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Well-defined surface catalytic sites for solar CO₂ reduction: heterogenized molecular catalysts and single atom catalysts

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Exciting progress has been made in the area of solar fuel generation by CO₂ reduction. New photocatalytic materials containing well-defined surface catalytic sites have emerged in recent years, including heterogenized molecular catalysts and single atom catalysts. This Feature Article summarizes our recent research in this area, together with brief discussions of relevant literature. In our effort to obtain heterogenized molecular catalysts, a diimine-tricarbonyl Re(I) complex and a tetraaza macrocyclic Co(III) compound were covalently attached to different surfaces, and the effects of ligand derivatization and surface characteristics on their structures and photocatalytic activities were investigated. Single atom catalysts combine the advantages of homogeneous and heterogeneous catalysis. A single-site cobalt catalyst was prepared on graphitic carbon nitride, which demonstrated excellent activity in selective CO₂ reduction under visible-light irradiation. Doping carbon nitride with carbon was found to have profound effects on the structure and activity of the single-site cobalt catalyst. Our research achievements are presented to emphasize how spectroscopic techniques, including infrared, UV-visible, electron paramagnetic resonance, and X-ray absorption spectroscopies, could be combined with catalyst synthesis and computation modeling to understand the structures and properties of well-defined surface catalytic sites at the molecular level. This article also highlights challenges and opportunities in the broad context of solar CO₂ reduction.

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Ehab Shaaban

Ehab Shaaban obtained his PhD degree from the University of New Hampshire in 2023 under the supervision of Prof. Gonghu Li. His dissertation was focused on highly dispersed metal sites on semiconductor surfaces for small molecule activation. He investigated structural and binding properties that emerge from metal-metal oxide interactions. His main research interests are on the development and investigation of nanostructured catalysts.



1. Introduction

Carbon dioxide (CO₂) is considered a renewable C1 feedstock for the production of valuable chemicals and fuels.^{1,2} Sufficient energy and catalysts are needed to activate CO₂, which is a very stable molecule. Solar-driven CO₂ reduction has been extensively investigated in recent years as it employs natural sunlight as a renewable energy source. Various photocatalysts capable of activating CO₂ under light irradiation have been reported in the literature, including molecular complexes, inorganic semiconductors, and other systems.^{3–7}

In molecular systems, the photocatalyst could be a transition metal complex that can harvest light and activate CO₂, or a molecular CO₂-reduction catalyst combined with a photosensitizer.^{8–11} An example of the former is Re(bpy)(CO)₃Cl (denoted “Re-bpy”), where bpy is 2,2′-bipyridine. This diimine-tricarbonyl Re(I) compound and its supramolecular complexes have demonstrated excellent reactivity and selectivity towards CO formation in photocatalytic CO₂ reduction.^{12,13} Examples of the latter include [Co(cyclam)Cl₂]Cl (denoted “Co-cyclam”), where cyclam is 1,4,8,11-tetraazacyclotetradecane. This macrocyclic complex requires the use of photosensitizers, such as *p*-terphenyl, to harvest light and generate electrons needed for CO₂ reduction.^{14,15} While these molecular

photocatalysts are highly active and selective towards CO₂ reduction, many suffer from poor stability under photochemical conditions. In addition, most molecular complexes require the use of non-renewable electron donors to maintain reactivity in CO₂-reduction photocatalysis.

In comparison, inorganic semiconductors,^{16–21} most notably titanium dioxide (TiO₂)-based materials, are robust heterogeneous photocatalysts for CO₂ reduction since the early work by Inoue and co-workers.²² One attractive feature of such photocatalysts is that some semiconductors (e.g. TiO₂) with appropriate band positions allow the use of water as a renewable electron donor for CO₂ reduction. However, most semiconductor photocatalysts are inefficient in converting photonic energy into chemical energy due to recombination of photo-generated charge carriers (electrons and holes) that prevails in the bulk.^{23,24} Furthermore, such heterogeneous photocatalysts often lack well-defined surface sites for selective CO₂ reduction.

Different strategies have been designed to prepare robust and efficient photocatalysts containing well-defined surface catalytic sites (Fig. 1). For instance, heterogenized molecular catalysts can be obtained by attaching molecular catalysts onto light-absorbing surfaces.^{25–31} Such photocatalysts generally show enhanced activity and improved photostability. Atomically dispersed catalysts or single-atom catalysts (SACs) are known to have well-defined active sites on a solid support.^{32–39} SACs have demonstrated distinct properties in a wide range of chemical reactions, including photocatalysis.^{40–43} Recently, there has been enormous interest in developing SACs for use in solar CO₂ reduction.^{44–48}

This Feature article summarizes our recent effort to construct well-defined catalytic sites on surface for use in solar CO₂ reduction. In particular, Re-bpy and Co-cyclam were attached to different surfaces to obtain heterogenized molecular catalysts. These two complexes were chosen in our studies due to their well-known activity in CO₂-reduction catalysis and their relatively simple molecular structures that allow facile ligand derivatization for surface attachment. Our research in this direction has been focused on the effects of ligand derivatization and surface characteristics on the photocatalytic activities of the heterogenized molecular catalysts. As will be discussed,



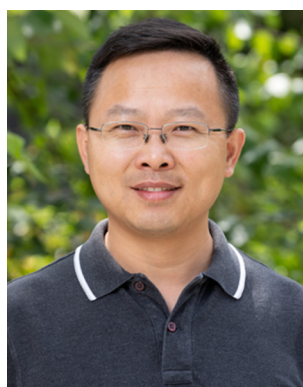
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Allison St. John

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Gonghu Li

Gonghu Li received his PhD degree in Chemistry from the University of Iowa in 2005 and conducted postdoctoral research at Northwestern University and Yale University. In 2009, he began his independent career in the Department of Chemistry at the University of New Hampshire, where he is now a Full Professor. His current research projects are focused on surface chemistry and photocatalysis for efficient solar fuel generation.



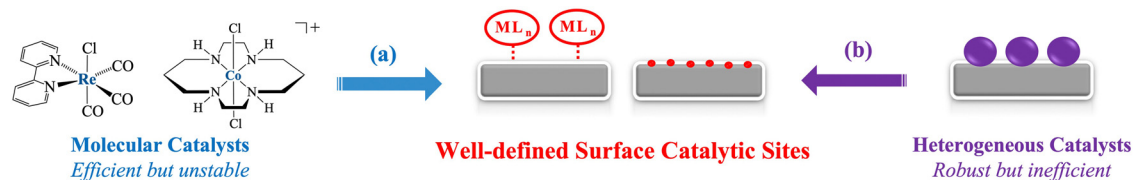


Fig. 1 Approaches to obtain well-defined surface catalytic sites for efficient solar CO₂ reduction: (a) heterogenization of molecular catalysts and (b) atomic dispersion of supported metal nanoparticles. Re-bpy and Co-cyclam are shown here as examples of molecular CO₂-reduction catalysts (ML_n). The third Cl[−] ligand for Co-cyclam is not shown here.

heterogenization of the molecular catalysts on surfaces allowed us to investigate their structures and complexation using spectroscopic techniques, which could be less informative and more challenging in studying the molecular catalysts in homogeneous solutions. Inspired by the CO₂-reduction activity of Co-cyclam and prior work by others in the area of SACs, we discovered a Co SAC on graphitic carbon nitride (C₃N₄) capable of catalyzing selective CO₂ reduction under visible-light irradiation. The Co SAC structure was interrogated by using a combination of synthesis, spectroscopy, and modeling studies. Relevant research results by others are also discussed to highlight the progress and challenges in the broad context of solar CO₂ reduction using innovative photocatalysts.

2. Solar CO₂ reduction using Re-bpy covalently attached to SiO₂

As a model molecular catalyst, Re-bpy has been covalently attached to semiconductor surfaces,^{49–57} including photoelectrodes,^{58,59} to achieve selective CO₂ reduction under solar irradiation. Despite the fact that Re-bpy itself is capable of catalyzing CO₂ reduction under light irradiation, the semiconductor support could further improve the photocatalytic performance of Re-bpy. For example, Windle and co-workers⁵² prepared a diimine-tricarbonyl Re(i) complex derivatized with phosphonate groups for immobilization on a TiO₂ surface. The heterogenized Re(i) catalyst demonstrated significantly higher activity than Re-bpy in solar CO₂ reduction under the same condition. Results obtained using transient absorption spectroscopy revealed that the enhancement in activity upon heterogenization was due to an increase in the lifetime of a reduced Re intermediate on the TiO₂ surface. A later study by Abdellah and co-workers⁵³ revealed the role of TiO₂ as an electron reservoir, slowing down charge recombination and improving photocatalysis.

We employ SiO₂ as a model support for Re-bpy to carry out spectroscopic investigations since (i) SiO₂ is almost “transparent” in studies using many spectroscopic techniques, and (ii) Re-bpy itself is capable of harvesting light and mediating CO₂ reduction. Our research in this area has been focused on the effects of ligand derivatization and surface water on the complexation and activity of heterogenized Re-bpy.

