sup>C, <sup>15</sup>N, and <sup>19</sup>F) were also collected

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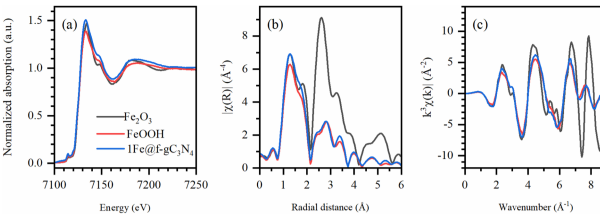


Fig. 4 (a) Normalized XANES spectra, (b) FT-EXAFS spectra (phase uncorrected) and (c)  $k^3$ -weighted EXAFS spectra at Fe K-edge for the 1Fe@fgC<sub>3</sub>N<sub>4</sub> and reference samples (Fe<sub>2</sub>O<sub>3</sub> and FeOOH).

Radial distance (Å)

to probe the local structure and to examine potential structural changes upon doping of the f-gC<sub>3</sub>N<sub>4</sub> with Fe<sup>n+</sup> species (Fig. S12†). The <sup>1</sup>H MAS, <sup>13</sup>C CPMAS, and <sup>15</sup>N CPMAS spectra were almost identical to NMR data we collected from the related polymeric carbon nitride catalysts and reported recently. 49,78 Therefore, we concluded that the overall structure and polymerization degree in these materials were not affected to any significant extent by doping with Fe<sup>n+</sup> ions. However, in the <sup>19</sup>F MAS NMR spectrum, the <sup>19</sup>F NMR shift of -120 ppm was distinct from that of -105 ppm observed by us for the undoped f-gC<sub>3</sub>N<sub>4</sub>, attributed to the presence of =CF<sub>2</sub> groups (Fig. S12b†).49,79,80 Moreover, an additional weak signal at -202 ppm was detected, which had not been observed for the undoped material. The additional <sup>19</sup>F MAS spectrum recorded using a short 0.2 s relaxation delay was collected to inspect if the appearance of the signal at -202 ppm could be related to the introduction of Fe<sup>n+</sup> ions in the material. Upon using a short relaxation delay of 0.2 s, the intensity of the signal at -202 ppm was almost unaffected, whereas the signal at -120 ppm was significantly saturated. This could be attributed to the paramagnetic relaxation enhancement of the 19F nuclei in close contact with paramagnetic Fe<sup>n+</sup> ions. The observed <sup>19</sup>F NMR shifts of -120 and -202 ppm were most probably affected by the induced paramagnetic NMR shift interactions.81

Energy (eV)

The XPS spectra were recorded in the Fe 2p, C 1s, and N 1s regions for both fresh and used samples of 1Fe@f-gC3N4 (Fig. S13†). The dominant component of the pristine 1Fe@fgC<sub>3</sub>N<sub>4</sub> was the g-C<sub>3</sub>N<sub>4</sub> phase, as confirmed by the C 1s and N 1s spectra. In the C 1s region, photoelectron emission, typical of sp<sup>2</sup>-bonded C atoms in N-C=N, was observed at 288.1 eV, while in the N 1s region three peaks at 398.6 eV (sp<sup>2</sup>-hybridized N atoms in C-N=C species), 400.1 eV (Odging N atoms N-(C)3 species) and 401.2 eV (N atoms in amino groups) were found. 82,83 Fe<sup>n+</sup> species also appeared on the surface of f-gC<sub>3</sub>N<sub>4</sub>. Due to the low content of this component, it was difficult to determine the chemical state of Fe unequivocally. Nevertheless, the Fe 2p<sub>3/2</sub> peak position at 710.6 eV with an apparent multiplet splitting and a low satellite at ca. 718 eV indicated the presence of high-spin Fe<sup>3+</sup> species. Another evidence confirming this

supposition was the orbital splitting of Fe 2p<sub>3/2</sub> and Fe 2p<sub>1/2</sub> of 13.8 eV, which was similar to that reported previously in the literature for Fe<sup>3+</sup>.84 The elemental composition of the photocatalyst surface did not change during the process, and the used sample exhibited the same peaks in the individual XPS spectra. However, their intensity decreased due to the deposition of an additional component on the surface. Its nature was revealed by the XPS C 1s spectrum, where two new peaks were distinguished at binding energies of 284.8 eV and 285.9 eV, respectively. The former was typical for C-C/C=C, while the latter was for the oxidized C forms, most likely C-O or C=O.85

