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Determining the structure and properties of CO₂ reduction photocatalysts: single atom cobalt atoms supported on various carbon nitrides†

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Carbon nitride materials paired with Co atoms have been shown to be effective for CO₂ photoreduction. However, the structures of Co/carbon nitrides are uncertain, especially when the degree of polymerization of the carbon nitrides is unknown. Literature has focused predominantly on the fully condensed carbon nitride, g-C₃N₄, despite evidence of other carbon nitrides being present in carbon nitride catalysts. We therefore used density functional theory (DFT) to model Co binding with molecular, partially condensed polymeric, and fully condensed polymeric carbon nitrides. We found strong binding of Co to the carbon nitrides, and larger coordination tended to lead to stronger binding of Co, as well as more cationic Co atoms. Co had a significant effect on photocatalytic processes. Co lowered band gaps significantly, enabling greater photoexcitation yields. Activation of CO₂ into a bent, anionic state is an important initial step during CO₂ reduction. Our calculations show that all Co/carbon nitride systems, except two-layer carbon nitrides, activated CO₂. For bent CO₂, strong interactions occurred between Co/carbon nitride systems, as evidenced by density of states plots and calculated interaction energies. On the other hand, weak interactions occurred with adsorbed linear CO₂. Activation energies of CO₂ spanned a wide range of values (−2.33 to −0.38 eV), and intermediate values over partially condensed carbon nitrides may be more likely to enable CO₂ reduction. Our work provides insights and understanding of Co/carbon nitride catalysts, and motivates study of carbon nitrides beyond g-C₃N₄ as photocatalysts.

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1 Introduction

Utilizing sunlight with efficient, low-cost photocatalysts is a vital step towards a clean, sustainable future. Carbon nitride materials have attracted much interest as photocatalysts,^{1–9} in part because these materials consist of cheap, abundant elements. They also have moderate band gaps for photoexcitation. These catalysts have been studied for a number of reactions, including water splitting,^{10–14} water purification,^{15–17} and CO₂ reduction.^{18–23} Reducing atmospheric CO₂ is an essential goal for mitigating climate change.

Carbon nitrides can have various compositions and structures.^{24–28} One of the most studied carbon nitrides is graphitic C₃N₄, or simply g-C₃N₄, which has a two-dimensional graphene-like structure with interconnected heptazine units. g-C₃N₄ may be considered a fully condensed two-dimensional material whereby melem molecules (C₆N₁₀H₆) combine together to release NH₃.^{28–30} Both experimental and computational work have shown that various polymeric carbon nitrides

form due to incomplete condensation under different reaction conditions.^{13,28,31–36} Density functional theory (DFT) work suggests that g-C₃N₄ may be hard to synthesize under typical synthesis conditions (*e.g.*, atmospheric pressure), and that partially condensed polymeric carbon nitride structures (*e.g.*, melon – [C₆N₉H₃]_n) may be more prevalent than many researchers have considered.^{31–33,35,37} Furthermore, most experimental carbon nitrides contain a significant amount of hydrogen^{25,28,35} and also deviate from the ideal g-C₃N₄ C : N ratio of 0.75,^{24,34} again indicating the likely presence of partially condensed polymers in many studies. Indeed recent DFT studies have attempted to more fully understand polymeric carbon nitrides such as melon.^{31,38–40} Other DFT work³³ suggests that polymeric carbon nitrides may have many different structural domains (both partially and fully condensed) rather than a single structure. Thus, there is a need to better understand and characterize polymeric carbon nitrides beyond just two-dimensional g-C₃N₄, particularly partially condensed carbon nitrides. In this work we concentrate on characterizing polymeric carbon nitride materials as photocatalysts for CO₂ reduction, especially when paired with single metal atoms.

One promising way to increase photocatalytic activity of carbon nitrides is to pair them with single metal atoms.^{41–48} Single atom catalysts (SACs) have attracted much interest

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because they can have high catalytic activity per atom, and can also have novel chemical or catalytic properties compared to bulk materials. The carbon nitride acts as a support and photoabsorber for these SACs. Synergy between the metal atoms and carbon nitride can decrease band gaps/increase photoabsorption,^{49–55} and also increase charge separation/lower charge recombination (necessary for long-lived charge carriers).^{42,46,55–59} Such carbon nitride/single metal atom catalysts may be active for reactions such as the H₂ evolution reaction^{52,58,60,61} or CO₂ reduction.^{62–66} Of particular interest is Co, which has demonstrated activity for CO₂ photoreduction.^{67–74} Using Co/carbon nitride catalysts for CO₂ photoreduction is a promising way to deal with the CO₂ problem. Co/carbon nitride catalysts have demonstrated better adsorption of CO₂, higher light utilization, and higher production of CO compared to pure carbon nitrides.⁷²

Identifying the geometry of Co/carbon nitrides is an important step towards connecting their structure to catalytic activity, as well as a precursor for simulating these materials using atomistic modeling methods. Without details on their structure, detailed reaction mechanisms and structure–activity relationships are unclear. Indeed, there is ambiguity about the structure of Co/carbon nitrides. For example, the Co–N coordination number of Co/carbon nitride catalysts has been reported to be in the range of two to four depending on the material.^{61,68,71–76} We have shown through theory and experiment a potential four-coordinated structure.⁷⁴ In contrast, Co may be four- or six-coordinated in bulk cobalt oxides. Other work involving doped carbon nitrides reported four-coordinated Co–P structures.⁷⁷ Besides details on the geometry of Co atoms, identifying the oxidation state, effect on photoabsorption, and potential reaction sites are important for clarifying structure–activity relationships. The modeling literature has predominantly focused on how Co SACs bind to g-C₃N₄.^{78–81} A few DFT papers have been published on Co SACs supported by melon, but these papers assumed a dehydrogenated form of melon,^{82,83} while another paper modeled SACs to one-layer melon.⁸⁴ Accordingly, it is critical to understand the nature of Co interacting with a full range of carbon nitrides, including molecular, partially condensed polymers, and fully condensed polymers, as well as multi-layer carbon nitrides, as all these types of carbon nitrides may be present in real-world catalysts.

In this work, we used DFT to model Co/carbon nitride catalysts in order to clarify their structures and determine potential photocatalytic activity, especially for CO₂ photoreduction. We modeled molecular carbon nitrides, partially condensed carbon nitrides, and fully condensed two-dimensional carbon nitrides. We assessed how different carbon nitride supports interact with Co atoms, leading to different complexes and Co oxidation states. We also elucidated how Co atoms may affect band gaps of these materials, which is important for photocatalysis. Finally we modeled the activation of CO₂ over these materials, a necessary first step towards CO₂ photoreduction. Our work provides fundamental details on these catalysts, and ties

together how different carbon nitride supports affect SAC stability, structure, and photocatalytic activity.