2.1. Covalent attachment of Re-bpy onto SiO₂ via amide linkages

In our initial study, Re-bpy was physically adsorbed on a hierarchical mesoporous zeolite.⁶⁰ Diffuse reflectance infrared

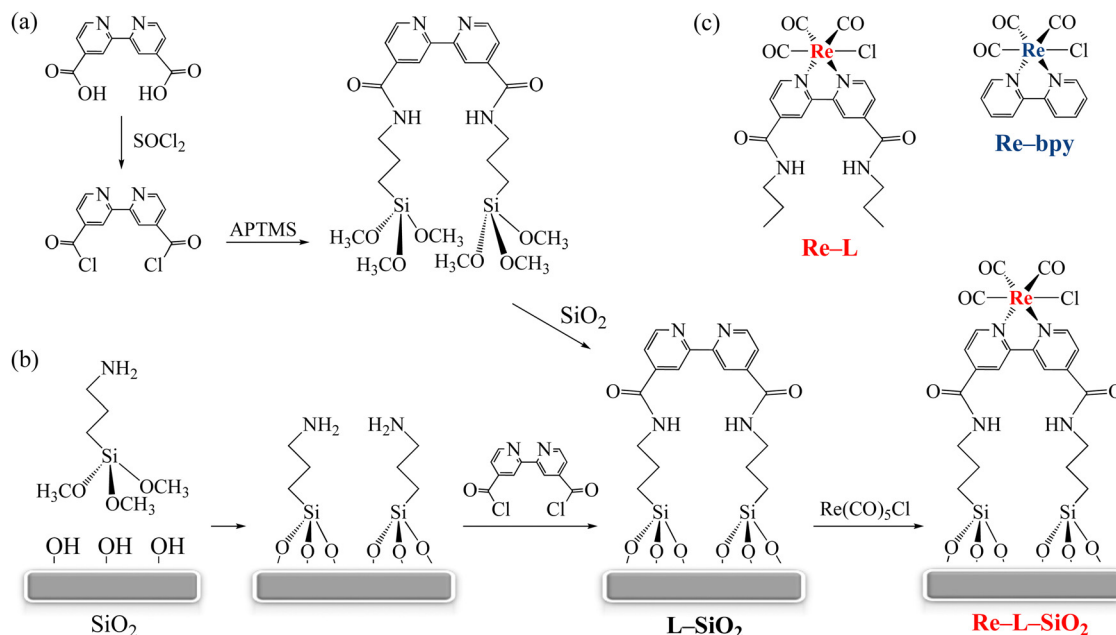
Fourier transform spectroscopy (DRIFTS) was employed to examine the adsorption and complexation of Re-bpy upon light irradiation. In the presence of CO₂ and an electron donor, the formation of important reaction intermediates, including Re-carboxylato and Re-formato species, was observed upon light irradiation.⁶⁰ This initial work allowed us to establish experimental protocols for investigating molecular catalysts on surfaces in the absence of organic solvents.

Covalent attachment of Re-bpy on SiO₂ was achieved *via* two approaches (Scheme 1), both of which involved the derivatization of bpy with chlorocarbonyl groups.^{61,62} In the first approach (Scheme 1a), a bpy-functionalized SiO₂ (denoted “L-SiO₂”) was obtained in the reaction between a SiO₂ surface and a bpy-containing silane coupling agent. In the second approach (Scheme 1b), the SiO₂ surface was functionalized with an amino silane coupling agent prior to reacting with the derivatized bpy ligand. The reaction between L-SiO₂ and Re(CO)₅Cl led to the covalent attachment of Re-bpy on SiO₂ *via* amide linkages.

The heterogenized Re-bpy, denoted “Re-L-SiO₂”, was characterized with DRIFTS to confirm the successful surface immobilization. Fig. 2 shows the DRIFTS spectra of SiO₂, Re-bpy physically adsorbed on SiO₂, L-SiO₂, and Re-L-SiO₂. An intense absorption at 3745 cm^{−1} featuring surface silanol (Si–OH) groups is present in the spectra of SiO₂ and Re-bpy physically adsorbed on SiO₂ (Fig. 2a and b). This peak is absent in the spectrum of L-SiO₂ shown in Fig. 2c since the silanol groups were consumed upon functionalizing the SiO₂ surface with the silane coupling agent. Further supporting this is the presence of infrared absorptions associated with the amide linkages at 1376 cm^{−1} (C–N stretch), 1548 cm^{−1} (N–H bend), and 1657 cm^{−1} (C=O stretch) as well as the methylene C–H stretching bands around 2800–3000 cm^{−1} (not labeled) shown in Fig. 2c. Compared to the spectrum of L-SiO₂, the spectrum of Re-L-SiO₂ contains two additional broad absorptions at 1903 cm^{−1} and 2026 cm^{−1} characteristic of surface Re(i)–carbonyl groups (Fig. 2d), indicating Re-bpy was successfully grafted on SiO₂ through amide linkages.⁶¹

The characteristic carbonyl infrared bands have been utilized to probe the configurations of Re(i) complexes on surfaces. Anfuso and co-workers⁵⁰ shown that a Re(i) complex bearing –COOH groups was bound to TiO₂ through the carboxylate groups in bidentate or tridentate linkage motifs. The surface Re(i) complex had the bpy moiety nearly perpendicular to the TiO₂ surface, exposing the Re(i) complex to the solution in a configuration suitable for catalysis.





Scheme 1 Two synthetic routes for covalently attaching Re-bpy to a SiO₂ surface containing silanol groups (a and b). APTMS: 3-amino-propyltrimethoxysilane. The molecular structures of two homogeneous Re(I) complexes are shown in (c).

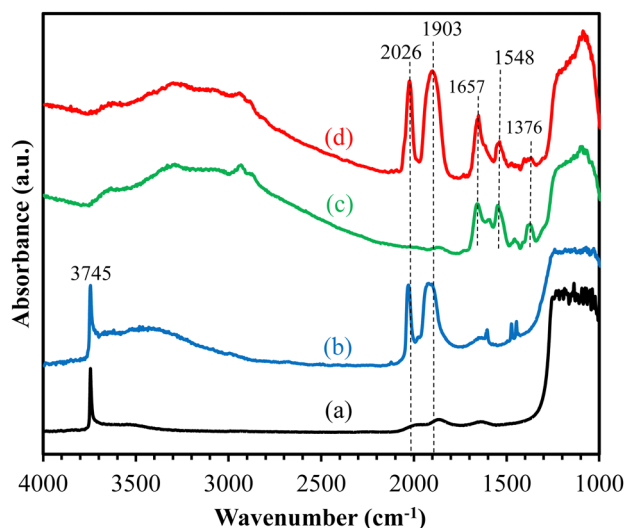


Fig. 2 DRIFTS spectra of (a) SiO₂, (b) Re-bpy physically adsorbed on SiO₂, (c) L-SiO₂, and (d) Re-L-SiO₂. Adapted with permission from *J. Mol. Catal. A*, 2012, **363–364**, 208–213. Copyright 2012 Elsevier.

2.2. Effects of derivatizing the bpy ligand with amide groups

Derivatization of the bpy ligand with amide groups allowed the successful surface immobilization of Re-bpy on SiO₂. However, such ligand derivatization resulted in significant changes in the photophysical and photochemical properties of Re-bpy. For instance, the ultraviolet-visible (UV-vis) spectrum of Re-bpy physically adsorbed on SiO₂ (Fig. 3a, solid trace) features an intense absorption at 365 nm, which is associated with the metal-to-ligand charge transfer (MLCT) transition of Re-bpy on the surface.⁶³ In comparison, the spectrum of the heterogenized

Re(I) catalyst, Re-L-SiO₂, has an MLCT band at 400 nm (Fig. 3b, solid trace).

In order to explore the origin of this observed red shift, a homogeneous compound, Re-L (see Scheme 1c for its structure), was synthesized by derivatizing Re-bpy with amide groups. The MLCT transition of the homogeneous Re-L occurs around 395 nm in contrast with 371 nm for Re-bpy (Fig. 3a and b, dotted traces). Therefore, the red shift observed in Fig. 3(a and b) originated from the ligand derivatization. It is noteworthy that the MLCT bands of surface Re(I) complexes (Fig. 3a and b, solid traces) are much broader than those of the corresponding homogeneous complexes (Fig. 3a and b, dotted traces). This peak broadening is a strong indication of heterogeneity of the surface Re(I) complexes.

We further revealed that the ligand derivatization significantly altered the electrochemical and photocatalytic properties of Re-bpy.⁶² The presence of electron-withdrawing amide groups in Re-L shifted the redox potentials of Re-bpy to more positive values. Consequently, the activity of Re-L was found to be much lower than Re-bpy in photocatalytic CO₂ reduction under simulated solar irradiation (Fig. 3c). Under the same experimental conditions, Re-L-SiO₂ displayed slightly higher activity than Re-L. Similar effects of ligand derivatization were observed in our effort to covalently attach Co-cyclam onto SiO₂, as will be described later in Section 3.1.

Effects of derivatizing Re-bpy with amide groups have been investigated in CO₂-reduction electrocatalysis. Recently, Jia and co-workers⁶⁴ prepared a series of heterogenized Re(I) catalysts on Si electrode surfaces. It was shown that covalent linkages containing the amide groups were not stable under highly reducing conditions, while alkyl linkages were more stable.