To obtain better insights into the mechanism of the reaction, the optical properties of the photocatalysts were characterized. Through UV/vis spectroscopy (Fig. S4a†), an absorption edge at 465 nm was observed in the case of gC<sub>3</sub>N<sub>4</sub>, whereas f-gC<sub>3</sub>N<sub>4</sub>, 0.5Fe@gC<sub>3</sub>N<sub>4</sub> and 0.5Fe@f-gC<sub>3</sub>N<sub>4</sub> exhibited a redshift of 30 nm, 34 nm and 34 nm, respectively (corresponding to 0.16 eV, 0.18 ev and 0.18 ev respectively). However, different Fe-loading had a negligible effect on the optical spectrum. The broad and intense absorption peak of the Fe-loaded f-gC<sub>3</sub>N<sub>4</sub> catalyst in the visible region inferred that the catalyst could absorb more photons, which was a consequent enhancement of the photocatalytic activity.86 The difference in the absorption band between gC<sub>3</sub>N<sub>4</sub> and Fe-loaded f-gC<sub>3</sub>N<sub>4</sub> could be due to the electrostatic interaction between Fe<sup>+n</sup> and f-gC<sub>3</sub>N<sub>4</sub> which promoted the electron delocalization throughout the heptazine framework by Fe<sup>+n</sup>.87 This was also evident from EPR spectroscopy, which showed a broadening of the 1Fe@f-gC3N4 signal (compared to f-gC<sub>3</sub>N<sub>4</sub> one), due to the mutual interaction between the paramagnetic species (Fig. S14†).

The photocatalysts' steady-state photoluminescence (PL) spectra (Fig. S17a†) revealed emission maxima of gC<sub>3</sub>N<sub>4</sub> at 471 nm and of Fe-gC<sub>3</sub>N<sub>4</sub> at 466 nm. The emission spectra of fgC<sub>3</sub>N<sub>4</sub> and Fe-loaded f-gC<sub>3</sub>N<sub>4</sub> were broad and centered around 513 nm. To investigate the charge carrier generation and dynamics, we measured the PL decays of the photocatalysts at their respective emission wavelength (Fig. S17b and c†). The PL decay profiles of all samples were recorded by time-correlated photon counting (TCSPC) spectroscopy and fitted using a tri**Edge Article Chemical Science** 

exponential decay equation. The average PL lifetimes and the fitting parameters are given in Table S3†. According to the lifetime fitting parameters, f-gC<sub>3</sub>N<sub>4</sub> and Fe-loaded f-gC<sub>3</sub>N<sub>4</sub> had longer  $\tau_1$  and  $\tau_2$  values compared to the gC<sub>3</sub>N<sub>4</sub> and Fe-gC<sub>3</sub>N<sub>4</sub> catalysts. Additionally, upon increasing Fe-loading in f-gC<sub>3</sub>N<sub>4</sub>, the carrier lifetimes exhibited an increasing trend of all  $\tau_1$ ,  $\tau_2$ , and  $\tau_3$  values. Although there is a slight difference in the average lifetime of the photocatalysts, the iron loading in fgC<sub>3</sub>N<sub>4</sub> increases the average lifetime, resulting in enhanced photocatalytic activity. This result supported the aboveexplained enhanced photocatalytic reactivity upon increasing the iron wt% due to a more efficient transfer of photogenerated electrons from f-gC<sub>3</sub>N<sub>4</sub> to Fe<sup>+n</sup>. Following the increase of the charge carrier lifetimes, the generated electrons were available for a longer time for the photocatalysis reaction to efficiently take place. Specifically, in 1Fe@f-gC<sub>3</sub>N<sub>4</sub>, the  $\tau_3$  value, reflecting the lifetime of the free charge carriers that can diffuse over longer distances, has significantly increased.