2 Methodology

2.1 Simulation parameters

We used the Vienna *Ab initio* Simulation Package (VASP)^{85–88} with the Perdew Burke Ernzerhof exchange-correlation functional⁸⁹ for all the DFT simulations. Valence electrons were represented by a plane wave basis set, while core electrons were represented by projector augmented wave (PAW) potentials.^{90,91} The number of valence electrons for each atom were: Co (9), C (4), N (5), H (1), and O (6). A cutoff energy of 450 eV was used for the plane waves. Criterion for convergence of electronic and ionic relaxations were 1×10^{-5} eV and $0.02 \text{ eV } \text{Å}^{-1}$. The Grimme D3 correction⁹² with Becke–Johnson damping⁹³ was used to incorporate van der Waals interactions. We also used Gaussian smearing with a σ value of 0.1 eV. Charges of atoms were determined using Bader charge analysis.^{94–96} We used the HSE06 hybrid functional,^{97–99} combined with Gaussian smearing and a σ value of 0.01 eV, for calculating higher resolution band gaps and the density of states. The VASPKIT code was used for post-processing of our simulations.¹⁰⁰ Images of structures were generated using VESTA.¹⁰¹

Binding energies ($\Delta E_{\text{binding}}$) of Co atoms were calculated by eqn (1):

$$\Delta E_{\text{binding}} = E_{\text{Co/Carbon Nitride}} - E_{\text{Carbon Nitride}} - E_{\text{Co}} \quad (1)$$

Here $E_{\text{Co/Carbon Nitride}}$ is the energy of the Co/carbon nitride complex, $E_{\text{Carbon Nitride}}$ is the energy of the carbon nitride material, and E_{Co} is the energy of a lone Co atom. Adsorption energies (ΔE_{ads}) of CO₂ were calculated by eqn (2):

$$\Delta E_{\text{ads}} = E_{\text{CO}_2/\text{Co/Carbon Nitride}} - E_{\text{Co/Carbon Nitride}} - E_{\text{CO}_2} \quad (2)$$

Here $E_{\text{CO}_2/\text{Co/Carbon Nitride}}$ is the energy of the CO₂/Co/carbon nitride complex and E_{CO_2} is the energy of a CO₂ molecule.

2.2 Structural models

In this work we modeled several carbon nitrides, including polymeric building blocks (melem and melem dimer), partially condensed polymers (melon), and fully condensed carbon nitrides (g-C₃N₄). Fig. 1 shows the carbon nitrides we modeled. These carbon nitrides have been previously modeled.^{31,32,40} Interestingly, recent simulations suggest that experimentally relevant carbon nitrides consist of both partially and fully condensed polymer structures.³³ Melem is a building block of polymeric carbon nitrides.^{24,25,102} Melem dimers may also form during the polymerization process. Melem and melem dimers were modeled in large supercells of size $30 \text{ Å} \times 30 \text{ Å} \times 30 \text{ Å}$. We used a single *k*-point for these calculations. Condensation of melem can form melon.^{24,33,102} Our melon model was based on Fina *et al.*³⁷ and is further discussed in the ESI Section 1.† As Fig. 1 shows, the stable melon structure consisted of melon



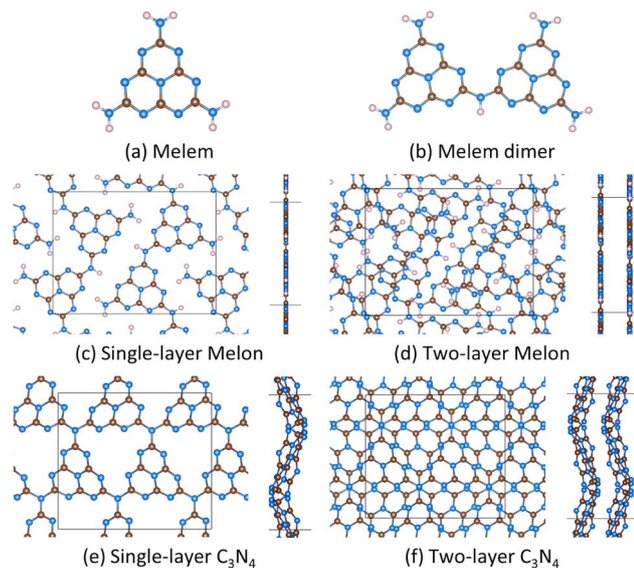


Fig. 1 Carbon nitrides molecules, melem (a) and melem dimer (b), modeled in this work. Periodic carbon nitride polymers, melon (c and d) and graphitic C_3N_4 (e and f), modeled in this work. Also shown are the simulation cells for the periodic structures. Brown spheres represent carbon atoms, blue spheres represent nitrogen atoms, and pink spheres represent hydrogen atoms.

strands arranged together within the simulation cell. The single-layer melon structure had lattice lengths of 12.8 Å and 16.8 Å (based on lattice optimization) with a vacuum spacing of ~ 30 Å, and included 72 atoms. Our two-layer melon structure was also based on Fina *et al.*'s melon structure.^{37,40} This simulation cell had lattice vectors of 12.8 Å and 16.7 Å after lattice optimization. The z lattice vector was set to ~ 30 Å, resulting in a vacuum spacing of ~ 27 Å.

Single-layer $g-C_3N_4$ (similar to Melissen *et al.*³¹) was modeled using a (2×2) orthorhombic supercell containing 56 atoms and with optimized lattice lengths of 11.8 Å and 13.1 Å. The vacuum space was ~ 30 Å. Corrugated $g-C_3N_4$ is more stable than planar $g-C_3N_4$, as found in literature³¹ and our own calculations, so we modeled all $g-C_3N_4$ as corrugated. We also modeled two-layer $g-C_3N_4$, as layers of $g-C_3N_4$ may be attracted by van der Waal forces. Two-layer $g-C_3N_4$ (taken from Botari *et al.*³²) was modeled with an orthorhombic cell containing 112 atoms with lattice parameters being 11.9 Å and 13.3 Å. The vacuum space was ~ 26 Å. Test calculations (see Table S1†) indicated that a $(2 \times 2 \times 1)$ k -point mesh was sufficient for all the periodic polymer calculations (*i.e.*, melon and $g-C_3N_4$), and we used such a k -point mesh in our work for all the periodic carbon nitrides. Analysis of our carbon nitride models (see the ESI Section 2†) indicates agreement with literature.

A challenge is quickly identifying whether Co–C or Co–N bonds occur in a structure, or the coordination number of Co. We used a simple procedure of designating a bond if the Co–X ($X = C, N$) bond distance was less than some threshold cutoff value, similar to previous work.^{103,104} Co–X bond distances in Co/carbon–nitrogen materials (such $g-C_3N_4$ and nitrogen doped graphene) were found to range from 2.02 to 2.44 Å (Co–C) and

2.04 to 2.37 Å (Co–N) (from both experimental^{61,75,105–108} and computational work^{61,75,105}). Therefore, we set the cutoff value for determining Co–C and Co–N bonds to 2.5 Å.

3 Results and discussion

3.1 Binding of Co to carbon nitride materials

3.1.1 Binding of Co to molecular carbon nitrides. We considered how single Co atoms bind to the different carbon nitrides, first examining Co bound to carbon nitride molecules. In order to identify the most stable structures, we modeled several possible Co/molecular complexes. For example, we modeled Co binding to two adjacent N atoms within a single carbon nitride molecule, as well as combinations in which N atoms from two carbon nitride molecules interacted with the Co atom. Furthermore, the Co atom was placed in different orientations, such as the same plane with the carbon nitrides or sandwiched between two molecules. We present those structures which are most stable after geometry optimization, or lowest in energy, in Fig. 2. Table 1 also summarizes details on these structures. We note that several structures had quite different geometries, but similar binding energies for Co. Thus, we present several different structures for each molecule, and label them based on the number of carbon nitride molecules and their configuration. A simplified representation of each structure showing only the local coordination around the Co atom is provided in Fig. S3.† For example, Co/melem-1 refers to the first structure with Co binding with one melem, while Co/melem-2 refers to the second structure with Co binding with one melem. Co/2-melem-1 refers to the first configuration of Co