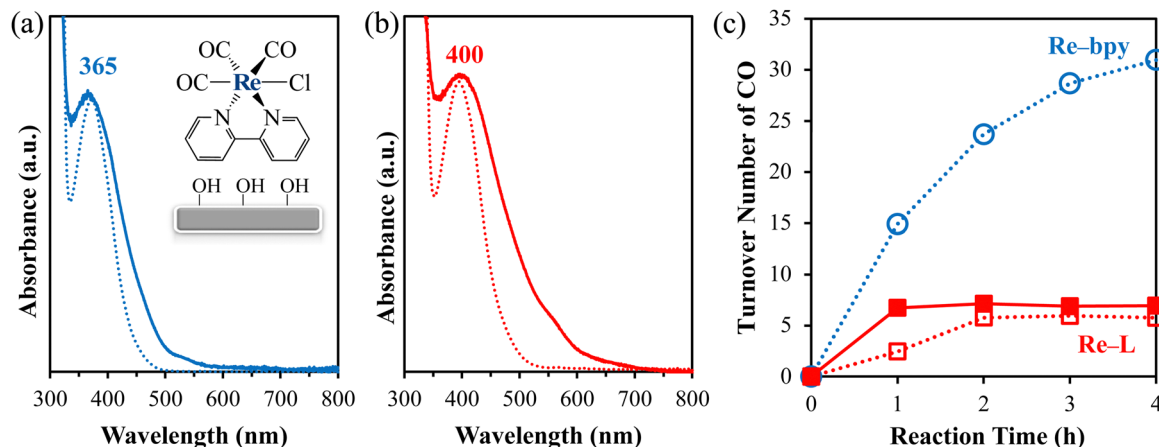


Fig. 3 Diffuse reflectance UV-vis spectra of (a) Re-bpy physically adsorbed on SiO₂ and (b) Re-L-SiO₂. The UV-vis spectra of corresponding molecular complexes (Re-bpy and Re-L) in organic solvents are also shown (dotted traces). Adapted with permission from *J. Mol. Catal. A*, 2012, **363–364**, 208–213. Copyright 2012 Elsevier. (c) Production of CO in solar CO₂ reduction using Re-bpy (open circle), Re-L (open square) and Re-L-SiO₂ (solid square). Adapted with permission from *ACS Catal.*, 2013, **3**, 655–662. Copyright 2013 American Chemical Society.

2.3. Effects of derivatizing the bpy ligand at different positions

Another effect of ligand derivatization was also discovered through studying heterogenized Re-bpy.⁶⁵ In this case, the bpy ligand was derivatized at different positions prior to being coupled with a SiO₂ surface through a dipodal silane coupling agent and being coordinated with Re(CO)₅Cl. In particular, 2,2'-bipyridine-4,4'-dicarboxylic acid and 2,2'-bipyridine-5,5'-dicarboxylic acid were used to prepare surface-immobilized Re(i) catalysts following the synthesis described in Scheme 1b, resulting in the formation of Re-4-SiO₂ and Re-5-SiO₂, respectively (see structures in Fig. 4a). Both Re-4-SiO₂ and Re-5-SiO₂ exhibited a strong MLCT absorption band around 400 nm.⁶⁵

During photocatalysis testing in the presence of triethanolamine (TEOA) as an electron donor, it was observed that the reaction suspension of Re-5-SiO₂ changed color from yellow to green. At room temperature, the green color gradually decayed over a few hours after the light was turned off, and re-appeared when light irradiation resumed.⁶⁵ In addition, the green color immediately changed back to yellow as soon as the reaction

suspension was exposed to air under continuous light irradiation. Interestingly, such color change was not observed for Re-4-SiO₂. Spectroscopic studies indicated that this green color was associated with an absorption around 620 nm (Fig. 4b). We further synthesized corresponding homogeneous complexes by coordinating 2,2'-bipyridine-4,4'-dicarboxylic acid and 2,2'-bipyridine-5,5'-dicarboxylic acid with Re(CO)₅Cl. A similar color change was observed for the Re(i) complex containing 2,2'-bipyridine-5,5'-dicarboxylic acid (Fig. 4c) upon light irradiation. Under the experimental conditions, the Re-bpy complex will be reduced, forming a one-electron reduced species. The green color observed for Re-5-SiO₂ is likely due to the electron density in the reduced Re-bpy being more localized on the 5,5'-derivatized bpy ligand, whereas in reduced Re-4-SiO₂ the electron density is more localized on the metal center.

This explanation was further supported by results obtained using electron paramagnetic resonance (EPR) spectroscopy.⁶⁵ In EPR studies, the heterogenized Re(i) catalysts in powder form was mixed with TEOA and purged with CO₂ prior to collecting EPR spectra. Fig. 4d shows EPR spectra of Re-4-SiO₂ and Re-5-SiO₂

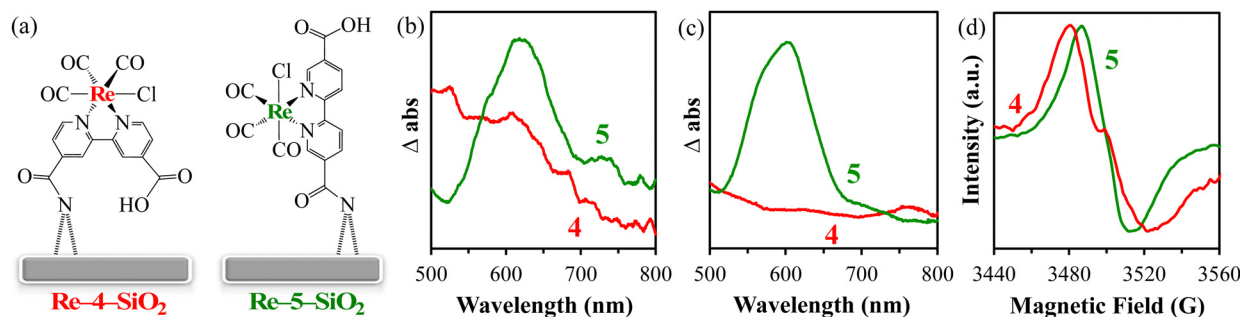


Fig. 4 (a) Structures of Re-4-SiO₂ and Re-5-SiO₂, (b) difference UV-vis spectra of the two heterogenized Re(i) catalysts suspended in dimethylformamide, (c) difference UV-vis spectra of corresponding homogeneous Re(i) complexes in dimethylformamide, and (d) EPR spectra of the two heterogenized Re(i) catalysts in the powder form. Spectra in (b–d) were collected upon light irradiation at room temperature in the presence of TEOA and CO₂. Adapted with permission from *J. Mol. Catal. A*, 2016, **411**, 272–278. Copyright 2016 Elsevier.

collected at room temperature and under continuous light irradiation. The spectrum of Re-4-SiO₂ shows some resonance features originated from hyperfine interactions between unpaired electrons with Re(I) nuclei (Fig. 4d, red trace). Less significant hyperfine coupling features are seen in the spectrum of Re-5-SiO₂ (Fig. 4d, green trace). Therefore, the electron density in reduced Re-5-SiO₂ is more localized on the bpy ligand while in reduced Re-4-SiO₂ the electron density is more localized on the metal center.

Effects of ligand derivatization at different positions were observed in a study by Argazzi and co-workers,⁶⁶ in which Ru(II) polypyridyl complexes were derivatized with -COOH groups for use in dye-sensitized solar cells. It was shown that moving the carboxylate moieties from the 4,4'-positions to the 5,5'-positions lowered the π^* energy of the bpy ligand. Subsequently, the Ru(II) photosensitizer containing 2,2'-bipyridine-5,5'-dicarboxylic acid was less efficient in converting photons into electrons because of more significant nonradiative decay of the excited states of this Ru(II) sensitizer. In a study by Guyot and co-workers,⁶⁷ carboxylic ester groups at the 5,5'-positions of the bpy ligand was found to have a drastic effect on the catalytic activity of Re-bpy toward electrochemical CO₂ reduction since the reducing equivalents are mainly accumulated on the electron-withdrawing ester, preventing the formation of Re(0) species and its interaction with CO₂. In comparison, alkyl-phosphonic ester substituents on the bpy ligand preserved the catalytic activity of Re-bpy. In our study, derivatizing the bpy ligand at different positions led to difference in charge density distribution within the reduced Re(I) complexes. This difference contributed to the higher activity of Re-4-SiO₂ relative to Re-5-SiO₂ in photocatalytic CO₂ reduction.⁶⁵

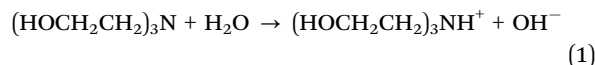
2.4. Effects of surface water

Some surface features could affect the structure, complexation, and behavior of heterogenized molecular catalysts. For example, Lian, Batista, and co-workers⁶⁸ demonstrated that the binding structure of Re-bpy catalysts can be significantly influenced not only by the nature and length of the covalent linkages, but also by the crystallographic facet of the surface. For our heterogenized molecular catalysts, the presence of surface water was found to have profound effects on the complexation of Re-bpy, while the presence of surface -OH groups dedicated the structure and activity of surface Co-cyclam (see Sections 3.1 and 3.2).