In addition, to investigate the separation efficiency of photogenerated electron-hole pairs, we recorded the photocatalyst's transient photocurrent responses under the irradiation of a Kessil lamp ( $\lambda = 427$  nm). Among all of them, 1Fe@f-gC<sub>3</sub>N<sub>4</sub> exhibited the quickest and highest photocurrent which remained stable up to 5 cycles which indicated a more efficient charge separation and faster electron transfer rate to trigger a superior photocatalytic reactivity of the catalyst (Fig.-S18b†). Furthermore, in situ EPR spectroscopy showed that the relative number of the photoexcited electrons of f-gC<sub>3</sub>N<sub>4</sub> was higher than 1Fe@f-gC<sub>3</sub>N<sub>4</sub> due to the facile electron transfer to Fe (Fig. S14†). Additionally, Electrochemical Impedance Spectroscopy (EIS) of gC<sub>3</sub>N<sub>4</sub>, f-gC<sub>3</sub>N<sub>4</sub> and all the Fe-loaded gC<sub>3</sub>N<sub>4</sub> and f-gC<sub>3</sub>N<sub>4</sub> exhibited decreasing charge transfer resistance, demonstrating the presence of covalent linking in 1Fe@f-gC<sub>3</sub>N<sub>4</sub> which significantly enhanced the conductivity (Fig. S18a†). The CO<sub>2</sub> reduction was confirmed by a spin-trapping experiment with DMPO (Fig. S16†). The holes in the valence band were quenched by the sacrificial reductant (TEA) and decreased the electron-hole recombination, as it is evident from the in situ EPR investigations (Fig. S15†). In the valence band, TEA reacted with holes and subsequently formed the α-amino radical and protons.88

Further to gather detailed information about the reaction mechanism, computational calculations were done (see in the ESI†) and after  $\approx$  50 ps of RMD, the sampled structures showed that the adsorption tendency of CO2 on the metal centers was mainly due to the coordination of one of its oxygens (Fe-O distance about 2.1 Å),89 and no reaction mechanisms were observed in those conditions. Indeed, the conversion started from an activated adsorption configuration where the carbon atom was connected to the metal center, and the CO2 molecule adopted a bent arrangement (CO2. -).90-92 Experimentally, this activation was obtained by light irradiation ( $\lambda = 427$  nm), which induced a charge transfer from the catalyst to the molecule, CO<sub>2</sub> chemisorption with the elongation of the C-O bond, bending of O-C-O angle, and finally, dissociation of CO<sub>2</sub> on the catalyst surface into CO and O species.93,94

We could mimic this process by including, in the RMD simulations, an electric field in the plane of the melem units (x direction,  $-0.01 \text{ V Å}^{-1}$ ). The effect of the external electric field is a perturbation of the atomic charges of the system and, thus, a distortion of the molecular charge distribution. The evolution of the dipole moments imitated a charge transfer from the metal center to the adsorbed CO<sub>2</sub>. This is apparent in the atomic charge distribution plots of Fig. S21,† where it is evident the charge transfer from Fe to CO<sub>2</sub> during the first 7.5 ps of the simulation (Fig. S21† - bottom) and the change of nature of the C atom when CO<sub>2</sub> reduces to CO (Fig. S21† - middle plot). The adsorbed CO2 in a bent geometry was negatively charged. Indeed, from the beginning of the polarized dynamics, CO<sub>2</sub> changed from an extended to a bent conformation and remained adsorbed on the Fe atom through its carbon (Fig. S20†). The stabilized complex received a proton from the solution and formed the adsorbed \*COOH species. This species was short-lived, and the OH was quickly released in solution and then protonated. In contrast, CO remained stably adsorbed on the metal center (Fig. S22†). Besides reproducing possible hydrogen exchanges between the solvent and the region around the COO-Fe complex, the mechanisms produced a water molecule that freely migrated in the solution, whereas CO remained connected to the metal.

To refine this picture, we extracted the MD snapshots describing the primary steps of the reaction mechanism, sizereduced them to three melem units with a chelated Fe atom and an adsorbed CO<sub>2</sub> molecule (high-coordination site) or two melem units with a chelated Fe-atom, the f-substituent, and an adsorbed CO2 molecule (low-coordination site), and carried out density functional theory (DFT) calculations with Gaussian 16.95 These were used to estimate minimum energy structures, charge analyses, possible reaction paths, and relative energy barriers (Fig. 5, 6 and S23†). We optimized the geometries in ACN through the integral equation formalism variant (IEFPCM) of the Polarizable Continuum Model (PCM), using the B3LYP-D3(BJ) functional with the Grimme D3 correction (Becke-Johnson parameters97), to account for the van der Waals interactions, and the 6-31(d,p) basis set for all the elements except Fe, which was described with the def2-TZVP basis set. 98 Charge analysis was performed using NBO (full Natural Bond Orbital).99,100