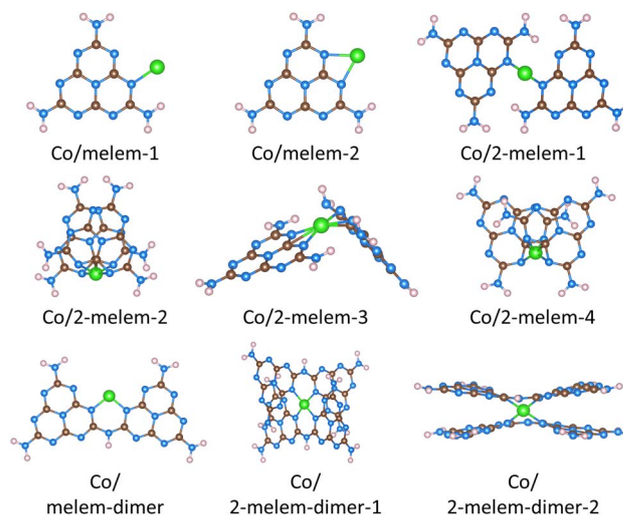


Fig. 2 Stable structures where Co was bound to carbon nitride molecules. Brown spheres represent carbon atoms, blue spheres represent nitrogen atoms, pink spheres represent hydrogen atoms, and green spheres represent cobalt atoms. The notation system is described in the main text. When two molecules are present, these are designated by the prefix '2-'. Each unique configuration is identified by the final number (*e.g.*, '-1', or '-2'). A simplified representation of each structure showing only the local coordination around the Co atom is provided in Fig. S3.†



Table 1 Results for Co bound to molecular carbon nitrides. Shown are calculated Co Bader charges, Co binding energies, Co coordination numbers (Co–N coordination number, followed by Co–C coordination number), and distances between Co and closest atoms (typically N). In some cases Co was bound to C atoms, as indicated in the table. Structures in the table correspond to the same structures shown in Fig. 2. Simplified images of the local structures are found in Fig. S3

Models	Co charge $ e^- $	$\Delta E_{\text{binding-Co}}$ (eV)	Co CN	Co–M distance (Å)
Co/melem-1	0.10	–1.11	1	2.07
Co/melem-2	0.15	–1.12	2	2.16 2.16
Co/2-melem-1	0.34	–3.14	2	1.86 1.86
Co/2-melem-2	0.68	–3.36	4/2	2.02 2.17 2.15 2.02 1.90(C) 1.90(C)
Co/2-melem-3	0.66	–3.31	4/2	2.03 2.10 2.00 2.10 2.45(C) 1.88(C)
Co/2-melem-4	0.70	–3.15	4/2	2.20 2.02 2.04 2.16 1.90(C) 1.90(C)
Co/melem-dimer	0.55	–2.78	2	1.86 1.86
Co/2-melem-dimer-1	0.96	–5.23	4	1.96 2.01 1.96 2.01
Co/2-melem-dimer-2	0.77	–5.28	4	1.91 1.91 1.91 1.91

binding with two melem molecules. Co/melem-dimer-1 refers to the first configuration with Co binding to a melem dimer, while Co/2-melem-dimer-1 refers to first configuration with Co binding to two melem dimers. In addition to the structures presented, we also identified other less stable structures, as well as structures that were similar to those in the main text. These alternative structures are included in Fig. S1.†

Binding energies for all molecular structures were found to be exothermic, ranging from –5.28 to –1.11 eV. The weakest binding occurred with just single melem molecules (e.g., –1.11 eV for Co/melem-1), where the Co interacted with just one or two N atoms. Notably, binding energies were more exothermic when Co interacted with two molecules (e.g., –3.36 eV for Co/2-melem-2), such as with two melem molecules or with two melem dimers. As noted above, Co binding with single melem molecules resulted in either one or two Co–N coordination. Co binding with other molecular carbon nitrides (such as multiple melems or dimers) typically resulted in larger coordination numbers (up to 6), as well as more exothermic binding energies (ranging from –5.28 to –2.78 eV). Notably, Co binding with two melems tended to form Co–C moieties, whereas no other carbon nitride molecules had such bonds.

Fig. S2† shows a comparison between binding energies and Co coordination number. R^2 values for a regression fit are around 0.5, indicating reasonably moderate correlation. Our results show that Co with higher coordination was generally most stable, as seen for example by Co/2 melems or Co/2 melem dimers, where the most stable structures have Co with four Co–N coordination (Co/2-melem-2 and Co/2-melem-dimer-2). Coordination values beyond four tend to have a small affect on binding energies (see Fig. S2†). We also observed a variety of geometries for the Co/melem complexes. Simplified representations indicating the local geometries and coordination coordination around the Co atom are provided in Fig. S3.† The local geometries are also listed in Table S4.† Notably, the bent and linear geometries had the weakest binding energies (–1.12 eV for Co/melem-2, –2.78 eV for Co/melem-dimer, and –3.14 eV for Co/2-melem-1), while the strained trigonal prismatic, tetrahedral, and square planar geometries had the strongest binding energies, ranging from –5.28 to –3.15 eV, likely due to the increased number of Co–X bonds in these later structures.

3.1.2 Binding of Co to partially condensed polymeric carbon nitrides. Fig. 3 shows structures for Co bound to a partially condensed polymeric carbon nitride, melon. Details of these calculations are summarized in Table 2. We present several different structures for Co binding to single-layer melon (denoted as Co/melon) and two-layer melon (denoted as Co/2-melon), with each configuration designated by a different number (e.g., 1, 2, etc.). The most stable structure with Co binding to single-layer melon was Co/melon-1 ($E_{\text{binding}} = -3.00$ eV), where Co was coordinated to two N atoms. An alternative structure (Co/melon-2) had a binding energy of –2.84 eV, and Co was coordinated to three N atoms. Simplified images of Co/melon structures illustrating the local geometries are found in

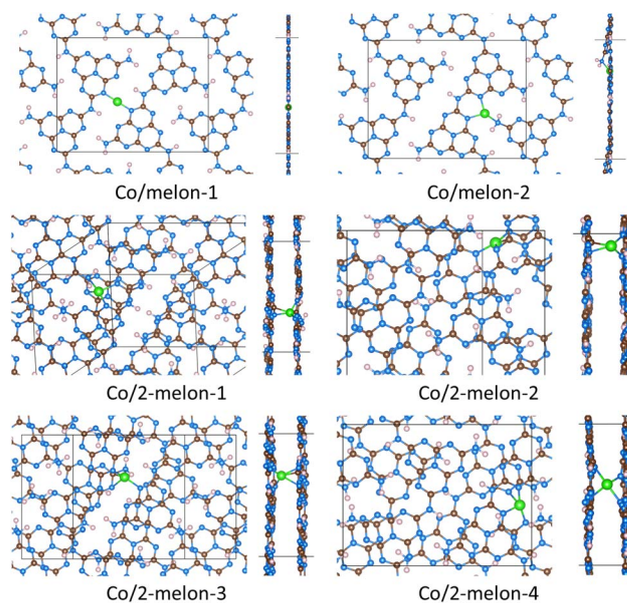


Fig. 3 Stable structures where Co was bound to melon strands. Brown spheres represent carbon atoms, blue spheres represent nitrogen atoms, pink spheres represent hydrogen atoms, and green spheres represent cobalt atoms. When two melons are present, these are designated by the prefix '2-'. Each unique configuration is identified by the final number (e.g., '-1', '-2', etc.). Top and side views are shown. Simplified images of the Co/melon structures are found in Fig. S4.†