The infrared spectrum of Re-bpy in the molecular state contains three carbonyl bands in the 1900–2100 cm⁻¹ range, corresponding to one high-energy, fully symmetric mode and two nearly degenerate lower-energy modes.^{56,69} The two lower-energy modes often coalesce to one broad feature, as seen in the DRIFTS spectrum of Re-L-SiO₂ (Fig. 2d). These two modes can be resolved by exposing the heterogenized Re-bpy to TEOA and CO₂.⁶² For example, three peaks at 2020, 2018, and 1896 cm⁻¹ are present in the DRIFTS spectrum of Re-bpy covalently attached to a non-porous SiO₂ (Fig. 5a). These absorption features are associated with the formation of a CO₂-bound Re(I) adduct, Re(I)-OOC-OCH₂CH₂NR₂ where R is CH₂CH₂OH, in the presence of TEOA and CO₂.⁷⁰ The Re(I)-OOC-OCH₂CH₂NR₂

species was suggested to be the real catalyst in many CO₂-reduction systems using diimine-tricarbonyl Re(I) complexes.

Interestingly, another group of three peaks at 1995, 1876, and 1849 cm⁻¹ are seen in the DRIFTS spectrum of Re-bpy covalently attached to a mesoporous SiO₂ (Fig. 5b). These three peaks are more prominent in the spectrum of Re-bpy covalently attached Kaolin, a hydrated aluminum silicate (Fig. 5c).⁷¹ These peaks are associated with a surface Re(I)-OH species, formed upon replacing the Cl⁻ ligand on Re(I) with -OH, which could be produced by a reaction between TEOA and water (reaction (1)). The comparison shown in Fig. 5 highlights the effect of surface water on the complexation of surface-immobilized Re-bpy. It was suggested that the presence of water in the interlayer space of Kaolin and mesopores of the mesoporous SiO₂ facilitated the formation of the Re(I)-OH species.⁷¹



Interconversion between the surface Re(I)-OOC-OCH₂CH₂NR₂ and Re(I)-OH species was followed by DRIFTS. For example, conversion of Re(I)-OH to Re(I)-OOC-OCH₂CH₂NR₂ was observed under a flow of gaseous CO₂ in the presence of TEOA.⁷¹ Under visible-light irradiation, the three peaks at 2020, 1918, and 1896 cm⁻¹ shifted to lower wavenumbers, while the three peaks at 1995, 1876, and 1849 cm⁻¹ did not shift but their intensities increased over time (Fig. 6a). These observations indicate that the surface Re(I)-OOC-OCH₂CH₂NR₂ species was at least partially converted to the Re(I)-OH species, which is not photoactive and might be associated with the deactivation of the heterogenized Re-bpy. Fig. 6b summarizes possible reactions of different surface Re(I) species based on

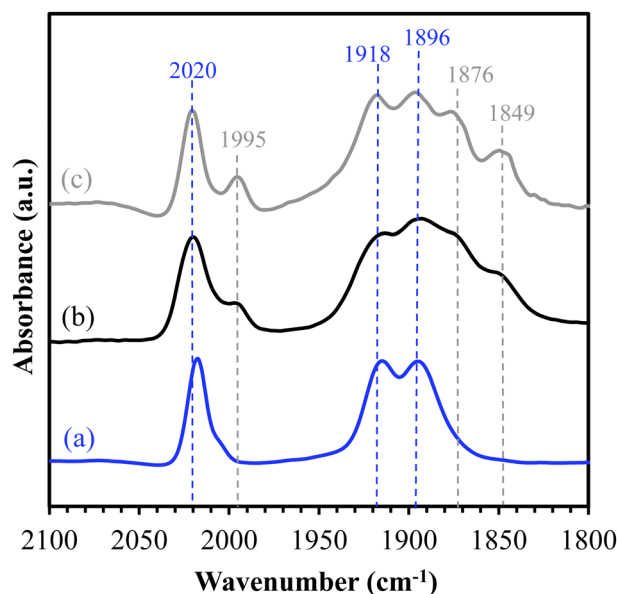


Fig. 5 DRIFTS spectra of Re-bpy covalently attached to (a) a non-porous SiO₂, (b) a mesoporous SiO₂, and (c) Kaolin. The samples in the powder form were mixed with TEOA and exposed to air prior to spectra collection. Adapted with permission from *J. Mol. Catal. A*, 2014, **395**, 145–150. Copyright 2014 Elsevier.



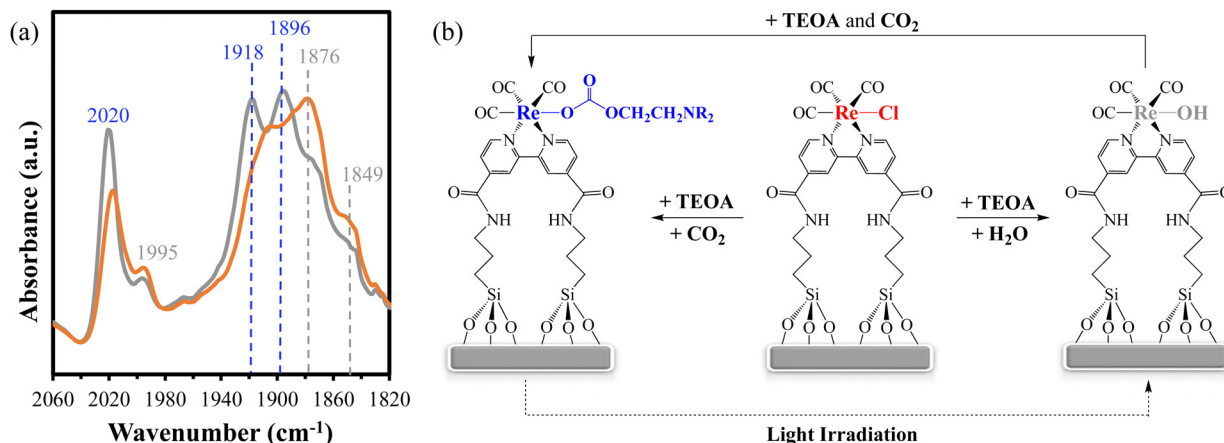


Fig. 6 (a) DRIFTS spectra of Re-bpy covalently attached to Kaolin before (gray) and after (orange) light irradiation in the presence of TEOA. (b) Possible pathways for the formation of different surface Re(I) complexes; R = CH₂CH₂OH. Adapted with permission from *J. Mol. Catal. A*, 2014, **395**, 145–150. Copyright 2014 Elsevier.

DRIFTS studies. Our studies suggest that the presence of surface water has a strong impact on the complexation and even photoactivity of surface Re-bpy.

3. Heterogenization of Co-cyclam for visible-light CO₂ reduction

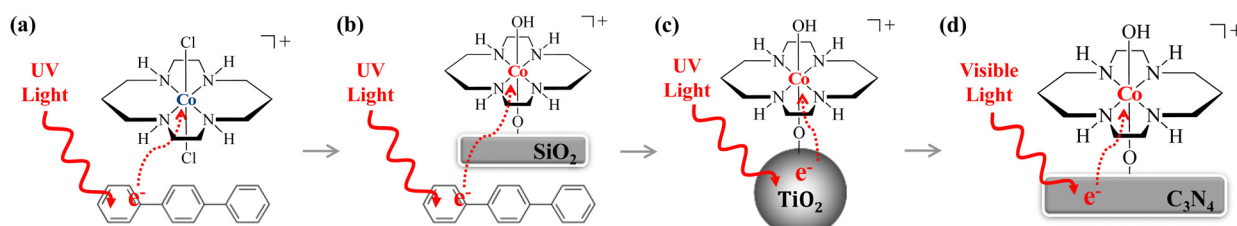
In addition to Re-bpy, other molecular catalysts have been covalently attached onto semiconductor surfaces for use in light-driven CO₂ reduction.^{72–79} Among these heterogenized molecular catalysts, many employed cobalt complexes because of their outstanding performance, low cost, and reasonable stability. Our research in this direction aims at achieving visible-light CO₂ reduction using heterogenized Co-cyclam on semiconductor surfaces. Co-cyclam demonstrated excellent activity in selective CO₂-to-CO conversion in the presence of a molecular photosensitizer, *p*-terphenyl, under UV irradiation (Scheme 2a). A serendipitous finding led to the deposition of Co-cyclam on SiO₂ via an oxo bridge (Scheme 2b). The resulting heterogenized Co-cyclam demonstrated better activity than the molecular Co-cyclam in CO₂ reduction under the same conditions. Co-cyclam was then deposited on TiO₂ (Scheme 2c) and C₃N₄ (Scheme 2d) which replaced *p*-terphenyl as the photosensitizers in CO₂. Our research also revealed the effects of ligand derivatization and surface –OH groups on the structure and activity of heterogenized Co-cyclam. This line of research

further inspired us to synthesize a Co SAC on C₃N₄ which showed excellent activity in visible-light CO₂ reduction (see Section 4).