Both high- and low-coordination sites provide a similar qualitative picture of the reaction mechanism, in agreement with the results of RMD simulations. In the absence of external stimuli, CO<sub>2</sub> molecule interacted weakly via one of its oxygens with the catalyst metal center (Fe-O equilibrium distance is 2.04 Å, in fair agreement with the results of RMD), which acted as a Lewis acceptor (see state 1 in Fig. 5), as confirmed by the small positive charge carried by CO<sub>2</sub> (about +0.2e) and the charge decrease on Fe after adsorption (about 1.1e, to be compared with 1.3e of a naked Fe atom - Fig. S23†). The  $CO_2^*$  activation happened after the addition of a H<sup>+</sup>/e<sup>-</sup> couple promoted by UV irradiation: a similar bent adsorption configuration obtained by RMD (and shown in state 2 of Fig. S23†) was stabilized by a flow of negative charge to both the  $CO_2$  molecule (charge of  $\approx -0.4e$ ) and the Fe center (charge decrease from  $\approx +1.0e$  to  $\approx +0.7e$ ). The

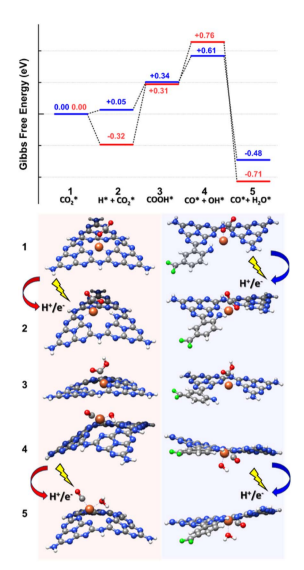


Fig. 5 Reaction mechanism showing  $CO_2$  reduction to CO involving Fe atom in a high-coordination (red path) and a low-coordination (blue path) sites. State 1 is chosen as a reference for energy estimation, whereas the free energies of the other states (from 2 to 5) are estimated at the DFT level by employing the Norskov model. Golor code: C gray, N blue, F green, Fe orange, O red, H white.

proton (as in RMDs) was adsorbed on a nitrogen atom at the edge of a neighboring melem unit.  $CO_2$  negative charging is also evident in the PDOS shown in Fig. S24.†

After proton migration to  $\mathrm{CO}_2^*$  we obtained the COOH\* configuration shown in state 3 (Fig. 5). This positive charge flow allows negative charge back-migration from Fe to the adsorbate (with COOH almost neutral), as shown in Fig. S23.† The elongation and weakening of the bond between the carbon and the hydroxyl group (induced by the catalyst-adsorbate charge transfer, populating the p\* LUMO of  $\mathrm{CO}_2$  (ref. 88)) can be exasperated until the breaking of the C–O bond realized the configuration shown in state 4 (Fig. 5), which presents CO and OH separately adsorbed on the Fe atom. The positive charge carried by CO was compensated by a reduced charge on Fe (relative to state 2), and the negative charge carried by OH\*

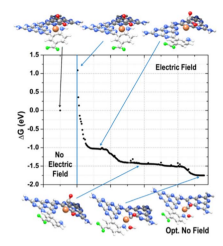


Fig. 6 Reaction path of the adsorbed  $\mathrm{CO}_2$  reduction to CO. The DFT minimum energy structures (initial and final points – top left and bottom right ball and stick models) are shown together with the intermediate species indicated by the blue arrows. Color code: C gray, N blue, F green, Fe orange, O red, H white.

(about -0.2e). States 3 and 4 are at higher energy relative to state 2 ( $\approx$ 1.08 eV for the high-coordination site and 0.56 eV for the low-coordination). The reduced energy difference characterizing the low-coordination site suggests that this can promote  $\mathrm{CO}_2$  reduction easier than the high-coordination site and will be further investigated in the following. The UV-vis spectra simulated for state 2 of the low-coordination site, calculated at the TD-DFT level (Fig. S25†), show a dominant peak at about 465 nm, corresponding to a charge transfer from the metal ion to the adsorbed  $\mathrm{CO}_2^*$  molecule, aiding the C–O bond breaking in overcoming the observed energy difference between states 2 and 4. This agrees with the experimental choice of a blue LED ( $\lambda$  = 427 nm) as the excitation source.