Table 2 Results for Co bound to carbon nitride polymers. Shown are calculated Co Bader charges, Co binding energies, Co coordination numbers (Co–N coordination number, followed by Co–C coordination number), and distances between Co and closest atoms (typically N). In some cases Co was bound to C atoms, as indicated in the table. Structures in the table correspond to the same structures shown in Fig. 3 and 4. Simplified images of the local structures are found in Fig. S4

Models	Co charge [e^-]	$\Delta E_{\text{binding-Co}}$ (eV)	Co CN	Co–M distance (Å)
Co/melon-1	0.49	–3.00	2	1.83 1.83
Co/melon-2	0.72	–2.84	3	2.06 1.90 1.92
Co/2-melon-1	0.78	–3.36	4/1	1.96 2.30 1.94 2.01 1.99(C)
Co/2-melon-2	0.77	–3.61	3/1	1.99 1.97 1.93 1.94(C)
Co/2-melon-3	0.78	–3.46	4/1	1.94 2.02 2.36 1.98 1.96(C)
Co/2-melon-4	0.86	–3.35	4	2.02 2.16 2.01 2.06
Co/C ₃ N ₄ -1	0.71	–2.88	2/1	1.87 1.89 1.93(C)
Co/C ₃ N ₄ -2	0.75	–3.06	3/1	1.95 2.06 1.95 1.96(C)
Co/2-C ₃ N ₄ -1	0.80	–3.84	4/2	2.08 1.96 2.05 2.02 1.95(C) 1.95(C)
Co/2-C ₃ N ₄ -2	0.77	–3.92	3/3	1.94 1.91 1.98 2.10(C) 2.06(C) 2.48(C)
Co/2-C ₃ N ₄ -3	0.77	–3.87	3/2	1.96 1.95 1.90 1.96(C) 2.09(C)
Co/2-C ₃ N ₄ -4	0.82	–3.66	3/1	2.04 1.92 1.97 1.97(C)

Fig. S4.† In Co/melon-1, Co was bound to nitrogen atoms from two melon strands, forming a linear structure. Co/melon-2 formed a trigonal planar structure interacting with two strands of melon.

The binding energies of Co interacting with two melon layers ranged from –3.61 to –3.35 eV. Co is therefore more stable interacting with two melon layers compared to just one melon layer, although the binding energies were not drastically more exothermic (–3.61 eV for Co/2-melon-2 and –3.00 eV for Co/melon-1). Co binding with two-layer melon formed either three or four Co–N bonds with both the layers, and up to two Co–C bonds. The most stable configuration was Co/2-melon-2 ($E_{\text{binding}} = -3.61$ eV), where the Co was coordinated to three N atoms and one C atom, having a square planar structure. The local geometries are listed in Table S4† and shown in Fig. S4,† indicating a variety of Co complexes, including linear, trigonal planar, square pyramidal, square planar, and tetrahedral. We also observed strained square pyramidal (Co/2-melon-1 and Co/2-melon-3, both with four Co–N and one Co–C bonds), and tetrahedral (Co/2-melon-4 with four Co–N bonds) geometries. Our calculations show that Co binding to two-layer melon tended to form Co–C bonds, whereas Co binding to single layer melon did not form any Co–C bonds.

3.1.3 Binding of Co to fully condensed polymeric carbon nitrides. Following the modeling of Co/melon structures, we explored Co binding with fully polymerized carbon nitride g-C₃N₄. Fig. 4 shows results for Co bound to polymeric (periodic) g-C₃N₄, and details of these calculations are summarized in Table 2. We found two stable Co/C₃N₄ structures. Co/C₃N₄-1 had a binding energy of –2.88 eV. Co/C₃N₄-2 was the most stable g-C₃N₄ structure identified ($E_{\text{binding}} = -3.06$ eV). We also modeled Co binding with two-layer g-C₃N₄, and the binding energies were between –3.92 and –3.66 eV. The coordination numbers were between three and four for single-layer C₃N₄, and between four and six for two-layer C₃N₄. Notably, all the stable geometries with Co bound to C₃N₄ had Co–C bonds (one to three Co–C bonds). It is also worth mentioning that Co binding with 2 molecular carbon nitrides is about two times more stable than

Co binding to just one molecule; however, Co binding with 2-layer C₃N₄ is only ~30% more stable than Co binding with 1-layer C₃N₄. For example, there is not a drastic change in coordination for Co/C₃N₄-2 (four coordination with 3 Co–N bonds) compared to Co/2-C₃N₄-2 (six coordination with 3 Co–N bonds). Coordination geometries of all these structures are listed in Table S4, and shown in Fig. S4.†

3.1.4 Connection to experimental work. Some experimental work has suggested four-coordinated bonding with N (Co–N₄) when Co interacts with polymeric carbon

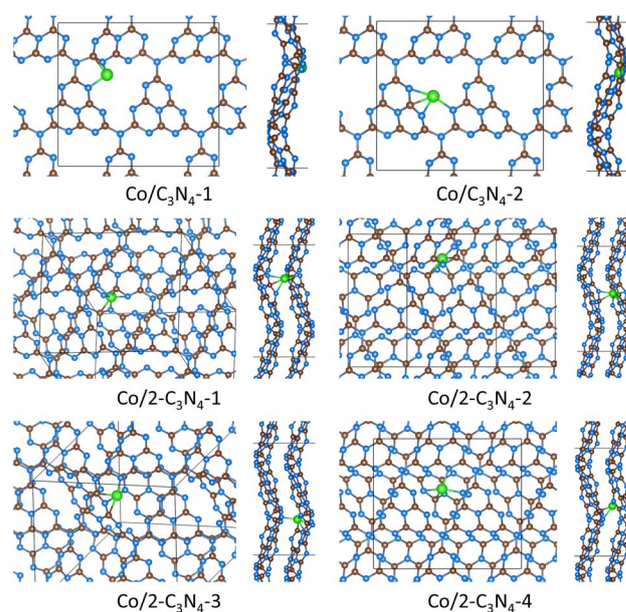


Fig. 4 Stable structures where Co was bound to g-C₃N₄. Brown spheres represent carbon atoms, blue spheres represent nitrogen atoms, pink spheres represent hydrogen atoms, and green spheres represent cobalt atoms. When two g-C₃N₄ sheets are present, these are designated by the prefix '2-'. Each unique configuration is identified by the final number (e.g., '-1', '-2', etc.). Top and side views are shown. Simplified images of Co/C₃N₄ structures are found in Fig. S4.†



nitrides.^{61,107,109} We observed four-coordinated Co in Co/C₃N₄-2, where 3 Co–N and 1 Co–C bonds formed. Fig. S1† shows that Co/2-melem-6 forms a Co–N₄ arrangement, although this structure was less stable than other melem structures. In our previous work,⁷⁴ we identified a similar possible Co–N₄ geometry, although this involved Co binding with two heptazine molecules (chosen to mimic carbon nitride edges), rather than melem, melon, or g-C₃N₄. It should be noted that experimental work, such as Extended X-ray Absorption Fine Structure (EXAFS), may have trouble distinguishing between Co–N and Co–C bonds.^{110–113} Thus experimental work from literature may be observing both Co–N and Co–C bonds (such as in Co/C₃N₄-2, or Co/2-melon-2) or Co interacting with carbon nitride edges (such as in our previous work⁷⁴). There may also be fitting errors in methods such as EXAFS due to low metal concentration.¹¹³ There is some literature that suggests the formation of M–C bonds. Fao *et al.*'s computation work¹¹⁴ reported Co coordinated to three N atoms and one C atom with a single-layer corrugated g-C₃N₄. Zheng *et al.*'s computation work⁷⁵ found Co binding with corrugated g-C₃N₄ to form three Co–N bonds and two Co–C bonds, although no bond lengths were reported. Wang *et al.*'s work¹¹⁵ combined experimental results, EXAFS and X-ray photoelectron spectroscopy (XPS), with DFT results to show Cu binding to g-C₃N₄ forming one Cu–C and two Cu–N bonds.