3.1. Surface immobilization of Co-cyclam on SiO₂

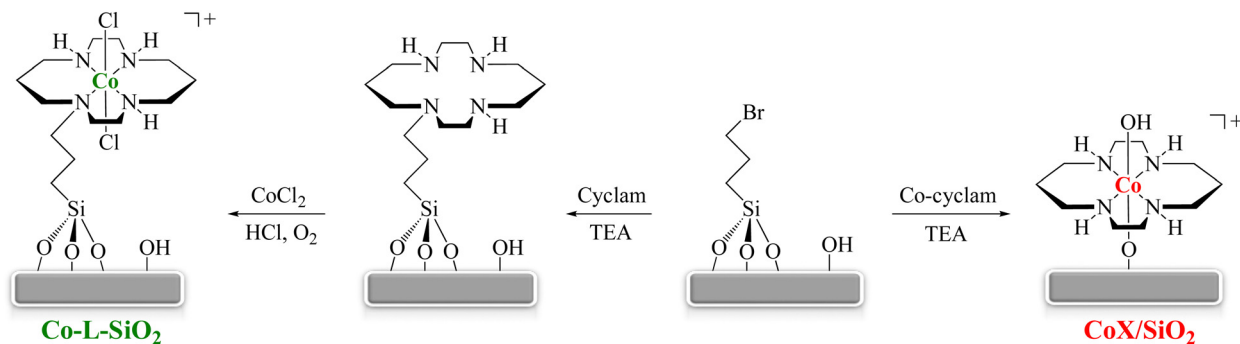
Sujandi and co-workers prepared Co-cyclam covalently attached onto mesoporous SiO₂ through N-functionalization of the cyclam ligand.⁸⁰ The resulting catalyst showed good activity in aerial oxidation of ethylbenzene. In our synthesis of heterogenized Co-cyclam, the SiO₂ surface was initially functionalized with a bromo silane coupling agent (Scheme 3, left). Cyclam-functionalized SiO₂ was then prepared by reacting bromo-functionalized SiO₂ with cyclam in the presence of triethylamine (TEA). Heterogenized Co-cyclam, denoted “Co-L-SiO₂”, was obtained as a green powder after reacting CoCl₂ with cyclam-functionalized SiO₂ in the presence of oxygen.⁸¹ The UV-vis spectrum of Co-L-SiO₂ features a distinct peak around 630 nm (Fig. 7, solid trace), characteristic of the *trans* isomer of Co-cyclam (Fig. 7, dotted trace).⁸²

The synthesized Co-L-SiO₂ was tested in photocatalytic CO₂ reduction using *p*-terphenyl as a molecular photosensitizer. Both CO and H₂ were detected as the major products in the presence of TEOA and methanol as electron donors. Under the same experimental condition, Co-L-SiO₂ demonstrated significantly lower activity and selectivity towards CO formation than Co-cyclam.⁸¹ A turnover number (TON) of 12.5 for CO formation was obtained using Co-L-SiO₂ after CO₂ reduction for



Scheme 2 Our research approach to obtain heterogenized Co-cyclam for visible-light CO₂ reduction.





Scheme 3 Synthetic routes to obtain Co-L-SiO₂ (left) and CoX/SiO₂ (right). TEA: triethylamine.

4 h, compared to 141.8 for Co-cyclam. The selectivity towards CO formation, as measured by the CO/H₂ ratio, was determined to be 7.8 and 15.1 for Co-L-SiO₂ and Co-cyclam, respectively. These results indicate that heterogenization through N-functionalization (Scheme 3, left) was detrimental to the catalytic property of Co-cyclam. In a study by Froehlich and co-workers,⁸³ a similar effect was observed in electrochemical CO₂ reduction using homogeneous Ni(II) cyclam catalysts, where N-functionalization on cyclam with methyl groups drastically altered the redox and catalytic properties of Ni(II) cyclam. Therefore, the observed detrimental effect originated from derivatizing one of the amines on the cyclam ring. Such a detrimental effect could be avoided by derivatizing on one of the C atoms of the cyclam ligand for surface immobilization. Recently, Forget and co-workers⁸⁴ prepared C-functionalized Ni(II) cyclam grafted onto a carbon electrode. The resulting catalyst demonstrated excellent activity in electrochemical CO₂ reduction.

In another attempt to synthesize Co-L-SiO₂, Co-cyclam was mixed directly with bromo-functionalized SiO₂ in the presence of TEA (Scheme 3, right). Interestingly, this reaction resulted in the formation of a different catalyst, denoted “CoX/SiO₂”, in which the Co(III) center is directly deposited on SiO₂ *via* an oxo bridge. Further characterization of CoX/SiO₂ suggested that the chloro ligands in Co-cyclam was replaced with –OH ligands upon surface deposition.⁸¹ Likely, Co-cyclam reacted preferentially with surface silanol (Si–OH) groups, instead of the bromo groups, on the functionalized SiO₂. Based on this discovery, we employed a microwave-assisted synthesis to achieve uniform deposition of Co-cyclam on an unfunctionalized mesoporous SiO₂.⁸⁵ The resulting CoX/SiO₂ demonstrated superior activity in CO₂ reduction using *p*-terphenyl as a photosensitizer. A TON more than 300 was obtained for CO after photocatalytic CO₂ reduction for 4 h (Fig. 8).

In order to further probe the structure of CoX on surfaces, Co-cyclam was deposited *via* the microwave method on a mesoporous SiO₂ with a surface area of 569 m² g^{−1} and a non-porous SiO₂ with a surface area of 200 m² g^{−1}. Under the same synthetic conditions, CoX deposited on the mesoporous SiO₂ (cobalt loading 19 μmol g^{−1}) displayed a green color, corresponding to three absorptions at 600 nm, ~460 nm, and ~380 nm in its UV-vis spectrum (Fig. 9a). In contrast,

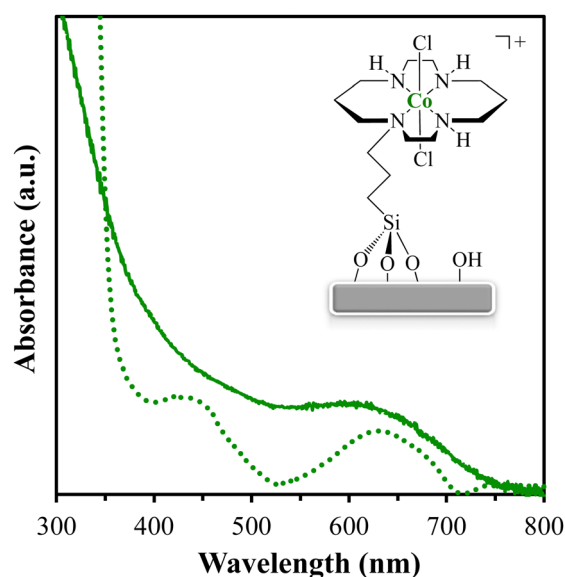


Fig. 7 UV-vis spectra of Co-L-SiO₂ in the powder form (solid trace) and Co-cyclam in H₂O (dotted trace). Adapted with permission from *J. Coord. Chem.*, 2016, **69**, 1748–1758. Copyright 2016 Taylor & Francis Group.

CoX deposited on the nonporous SiO₂ (cobalt loading 13 μmol g^{−1}) had a brown color, and the three peaks are not distinguishable in its UV-vis spectrum due to peak broadening and overlapping (Fig. 9b). Likely, CoX exists on the nonporous SiO₂ as a peroxo Co(III) complex featuring a Co–O₂–Co unit.⁸⁶ In photocatalytic CO₂ reduction using *p*-terphenyl as the photosensitizer, CoX on the mesoporous SiO₂ demonstrated significantly higher activity than on the nonporous SiO₂.

3.2. Deposition of Co-cyclam on TiO₂

Following the same microwave synthesis, CoX was deposited on TiO₂ (Scheme 2c) in order to eliminate the use of molecular photosensitizers such as *p*-terphenyl.⁸⁷ Upon UV light irradiation, photoexcited electrons in TiO₂ nanoparticles were transferred to the surface CoX for CO₂ reduction. On the surface of a commercial TiO₂ material, Evonik P25 consisting of ~80% anatase and ~20% rutile, a TON of 11.2 for CO production was obtained in photocatalytic CO₂ reduction in the presence of TEOA as the electron donor (Fig. 10a). In comparison, no



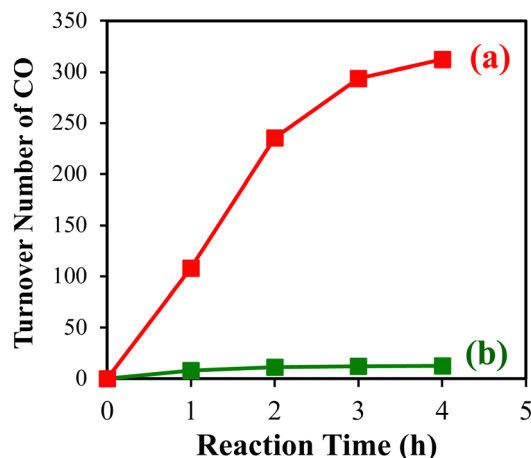


Fig. 8 Photocatalytic CO₂ reduction using (a) CoX/SiO₂ and (b) Co-L-SiO₂. The reaction was carried out using *p*-terphenyl as a photosensitizer and TEOA and methanol as electron donors. Adapted with permission from *J. Coord. Chem.*, 2016, **69**, 1748–1758. Copyright 2016 Taylor & Francis Group.