Further addition of a H<sup>+</sup>/e<sup>-</sup> couple (promoted by UV excitation) led to the configuration of state 5 (Fig. 5), which corresponded to adsorbed CO and H2O on the Fe catalyst. H2O carried a slight positive charge (about +0.2e), and CO kept its positive charge of about +0.4e. The relatively strong dipole between Fe and CO is a clear signal of a strong CO anchoring to the catalytic site, which can be disrupted if the charge flows from Fe to CO. The Gibbs free energy of this last state is lower than that of any other state investigated, confirming the thermodynamic tendency of this system to promote CO<sub>2</sub> reduction. This sequence of events is the same as observed in RMD. We resorted to the electric field option already used for the RMD to recreate a possible dynamics mechanism at the DFT level instead of performing computationally expensive excited-state quantum chemistry simulations. Starting from the optimized complex of the low-coordination corresponding to state 2 (Fig. 5), where the CO<sub>2</sub> molecule was adsorbed with the carbon atom on the metal ion, we emulated the first few steps of the reduction mechanism by applying an external electric field in the -x direction with a magnitude of 0.008 au.

From the examination of the free energy difference plot, it is evident that a low activation energy of  $\approx 1.1$  eV was necessary to

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start the process; this activation energy is reasonable when compared to the energy difference of  $\approx 0.6$  eV between states 2 and 4, estimated at the static level (Fig. 6). Then, the mechanism proceeded barrierlessly, passing through two intermediate metastable geometries (plateau regions). The first one is characterized by the appearance of the adsorbed COOH\* species (state 3 in Fig. 5), whereas in the second one, the OH detached from C and became connected to the metal center (at a C-Fe-OH angle of about 103°, state 4 in Fig. 5), forming a bond with Fe (with an average Fe-O length of approximately 1.8 Å) and a hydrogen bond with the nitrogen atom of the nearby triazine ring (with an average OH-N distance of roughly 1.8 Å and a donor-H-acceptor angle of about 140°). In the final stable configuration, the hydrogen bond was lost, and the OH group moved to a farther N-O separation of about 3 Å but remained connected to the metal center. Interestingly, when simulated under the effect of the electric field, the energies of states 3 and 4 become lower than that of state 2, indicating a stabilizing effect played by the external perturbation on configurations which, when analyzed in their ground state, resulted instead higher in energy.

#### Conclusions

In conclusion, we have successfully demonstrated that coupling amorphous FeOOH with f-gC<sub>3</sub>N<sub>4</sub> exhibited an efficient visible-light-mediated CO<sub>2</sub> conversion into CO. The presence of FeOOH in functionalized gC<sub>3</sub>N<sub>4</sub> modulated the electronic interaction between Fe species and semiconductor, making an efficient photocatalyst for CO<sub>2</sub> reduction. This FeOOH/f-gC<sub>3</sub>N<sub>4</sub> composite heterogeneous photocatalyst has shown the highest CO evolution rate (so far reported among Fe-based heterogeneous photocatalysts) of 304  $\mu$ mol g<sup>-1</sup> h<sup>-1</sup> with an excellent selectivity of >99%. This earth-abundant and low-cost photocatalyst also exhibited excellent reactivity and stability for reducing car exhaust gas, which clearly depicted the strong application potential of this chemistry. We strongly believe that our photocatalytic system will open a new strategy in the field of photocatalytic CO<sub>2</sub> reduction.

#### Author contributions

T. G., P. R. and S. D. conceptualized the project. S. D. supervised the project. T. G and P. R. synthesized the catalysts, conducted the catalytic experiments and the related data processing, and performed materials characterization and analysis with the help of P. F., A. J., A. R., P. K., L. C. and E. D. Furthermore, A. J. collected solid-state NMR spectra and J. R. performed EPR investigations. M. T. and S. B. conducted high-resolution, high angle annular dark-field transmission electron microscope (HAADF-STEM) spectroscopy. G. B. and S. M. performed the theoretical studies. J. S.-A., A. S. and L. S. performed X-ray absorption near edge structure (XANES) and extended X-ray absorption fine structure (EXAFS) measurement and analysis. A. C and E. D performed the steady state spectroscopy. A. R and P. K performed the X-ray photoelectron spectroscopy (XPS). The

manuscript was written through the contributions of all authors.

#### Conflicts of interest

There are no conflicts to declare.

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