Identification of the exact carbon nitride structure from experiment is not trivial, and it is possible that some literature work has misidentified their carbon nitride structure. See the introduction for discussion on partially condensed carbon nitrides and the difficulty of synthesizing g-C₃N₄. For example, temperature plays a vital role in what carbon nitride structure is synthesized and observed.⁴⁰ It is possible that experimental papers finding four-coordinated Co atoms could be Co binding to partially condensed carbon nitrides. For example, Huang *et al.*'s experimental work¹¹⁶ argued that Co formed four identical Co–N bonds when incorporating Co into a carbon nitride. Their experimental work identified a significant number of NH₂ groups in the carbon nitride, which suggests a structure other than fully condensed g-C₃N₄. This configuration resembles Co/2-melem-dimer-2 found in our study, as our structure also had four identical Co–N bonds. Ding *et al.*'s work¹⁰⁹ presented a symmetric Co–N₄ structure with 1.98 Å, similar to Co/2-melem-dimer-1, which has four Co–N bonds all similar in length. However, in Xiong *et al.*'s work,¹¹⁷ their experimental results indicted a Co coordination of 2.3, with lengths of 1.95 Å, similar to several of our modeled carbon nitrides: Co/melem-2, Co/2-melem-1, or Co/melem-dimer.

3.2 Electronic properties of Co-carbon nitride materials

3.2.1 Bader charge analysis. We further examined the electronic properties of the Co/carbon nitride structures. Tables 1, 2, and S4† provide Co Bader charges of different structures, which indicate the oxidation state of the Co atoms. Co became positively charged in all cases once it interacted with the carbon nitride supports. This agrees with experimental^{61,67,118} and computational^{75,119} observations on the cationic nature of Co. Larger Co Bader charge indicated more oxidation of Co, and

presumably more chemical interactions between the Co and carbon nitride substrate. The Co charges in Co/molecular carbon nitrides ranged from +0.1 to +0.96 |e⁻|, with an average value of +0.55 |e⁻|. With polymeric carbon nitrides, Co charges were similar for single-layer and two-layer structures, ranging from +0.72 to +0.86 |e⁻|, except for Co/melon-1, which had a Co charge of +0.49 |e⁻|. The average Co charge with the polymeric carbon nitrides was +0.75 |e⁻|. Thus, the Co atoms were all cationic when bound to carbon nitride molecules, and Co atoms bound to polymeric carbon nitrides were generally more cationic than Co bound to molecular carbon nitrides.

We next determined the relationships between the Co charges and structure of the complexes. Fig. 5 illustrates how the charges correlate to Co coordination number, binding energies, and structure of the carbon nitride support. The plots (a) and (b) show general trends that the Co charge increases (*i.e.*, becomes more oxidized) with both increasing Co–N coordination number and total Co coordination number. This connects with our observation that the Co charge was larger when Co was bound to more molecules (*e.g.*, two melem molecules compared to one molecule), indicating that more chemical interactions (as evidenced through increased coordination number) lead to stronger charge transfer between Co and neighboring atoms. Indeed, Fig. S2† demonstrates how coordination number correlates to binding energy. Plot (c) shows that the Co charge increases with stronger binding (*i.e.*, more negative binding energy). These results again indicate that stronger interactions between the Co atom and nearby bonding atoms lead to more charge transfer. Finally, we show in plot (d) how the size of the carbon nitride substrate correlates to Co charge. Larger carbon nitrides have more triazine rings, and generally bind stronger to Co (see Table 2), leading to more oxidation of the Co atom. Our results connecting the Co charge to structure are especially important in deciphering experimental data from SACs supported on carbon nitrides. For example, these trends could help interpret coordination numbers and/or oxidation states obtained through methods, such as from XAS.¹²⁰ Indeed, the oxidation state of Co has been linked to its reactivity.⁶⁷

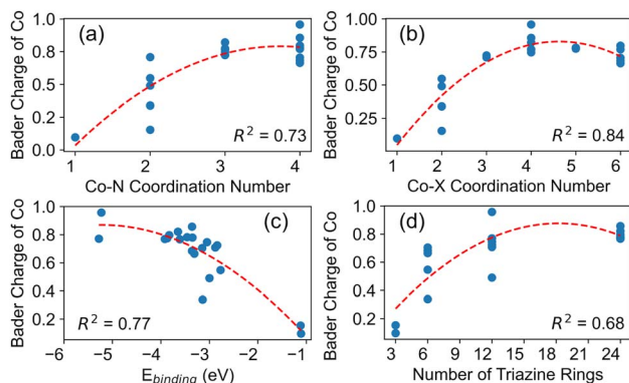


Fig. 5 Correlations between the Bader charges of Co atoms and (a) Co–N coordination numbers, (b) total Co coordination numbers (both C and N), (c) Co binding energies, and (d) size of the carbon nitride (*i.e.*, number of triazine rings). The red dashed lines are second-order polynomial fits.



3.2.2 Band gaps. A key property for photocatalysis is the band gap, as a lower band gap enables a broader spectrum of light to initiate photoexcitation. Co atoms may create favorable reaction sites (as discussed in Section 3.3), but also facilitate increased photoexcitation in Co/carbon nitride photocatalysts. Experimental work shows Co supported on carbon nitrides leads to a reduction in band gaps,^{118,121–123} although the exact structures of the carbon nitrides in several of these papers still need clarification. The reduction of the band gap caused by binding of a Co atom to carbon nitrides has also been demonstrated in computational work involving g-C₃N₄.^{78,79} On the other hand, computational work on the effect of Co paired with other carbon nitrides (such as melem or melon) is missing. Therefore, we calculated band gaps of Co interacting with a variety of carbon nitrides.

We show the band gaps of pristine carbon nitrides and the most stable Co/carbon nitrides calculated using the HSE06 hybrid functional in Fig. 6. Our results demonstrate that the band gap decreases with increased polymerization of the carbon nitrides (see left panel of Fig. 6), as already suggested in literature.³⁶ Notably, the incorporation of a Co atom significantly reduces the band gaps (see right panel of Fig. 6). For example, the band gap of Co/2-melem-2 is 2.61 eV, which is 2 eV lower than a pristine melem molecule. Other Co/carbon nitride band gaps are much smaller, being in the range of 0.15 to 1.49 eV. There does not appear to be a clear relationship between carbon nitride size and Co/carbon nitride band gap. All Co/carbon nitrides have band gaps < 1.2 eV, with the exception of Co/2-melem-2 (2.61 eV) and Co/2-C₃N₄-2 (1.49 eV). However, our results do show that Co does indeed cause band gap reduction for Co supported on a variety of carbon nitrides.