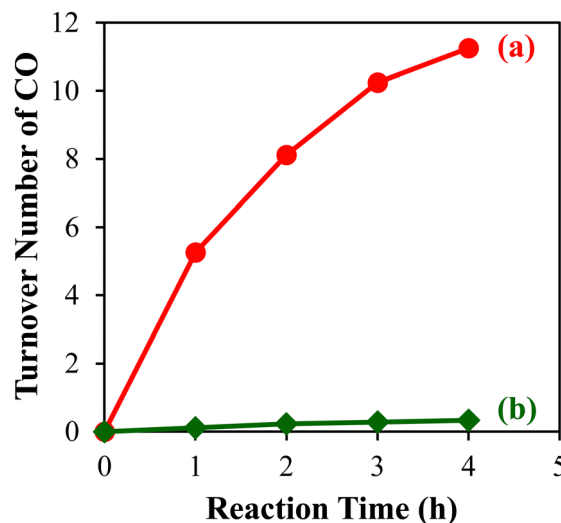


Fig. 10 Photocatalytic CO₂ reduction using (a) CoX/P25 and (b) CoX/Rutile. The reaction was carried out under UV light using TEOA as an electron donor. Adapted with permission from *J. Mol. Catal. A*, 2016, **423**, 293–299. Copyright 2016 Elsevier.

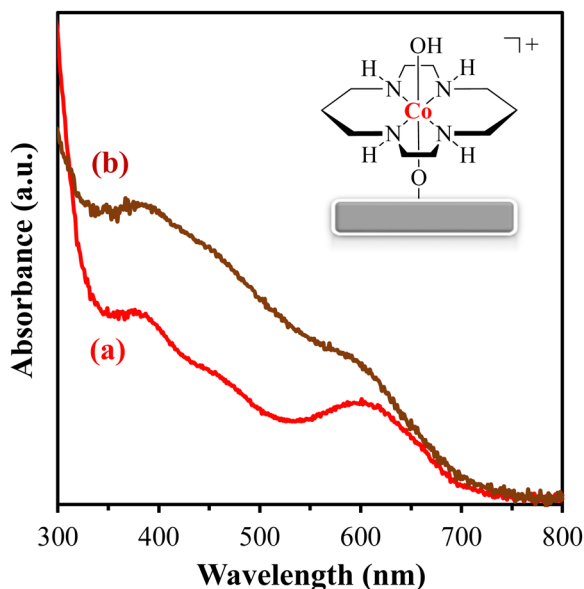


Fig. 9 UV-vis spectra of CoX deposited on (a) a mesoporous SiO₂ and (b) a non-porous SiO₂ via the microwave method. Adapted with permission from *Dalton Trans.*, 2017, **46**, 10721–10726. Copyright 2017 Royal Society of Chemistry.

significant CO production was observed using a suspension containing P25 TiO₂ nanoparticles and Co-cyclam.

The presence of certain surface –OH groups on TiO₂ was found to be essential for the molecular deposition of Co-cyclam.⁸⁸ We prepared two photocatalysts, CoX/P25 and CoX/Rutile, which were prepared by depositing CoX on two commercial TiO₂ materials, P25 and Rutile nanopowder (Sigma-Aldrich), respectively. In photocatalytic CO₂ reduction, CoX/P25 exhibited very good activity, while negligible CO production was observed using CoX/Rutile (Fig. 10). DRIFTS

studies indicated the presence of terminal Ti–OH groups on P25, while the surface of Rutile lacked such –OH groups. In a study by Petsi and co-workers,⁸⁹ surface –OH groups were found to be important for the deposition of [Co(H₂O)₆]²⁺ on TiO₂ nanoparticles. In our study, the terminal –OH groups are likely essential for the molecular deposition of catalytically active CoX on P25 via Ti–O–Co linkages. The lack of sufficient Ti–O–Co linkages in CoX/Rutile led to inefficient photoexcited electron transfer from Rutile to surface CoX, resulting in its poor activity in photocatalytic CO₂ reduction (Fig. 10b).

Additional characterization confirmed that CoX exists in different forms on CoX/P25 than on CoX/Rutile. For example, the UV-vis spectrum of CoX/P25 contains two broad absorptions at ~460 nm and 600 nm (Fig. 11a), indicating the presence of uniform, molecularly deposited CoX.⁸⁸ However, these features are absent in the spectrum of CoX/Rutile (Fig. 11b). Microscopic images showed that amorphous, nano-sized aggregates of CoX are present in CoX/Rutile, but not in CoX/P25.

Based on the specific surface areas of the TiO₂ materials and elemental analysis results, surface catalyst densities were estimated to be 6–7 Co per nm² surface in CoX/P25 and ~26 Co per nm² surface in CoX/Rutile.⁸⁸ These numbers further support our conclusion that CoX is molecularly deposited on P25. The absence of terminal –OH groups for uniform deposition of CoX on Rutile resulted in the formation of CoX in the aggregated form as observed in microscopic images.

Characterization with EPR spectroscopy confirmed the fact that surface cobalt sites exist in different states on Rutile than on P25. Upon deposition on TiO₂ surface in the presence of TEA, some Co(III) centers were reduced to Co(II), which served as a sensitive probe for EPR studies.⁸⁸ As shown in Fig. 12, the spectrum of CoX/P25 under N₂ (Fig. 12a, gray trace) contains resonances corresponding to mononuclear high-spin Co(II) (*S* = 3/2) centers at *g_x* = 5.89, *g_y* = 3.40, and *g_z* = 2.22 (not labeled). The same



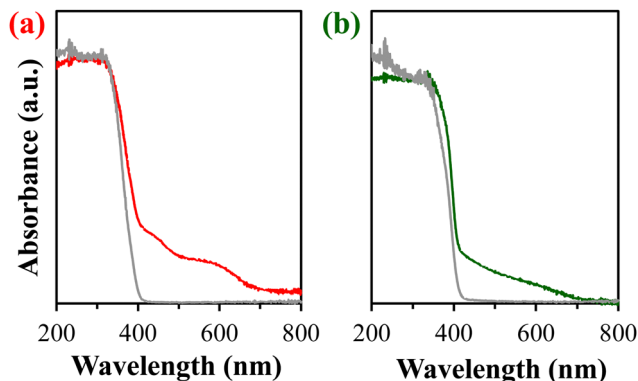


Fig. 11 UV-vis spectra of (a) CoX/P25 and (b) CoX/Rutile. The spectra of bare P25 and Rutile are included as gray traces. Adapted with permission from *J. Mol. Catal. A*, 2016, **423**, 293–299. Copyright 2016 Elsevier.

spectrum also contains resonances near $g = 2$ (not labeled) that are associated with trapped electrons and holes in P25^{90,91} upon exposure to ambient light.

In the spectrum of CoX/Rutile (Fig. 12b, gray trace), however, a very different EPR spectrum was observed in the high-spin region featuring a broad Co(II) peak with $g = 5.35$ due to averaging of g_x and g_y components indicative of Co(II) in a broad range of coordination geometries. In addition, strong eight-line signals associated with oxygen-centered radicals were observed in the region of $g = 2$. Likely, some of the CoX exists as a peroxo complex featuring a Co–O₂–Co unit on Rutile, similar to that on the nonporous SiO₂ (Fig. 9).

Upon CO₂ adsorption, the Co(II) sites in CoX/P25 displayed a high-spin signal at $g_x = 5.67$ (Fig. 12a), indicating that the adsorbed CO₂ on CoX/P25 took electron density away from the Co(II) centers. On CoX/Rutile (Fig. 12b), CO₂ adsorption did not

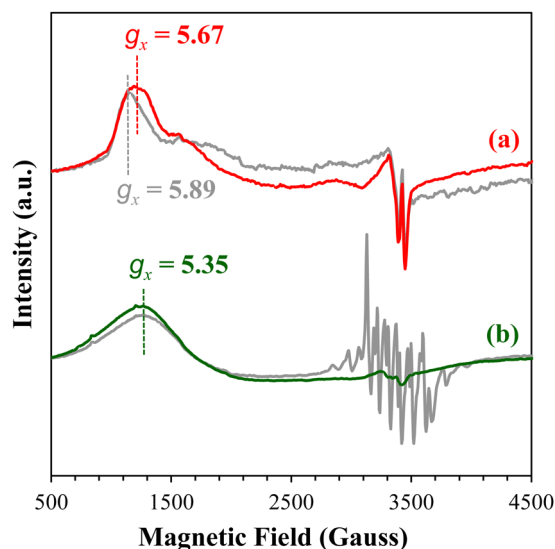


Fig. 12 X-band EPR spectra of (a) CoX/P25 and (b) CoX/Rutile under CO₂. Corresponding spectra under N₂ are also included (gray traces). The spectra were collected at liquid He temperature after purging the samples under N₂ or CO₂ at room temperature. Adapted with permission from *J. Mol. Catal. A*, 2016, **423**, 293–299. Copyright 2016 Elsevier.

shift the high-spin Co(II) signal at $g_x = 5.35$. Instead, the hyperfine splitting of the oxygen radicals vanished and an anisotropic signal appeared near $g = 2$ (not labeled). This observation implies that the electron density on the oxygen radicals adjacent to Co(II) was preferentially transferred to CO₂ and adsorption of CO₂, in fact, removed electron exchange of Co(II) with oxygen atoms on the Rutile surface. Based on these observations, it can be concluded that the Co(II) sites in CoX/Rutile are in a broad range of coordination geometries different than those in CoX/P25.

3.3. Deposition of Co-cyclam on C₃N₄

An inorganic polymeric semiconductor, C₃N₄ has attracted enormous interest due to its low cost, high thermal and chemical stability, nitrogen-rich structure, and suitable band-gap (2.7 eV).^{92–94} In CO₂ reduction, C₃N₄-based materials have been employed as photocatalysts^{95–97} or coupled with metal-ligand complexes^{76–79,98–104} to achieve enhanced catalysis. For example, Roy and co-workers⁷⁶ deposited cobalt phthalocyanine cyanine on mesoporous C₃N₄ via a weak π – π stacking interaction for visible-light CO₂ reduction. Wei and co-workers⁷⁹ attached a cobalt quaterpyridine complex onto C₃N₄ through an amide linkage, which showed remarkable selectivity (98%) towards CO production under visible-light irradiation. In the recent work by Shang and co-workers,¹⁰⁴ a cobalt phthalocyanine complex was deposited on C₃N₄ for photocatalytic CO₂ reduction. The cobalt phthalocyanine complex contains –COOH groups, which enabled high surface coverage of the cobalt complex and promoted its catalytic property.

In our study, Co-cyclam was deposited on C₃N₄ (Scheme 2d) in the presence of TEA, forming CoX/C₃N₄ as indicated by a broad peak around ~ 600 nm in its UV-vis spectrum (Fig. 13). A TON of more than 160 for CO production was obtained using Co-cyclam deposited on a modified C₃N₄ after CO₂ reduction for an extended period under visible-light irradiation ($\lambda > 420$ nm).¹⁰⁵ Under the same reaction condition, no CO was produced using a physical mixture of Co-cyclam and C₃N₄.

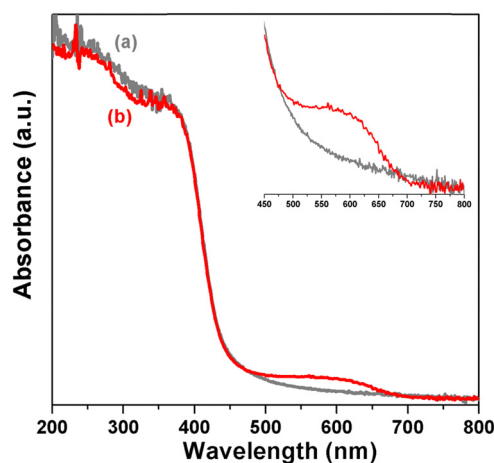


Fig. 13 UV-vis spectra of (a) bare C₃N₄ and (b) CoX/C₃N₄. Insert: A close view of the spectra.



In combination with computational modeling, X-ray absorption spectroscopy (XAS) was employed to investigate the structure of CoX on C_3N_4 .^{105,106} For metal-based catalytic materials, the extended X-ray absorption fine structure (EXAFS) analysis can provide valuable insights into the local geometric structure and coordination environment of the metal sites,¹⁰⁷ while the X-ray absorption near edge structure (XANES) spectroscopy is particularly sensitive to the electronic structure of the metal center and its local environment.¹⁰⁸ As can be seen in Fig. 14a, the Co K-edge XANES spectrum of CoX/ C_3N_4 has different features than that of Co-cyclam, confirming that the coordination environment of cobalt significantly changed upon deposition on C_3N_4 . Based on prior studies,^{87,88,101} we hypothesize that the two axial Cl ligands of Co-cyclam were replaced with one OH group and one surface -O- or -NH- group upon deposition on C_3N_4 in the presence of TEA.

Our hypothesis was supported by the Co K-edge EXAFS spectra of Co-cyclam and CoX/ C_3N_4 (Fig. 14b), which clearly indicate the absence of Co-Cl bonds in CoX/ C_3N_4 . Fitting the first peak of the EXAFS spectrum of Co-cyclam led to coordination numbers of 3.5 ± 1.2 and 2.0 ± 1.2 for Co-N and Co-Cl bonds, respectively. The Co-O scattering path was chosen to fit the first peak of CoX/ C_3N_4 , resulting in a coordination number of 6.3 ± 1.4 for Co-O nearest neighbors.¹⁰⁵ Since the bond lengths of Co-N and Co-O are very close to each other, our experimental results were not able to distinguish between contributions from N and O atoms. We further carried out density functional theory (DFT) simulations, in which anchoring of CoX to C_3N_4 through either a surface -O- or -NH- group represents the most probable modes. Similar energies and distances between the Co center of CoX and C_3N_4 were obtained for these two modes. Therefore, it remains unclear whether CoX attaches to C_3N_4 through the -O- or -NH- group. Nevertheless, the structure shown in Scheme 2d represents a plausible model of CoX/ C_3N_4 . Fig. 14c shows a DFT optimized structure of CoX attached to C_3N_4 through an oxygen atom.

A similar macrocyclic complex, [Co(HMD)Cl₂]Cl where HMD is 5,7,7,12,14,14-hexamethyl-1,4,8,11-tetraazacyclotetradeca-4,11-diene, is also known for its catalytic activity in selective CO₂ reduction.¹⁰⁹ We have deposited this cobalt complex onto different semiconductor surfaces (TiO₂, N-Ta₂O₅, and C_3N_4).

The resulting heterogenized catalyst on TiO₂ demonstrated lower activity than CoX/TiO₂ in CO₂ reduction under UV irradiation, while on C_3N_4 it showed significantly higher activity than CoX/ C_3N_4 under visible-light irradiation.¹⁰¹

4. Single atom catalysts in C_3N_4 for visible-light CO₂ reduction

Crystalline porous materials, particularly metal-organic frameworks (MOFs), are excellent support for constructing well-defined catalytic sites.^{110–112} A variety of molecular catalysts, including Re-bpy, have been incorporated in MOFs with demonstrated activity towards photocatalytic CO₂ reduction.^{113–116} Other examples of such catalysts in porous materials include metal-to-metal charge transfer units^{117–119} and single atom catalysts (SACs).^{120–123} Porous materials represent an ideal platform for studying effects of chemical microenvironment on the activity of well-defined catalytic sites.^{124–126}

SACs have been extensively explored in a variety of applications including photocatalytic^{44–48} and electrocatalytic CO₂ reduction.^{127–129} Two-dimensional materials, including N-doped graphene and C_3N_4 , have been employed as the support for SACs.^{130–132} The N-rich framework of C_3N_4 enables it to serve as a photoactive support for SACs featuring metal (M)-N coordination.^{133–135} In 2018, we reported a Co SAC on C_3N_4 that demonstrated interesting activity in selective CO₂ reduction under visible-light irradiation.¹³⁶ Since then, many studies have examined SACs on C_3N_4 for solar CO₂ reduction.^{137–151} For example, Wang and co-workers¹⁴⁹ synthesized an Ni SAC anchored by boron-oxo species on C_3N_4 nanosheets. The Ni SAC was shown to possess a unique Ni-O₆ coordination and demonstrated excellent activity towards CO and CH₄ production in photocatalytic CO₂ reduction. Sun and co-workers¹⁵⁰ prepared two Cu SACs with different Cu configurations (Cu₁N₃ and Cu₁P₃) on phosphorus-doped C_3N_4 . In visible-light CO₂ reduction in the presence of H₂O, the Cu SAC containing the Cu₁N₃ unit was found to be active towards CO production, while the Cu SAC containing the Cu₁P₃ unit preferentially produced H₂ instead of CO. Ou and co-workers¹⁵¹ synthesized a SAC with two active centers, Mn and Co, on C_3N_4 .

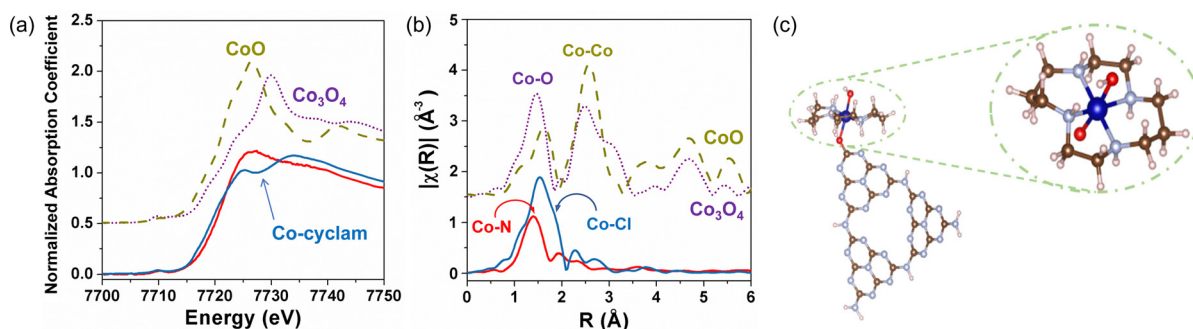


Fig. 14 (a) XANES and (b) EXAFS spectra of Co-cyclam (blue), CoX/ C_3N_4 (red), and CoO and Co_3O_4 as references. (c) An optimized structure of CoX attached to C_3N_4 through an oxygen atom. Color scheme: Co (blue), N (silver), C (brown), O (red), and H (pink). Adapted with permission from *J. Phys. Chem. C*, 2023, **127**, 3626–3633. Copyright 2023 American Chemical Society.



Experimental results and DFT calculations showed an interesting synergy in photocatalysis using the Mn–Co SAC, where H₂O oxidation occurred on the Mn center and CO₂ reduction occurred on the Co center.