3.2.3 Electronic states. In the following section we briefly analyze the electronic states to better understand how Co leads to diminished band gaps and better photocatalysis. We show the density of states (DOS) using the HSE06 functional of carbon nitride molecules in Fig. S6 and polymers in S7.† The valence bands of the bare carbon nitrides consist mainly of states from nitrogen, and the conduction bands consist of states from carbon and nitrogen. Upon addition of Co, however, states arise in the carbon nitride band gaps consisting primarily

of Co electronic states. These states effectively lower the band gaps as lower energies are needed to excite electrons from these gap states to the conduction bands. Previous literature indicated that similar gap states could arise over g-C₃N₄,¹²⁴ and a melon-like structures.⁸² Our work shows that these gap states arise, regardless of the degree of polymerization (*i.e.*, molecular, partially condensed, and fully condensed carbon nitrides). Experiment also indicates that Co lowers electron-hole recombination when added to carbon nitrides.¹²³ This lowered recombination could be facilitated by the midgap states created by Co in all carbon nitrides.

3.3 CO₂ activation over Co-carbon nitride materials

CO₂ photoreduction involves many different steps, and a key step is the activation of CO₂. The formation of bent, anionic CO₂ is important, as this enables the further reactivity of this otherwise stable linear molecule.¹²⁵ If formation of bent, adsorbed CO₂ is unfavorable, then CO₂ reduction will be slow or hindered. We note that CO₂ adsorption is weak on pure carbon nitrides, as the process has been reported for single-layer g-C₃N₄ to be slightly exothermic (−0.42 eV)¹²⁶ or endothermic (0.02 eV,⁷⁸ and 0.24 eV).⁷² Thus, improving the adsorption and reactivity of carbon nitrides is needed, such as pairing them with SACs. Experimental work on Co/carbon nitride catalysts shows these to be potential catalysts for CO₂ reduction.^{67,68,71–74} In this section we examine the formation of activated CO₂ over the various Co/carbon nitride structures.

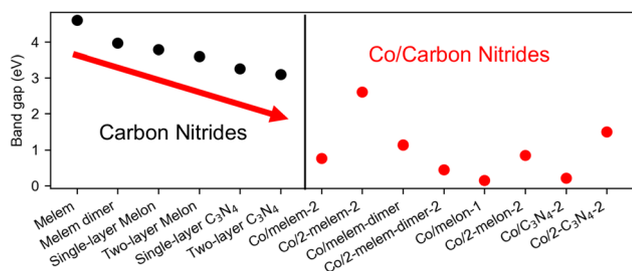


Fig. 6 Band gaps of pristine carbon nitrides and the most stable Co/carbon nitrides calculated using the HSE06 functional. The left graph shows the band gaps for pristine carbon nitrides, while the right graph shows the band gaps for the most stable Co/carbon nitrides. The arrow shows the trends in band gaps for carbon nitrides, which decrease with increasing polymerization.

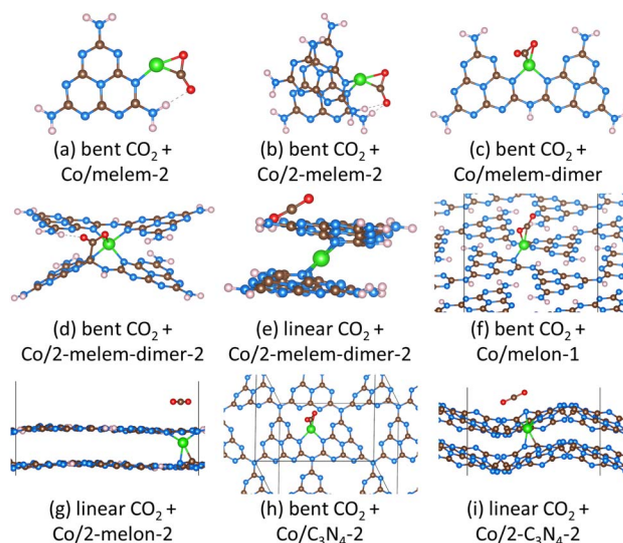


Fig. 7 Most stable structures after CO₂ adsorption to Co-carbon nitride materials. Shown are (a) bent CO₂ + Co/melem-2, (b) bent CO₂ + Co/2-melem-2, (c) bent CO₂ + Co/melem-dimer, (d) bent CO₂ + Co/2-melem-dimer-2, (e) linear CO₂ + Co/2-melem-dimer-2, (f) bent CO₂ + Co/melon-1, (g) linear CO₂ + Co/2-melon-2, (h) bent CO₂ + Co/C₃N₄-2, and (i) linear CO₂ + Co/2-C₃N₄-2. Brown spheres represent carbon atoms, blue spheres represent nitrogen atoms, pink spheres represent hydrogen atoms, red spheres represent oxygen atoms, and green spheres represent cobalt atoms.



Table 3 Adsorption energies and deformation energies of linear and bent CO₂ on different Co/carbon nitride systems. The most stable CO₂ for each Co/carbon nitride is marked in gray. E_{ads} refers to the adsorption energy, $E_{\text{Co/CN deform}}$ is the deformation energy of the Co/carbon nitride $E_{\text{CO}_2\text{deform}}$, represents the deformation energy of CO₂, and E_{int} is the interaction energy between the adsorbent and CO₂

Co/CN structures	CO ₂ configuration	O–C–O angle (°)	CO ₂ charge $ e^- $	E_{ads} (eV)	$E_{\text{Co/CN deform}}$ (eV)	$E_{\text{CO}_2\text{deform}}$ (eV)	E_{int} (eV)
Co/melem-2	Bent	141.0	−0.67	−2.33	−0.03	1.55	−3.85
	Linear	177.7	−0.03	−0.16	0.01	0.01	−0.18
Co/2-melem-2	Bent	139.0	−0.74	−1.12	1.29	1.57	−3.98
	Linear	176.9	−0.02	−0.23	0.03	0.01	−0.27
Co/melem-dimer	Bent	141.6	−0.60	−1.29	0.22	1.38	−2.89
	Linear	177.7	−0.10	−0.44	0.03	0.01	−0.48
Co/2-melem-dimer-2	Bent	128.5	−0.63	−0.38	1.22	2.25	−3.86
	Linear	175.7	−0.04	−0.38	−0.01	0.01	−0.38
Co/melon-1	Bent	141.1	−0.60	−0.81	0.66	1.44	−2.91
	Linear	180.0	0.00	−0.05	0.00	0.00	−0.05
Co/2-melon-2	Bent	141.7	−0.60	0.02	0.74	1.21	−1.93
	Linear	179.1	0.00	−0.21	0.00	0.00	−0.20
Co/C ₃ N ₄ -2	Bent	145.0	−0.47	−0.79	0.34	1.17	−2.30
	Linear	175.8	−0.01	−0.31	0.11	0.02	−0.45
Co/2-C ₃ N ₄ -2	Bent	141.5	−0.57	1.13	2.00	1.28	−2.16
	Linear	179.0	0.00	−0.31	0.00	0.00	−0.31

3.3.1 Formation of bent CO₂. We modeled both linear and bent CO₂ adsorbed on our most stable Co/CN structures. Fig. 7 presents only the most stable configurations (*i.e.*, lowest CO₂ adsorption energies) for adsorbed CO₂, while Fig. S10† shows all the CO₂ structures we identified. Table 3 also provides adsorption energies and Bader charges for linear and bent CO₂ on the different Co/CN structures. All the Co/carbon nitrides activated CO₂, except for two-layer melon (Co/2-melon-2) and two-layer C₃N₄ (Co/2-C₃N₄-2). Several adsorption energies of bent CO₂ were quite exothermic (from −2.33 to −1.12 eV), including Co/melem-2, Co/2-melem-2, and Co/melem-dimer. In all cases of bent CO₂, the CO₂ molecules exhibited O–C–O angles between 128.5° to 145.0°, and the C–O bonds were elongated to 1.20–1.30 Å (compared to the gas-phase CO₂ bond length of 1.17 Å), indicating activated states, similar to previous work on other materials.^{127–129} The Bader charge analysis also shows that bent CO₂ molecules were negatively charged from −0.74 $|e^-|$ to −0.47 $|e^-|$, while the linear CO₂ molecules were close to neutral (−0.10 $|e^-|$ to 0.00 $|e^-|$). This again, confirms the anionic, activated state of bent CO₂.

These results demonstrated that Co could significantly improve CO₂ adsorption and activation on carbon nitride materials. The Co atoms can facilitate charge transfer and serve as active centers to adsorb and activate CO₂ molecules. As we mentioned previously, adsorption of CO₂ on pristine g-C₃N₄,^{72,78,126,130} and melon¹³¹ was reported to form linear, weakly bound CO₂. Co/molecular carbon nitrides and Co/single-layer polymeric carbon nitrides could however activate CO₂ into bent, exothermic structures. The adsorption energies of linear and bent CO₂ over Co/2-melem-dimer-2 were equivalent to each other. We modeled both linear and bent CO₂ adsorbed to the Co atom in several different orientations. CO₂, however, was only weakly absorbed to Co in the bent structure (Fig. 7d), likely due to steric hindrance as the CO₂ situated between two carbon nitride molecules. In the linear structure, the Co atom preferred to interact only with the carbon nitride (Fig. 7e), despite the

initial geometry with CO₂ placed interacting with the Co atom. Again, steric hindrance prevents linear CO₂ from interacting with the Co atom. In addition, we found that Co/two-layer polymeric carbon nitrides (melon and g-C₃N₄) only weakly absorbed linear CO₂, and activation of CO₂ into the bent form was endothermic. The Co atoms are sandwiched between the two carbon nitride layers for both carbon nitrides. A linear CO₂ molecule placed between the two carbon nitride layers, to interact with the Co atom, was sterically hindered, and the CO₂ molecules moved to the locations shown in Fig. 7g and i. Our efforts to form bent CO₂ in these two-layered structures led to displacement of Co, and weakening of Co-carbon nitride interactions, as illustrated in Fig. S10.† As shown, in Co/2-melon-2 the Co atom migrates away from one melon layer to bond with a bent CO₂. The Co atom after CO₂ adsorption only interacts with one carbon nitride layer, and is therefore weakened compared to its pre-adsorbed state (Co interacting with two carbon nitride layers). In the case of Co/2-C₃N₄-2, an adsorbed, bent CO₂ interacts with the Co atom as it sits between the two carbon nitride layers. The CO₂ molecule is thus confined between the two layers and experiences steric repulsion, making activation of CO₂ unfavorable.

In order to further understand the role of the carbon nitride support, we plotted the stable bent CO₂ adsorption energies *versus* carbon nitride size in Fig. 8. These results show that smaller carbon nitride supports have the strongest adsorption of bent CO₂, while larger carbon nitrides have the weakest CO₂ adsorption energies. The Sabatier principle prescribes that reactant adsorption energies should neither be too strong or too weak. In this case, intermediate adsorption energies occur for Co/melon-1 (−0.81 eV), Co/2-melem-2 (−1.12 eV), and Co/melem-dimer (−1.29 eV). Thus, based on the Sabatier principle, some molecular and partially condensed complexes may give the best CO₂ reduction activity, although further clarification of the reaction mechanism is needed. We should also note that molecular carbon nitrides or melon strands may be



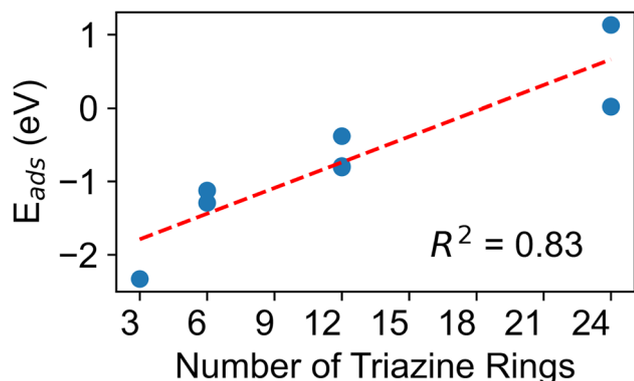


Fig. 8 Correlation between the adsorption energies of CO_2 over Co/carbon nitrides and the size of the carbon nitride (i.e., number of triazine rings). The red dashed line shows a linear fit.

representative of exposed carbon nitride edges, which may be a very reactive region.¹³²

3.3.2 The nature of linear and activated CO_2 . Generally bent, activated CO_2 was more stable on the Co/carbon nitrides compared to linear CO_2 , with the exception of two-layer structures, Co/2-melon-2 and Co/2- C_3N_4 -2. We next analyze the different structures to further explain the nature of linear *versus* bent CO_2 . For bent CO_2 there were significant changes in Co geometry after CO_2 adsorption. For structures that activated CO_2 (all but two-layer carbon nitrides), we observed elongation of the original Co-N bonds and breaking of the Co-C bonds. But, there were also newly formed Co-C and Co-O bonds with the CO_2 molecule. For example, in Co/2-melem-2, the Co atom had four Co-N and two Co-C bonds. After CO_2 adsorbed, the number of Co-N bonds reduced to two, and the two Co-C bonds broke, while new Co-C and Co-O bonds formed with CO_2 . For structures that favorably bound linear CO_2 (Fig. 7g and i), the geometries and Co coordination environment remained essentially the same as the pre-adsorption state. We also found that Co became more cationic upon interacting with CO_2 . For instance, the average charge of Co over carbon nitrides that activated bent CO_2 increased by $+0.22|e^-|$. On the other hand, the average Co charge for those structures that preferred linear CO_2 did not change, again indicating weak interactions.

In order to better understand why certain Co/carbon nitrides preferred linear over bent CO_2 , we examined the deformation and interaction energies of the complexes, which are listed in Table 3. The adsorption energy reflects a balance between the deformation of both the adsorbate and substrate upon adsorption, and their chemical interactions. The deformation energies embody the geometric changes of the adsorbate and substrate from their pre-adsorbed to post-adsorbed states. The interaction energy represents the bond formation energy between the adsorbate and substrate.¹³³⁻¹³⁶ The adsorption energy can be written as: $\Delta E_{\text{ads}} = E_{\text{int}} + E_{\text{Co/CN deform}} + E_{\text{CO}_2 \text{ deform}}$. Thus, whether adsorption is exothermic or not depends on the nature of structural changes upon adsorption and the nature of favorable (or unfavorable) electronic interactions. Deformation

energies (E_{deform}) of Co/carbon nitrides and CO_2 adsorption were calculated by eqn (3) and (4):

$$E_{\text{Co/CN deform}} = E'_{\text{Co/Carbon Nitride}} - E_{\text{Co/Carbon Nitride}} \quad (3)$$

$$E_{\text{CO}_2 \text{ deform}} = E'_{\text{CO}_2} - E_{\text{CO}_2} \quad (4)$$

Here $E'_{\text{Co/Carbon Nitride}}$ is the energy of the Co/carbon nitride complex after CO_2 adsorption, but with CO_2 removed. $E_{\text{Co/Carbon Nitride}}$ is the energy of the geometry of the optimized complex with no CO_2 present. E' is the energy of the adsorbed CO_2 minus the Co/carbon nitride complex. E_{CO_2} is the energy of gas-phase CO_2 . The interaction energy (E_{int}) was calculated by eqn (5):

$$E_{\text{int}} = \Delta E_{\text{ads}} - E_{\text{Co/CN deform}} - E_{\text{CO}_2 \text{ deform}} \quad (5)$$

CO_2 deformation energies were always positive for bent adsorption (average value of 1.48 eV), indicating an energy cost to bend the molecule into its activated state. The Co/carbon nitride deformation energies for the bent CO_2 case were quite varied, ranging from -0.03 to 2.00 eV, with an average value of 0.81 eV. Since deformation energies for the bent CO_2 case were positive, interaction energies must be sufficiently negative for adsorption to be exothermic. Indeed, our results show this to be the case, as these interaction energies were all negative (ranging from -1.93 to -3.98 eV). This strong interaction between bent CO_2 and the Co/carbon nitride overcomes geometric distortions to enable exothermic adsorption of CO_2 . Indeed, bent CO_2 becomes anionic, further indicating strong interactions occurring with bent CO_2 . The significantly more exothermic adsorption energies of bent CO_2 molecules on Co/melem-2, Co/2-melem-2, and Co/melem-dimer can be attributed to their more negative interaction energies (from -2.89 to -3.98 eV) compared to Co/melon-1 and Co/ C_3N_4 -2 (-2.91 eV and -2.30 eV).

On the other hand, interaction energies for linear CO_2 were much weaker, ranging from -0.05 to -0.48 eV. Despite CO_2 deformation energies for linear CO_2 being close to zero, the interaction energies were not exothermic enough to lead to strong linear CO_2 adsorption. CO_2 charges were close to neutral

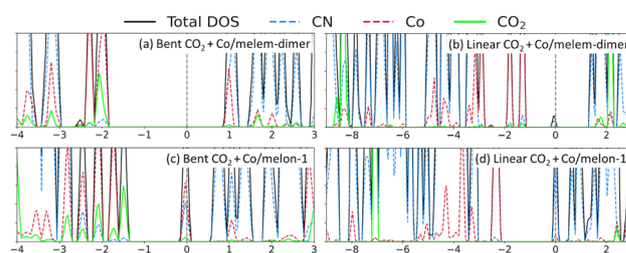


Fig. 9 Example density of states (DOS) plots for adsorbed CO_2 , indicating how linear and bent CO_2 interact with Co/carbon nitrides: (a) bent CO_2 + Co/melem-dimer, (b) linear CO_2 + Co/melem-dimer, (c) bent CO_2 + Co/melon-1, and (d) linear CO_2 + Co/melon-1. The total, carbon nitride (CN), Co, and CO_2 DOS are shown. The energies are relative to the Fermi level, which has been shifted to 0 eV and indicated by a dashed vertical line. Due to smearing of the electronic states, the Fermi level appears within the valence band, even though the HOMO energy level is below the Fermi level.



for linear CO₂, indicating little charge transfer and weak interactions between CO₂ and the Co/carbon nitrides.

DOS plots also provide insight on the nature of interactions between CO₂ and the Co/carbon nitride catalysts. We show in Fig. 9 example DOS plots for linear and bent CO₂ interacting with different Co/carbon nitride complexes. Other DOS plots are shown in Fig. S8 and S9.† These plots show that CO₂ in the bent position interacts strongly with Co. See for example the many overlapping peaks involving Co and CO₂ near -2 , -3.3 , and -3.8 eV for Co/melem-dimer. Similar overlapping peaks occur at -1.8 , -2.1 , -2.5 , and -2.9 eV for Co/melon-1. These interactions are due to bond formation between the CO₂ molecule and the cobalt atom. On the other hand, such interactions are missing for linear CO₂. The CO₂ electronic levels are all much lower in energy and overlap with Co is minimized. Both the interactions energies and density of state plots confirm that bent CO₂ forms strong bonds with the Co/carbon nitride while linear CO₂ does not.

3.4 Implications for photocatalysis

Experimental work shows that addition of Co to carbon nitrides improves photoactivity.^{67,78,116} The exact structure of carbon nitride in these experiments is not fully resolved. We modeled non-polymerized, partially condensed, and fully condensed carbon nitrides, which should provide details on several possible carbon nitrides that could be present in real-world catalysts. Our results showed that Co/carbon nitrides have better photoexcitation compared to pristine carbon nitrides, through more narrow band gaps. This should lead to more photoexcited electrons that may enable photocatalytic reduction. Experiment also indicates that Co lowers electron-hole recombination when added to carbon nitrides.¹²³ This lowered recombination could be facilitated by the mid-gap states created by Co, which we observed. Our results also indicate that in addition to facilitating a larger number of charge carriers to participate in photocatalytic reactions, Co/carbon nitride catalysts also are effective for CO₂ activation. In particular, we found that molecular and partially condensed carbon nitride complexes with CO₂ may have moderate CO₂ adsorption energy (neither too strong nor too weak) to enable CO₂ reduction. On the other hand, multi-layer carbon nitrides were found to have difficulties activating CO₂. Future work may assess the full catalytic mechanism, but in the least, CO₂ activation is a crucial reaction step which Co/carbon nitrides readily enable. All these different effects caused by Co paired with different carbon nitride supports (increased photoexcitation, decreased charge recombination, and increased reactivity) lead to better CO₂ reduction and photocatalytic activity (as observed in experiments⁶⁷⁻⁷⁴).

4 Conclusions

Literature has largely focused on two-dimensional g-C₃N₄ as a support for SACs, despite evidence showing that partial condensation is likely common in working carbon nitride catalysts. We assessed single Co atoms supported on a variety of

carbon nitrides as potential photocatalysts. We modeled single Co atoms on six different carbon nitrides with varying degrees of polymerization, including molecular (melem and melem dimer), partially condensed single-layer and two-layer melon, and fully condensed (single-layer and two-layer C₃N₄). Several stable Co/carbon nitrides were identified with Co forming up to four Co-N bonds, and three Co-C bonds having various Co local geometries. We also discovered correlations between Co coordination number, Co charge, and binding energy. As the Co coordination number increased, the Co atom generally became more oxidized, and the binding energy became more exothermic. Furthermore, binding of Co tended to be strongest with larger carbon nitrides.

In addition, we predicted how Co supported on a carbon nitride could function as a photocatalyst for CO₂ reduction. Adding Co to the various carbon nitrides lowered the band gaps, which should lead to better photoexcitation yield. Co atoms also helped activate CO₂ into a bent, anionic state, an essential step for CO₂ reactivity. The activation of CO₂ is facilitated by strong interactions between the CO₂ molecule and Co/carbon nitride, which is required to overcome unfavorable bending of the CO₂ molecule. For example, the interaction energy between a bent CO₂ molecule and Co/melem-2 is -3.85 eV, while the deformation energy of the bent CO₂ molecule is 1.55 eV, leading to an overall favorable activation of CO₂. Exothermic bent CO₂ adsorption energies varied between -2.33 and -0.38 eV, and likely a moderate CO₂ adsorption energy (not too strong and not too weak, such as near -1 eV) would best facilitate CO₂ reduction. Two melem molecules, a melem dimer, melon, and single-layer g-C₃N₄ all have moderate CO₂ adsorption energies, and thus may be good candidates for CO₂ photoreduction. Our work shows that carbon nitrides other than g-C₃N₄ when paired with single metal atoms have potential as photocatalysts, and therefore should be studied further.

Data availability

Data from the various tables and figures for this article have been included in the ESI.† Simulation files, including geometries, can be found at <https://github.com/Deskins-group/Structure-Files/tree/master/Co-Carbon-Nitrides>.

Conflicts of interest

There are no conflicts to declare.

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