In 2016, Gao and co-workers¹⁵² predicted that efficient visible-light CO₂ reduction can be achieved using single Pd and Pt atoms supported on C₃N₄. Inspired by this report and other prior work on SACs, we prepared a Co SAC on C₃N₄ (Fig. 15a) by mixing CoCl₂ with C₃N₄ in acetonitrile, followed by microwave heating in the presence of TEA.¹³⁶ The typical SAC behavior was observed in photocatalytic CO₂ reduction using the synthesized material (Fig. 15b). At relatively low cobalt loadings, the presence of single Co²⁺ sites was confirmed with XAS (Fig. 15c, red trace). Catalytically inactive cobalt oxide nanoparticles were produced at high cobalt loadings (Fig. 15c, blue trace).

Different structural modes have been proposed for SACs on C₃N₄, including in-plane M–N₃ and inter-layer M–N₄ coordination,¹⁵³ C–M–N₂ coordination,¹³⁷ M–N₄ coordination in the presence of N vacancies,¹⁵⁴ and other possibilities.^{155,156} Results from fitting the EXAFS spectrum shown in Fig. 15c indicated that our Co SAC has four equivalent Co–N bonds. However, computational modeling failed to reach such a Co–N₄ structure inside the “pocket” of pristine C₃N₄ (Fig. 16a and b). Instead, a Co–N₂₊₂ coordination at C₃N₄ edge sites appeared to be the most probable structure (Fig. 16c).¹⁵⁷ For this structure, the calculated Co–N bond length is in the range of 2.07–2.10 Å, in excellent agreement with the value ($d_{\text{Co–N}}$ 2.08 Å) obtained by fitting the EXAFS spectrum. This mode of coordination was further supported by our experimental results obtained using a

Co SAC in C₃N₄ treated with NH₃, which contains more edge sites than untreated C₃N₄ and subsequently allows a significantly higher loading of single Co²⁺ sites. Similar SAC structures have been reported, such as the Fe–N₂₊₂ sites of Fe SACs confined in N-doped carbon materials.^{129,158,159}

Doping C₃N₄ with an appropriate amount of carbon significantly enhanced the photocatalytic activity of the Co SAC on C₃N₄.¹⁶⁰ In our study, carbon doping was achieved by adding a small amount of dextrose to urea (5–400 mg dextrose per 20 g urea) for synthesizing C₃N₄ *via* pyrolysis. Fig. 17a shows the pictures of synthesized C₃N₄ containing varying amounts of carbon dopant. While pristine C₃N₄ synthesized from pure urea displays a light yellow color, C-doped C₃N₄ becomes darker in color with increasing amounts of dextrose used in synthesis. These C₃N₄ materials were used to prepare Co SACs following our established method, which was then tested in photocatalytic CO₂ reduction to probe the effect of carbon doping. As can be seen from Fig. 17b, an optimal level of carbon doping exists since the highest activity was obtained using a Co SAC on a C-doped C₃N₄ material synthesized from 20 mg dextrose and 20 g urea. In subsequent studies, this particular C-doped C₃N₄ was utilized to prepare Co SACs.

At a cobalt loading $\sim 0.014 \mu\text{mol mg}^{-1}$, TONs of 4.4 and 61.6 for CO production were obtained using a Co SAC on pristine C₃N₄ (denoted “Co²⁺@C₃N₄”) and a Co SAC on C-doped C₃N₄ (denoted “Co²⁺@C-C₃N₄”), respectively (Fig. 18a).¹⁵⁷ The XANES spectrum of Co²⁺@C₃N₄ is almost identical to that of Co²⁺@C-C₃N₄ (Fig. 18b), suggesting that the two Co SACs have similar electronic and geometric structures. Additional characterization was done to explore possible reasons for the

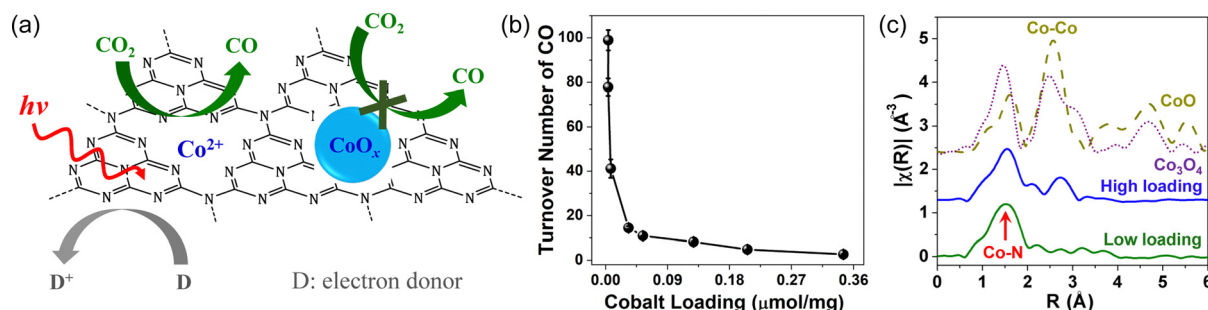


Fig. 15 (a) Schematic of photocatalytic CO₂ reduction mediated by a Co SAC on C₃N₄; (b) TONs of CO after CO₂ reduction for 2 h as a function of cobalt loading; (c) EXAFS spectra of Co²⁺ on C₃N₄ at low and high loadings, and CoO and Co₃O₄ as references. Adapted with permission from *J. Am. Chem. Soc.*, 2018, **140**, 16042–16047. Copyright 2018 American Chemical Society.

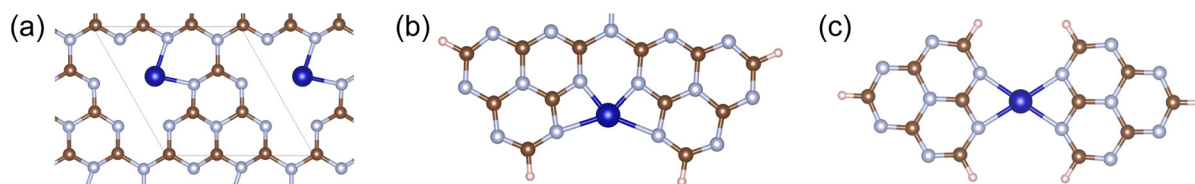


Fig. 16 Structure models of single Co²⁺ sites coordinated with N atoms in C₃N₄: (a) Co–N₂, (b) Co–N₄, and (c) Co–N₂₊₂ at edge sites. A $4 \times 4 \times 1$ *k*-point mesh and a force convergence of 10^{-5} eV Å⁻¹ was used in modeling studies. Color scheme: Co (blue), N (silver), C (gold), and H (light brown). Adapted with permission from *J. Phys. Chem. C*, 2022, **126**, 8596–8604. Copyright 2022 American Chemical Society.



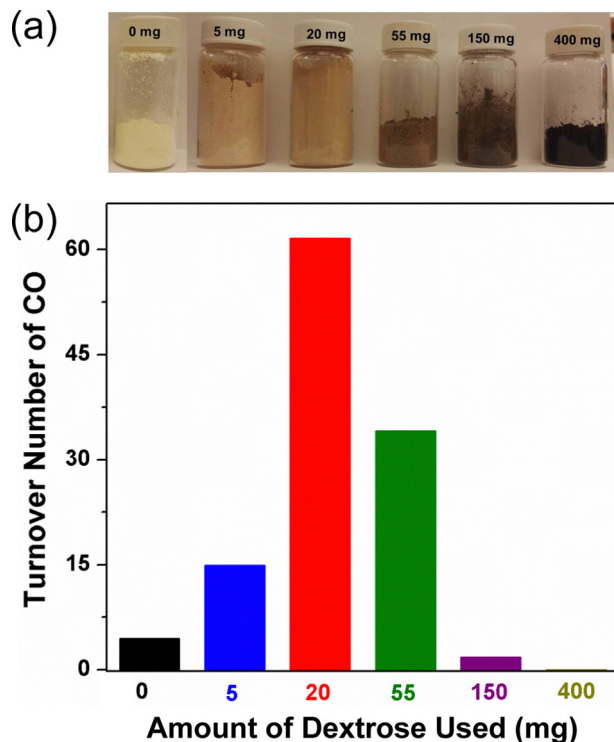


Fig. 17 (a) Pictures of C₃N₄ synthesized in the presence of different amounts of dextrose. (b) TONs for CO production after photocatalytic CO₂ reduction for 2 h using Co SACs on different C-doped C₃N₄. Adapted with permission from *ChemNanoMat*, 2021, **7**, 1051–1056. Copyright 2021 Asian Chemical Editorial Society.

observed enhancement in photocatalysis. The synthesized C-C₃N₄ has greater absorption than pristine C₃N₄ in the visible-light region (400–800 nm, Fig. 18c). The different photoluminescence spectra of C-C₃N₄ and C₃N₄ (Fig. 18d) indicate that doping C₃N₄ with carbon significantly inhibited photo-induced charge recombination. It has been reported that appropriate carbon doping can enhance the photocatalytic properties of C₃N₄ materials by improving its photoresponse and inhibiting charge recombination.¹⁶¹ These factors should account for, at least partly, the observed higher activity of Co²⁺@C-C₃N₄ than Co²⁺@C₃N₄ in our study.

Our experimental results suggest that other factors also contributed to the enhanced activity of Co²⁺@C-C₃N₄ in photocatalytic CO₂ reduction. In particular, Co-cyclam deposited on C-C₃N₄ displayed a 2-fold enhancement in activity in comparison with Co-cyclam deposited on C₃N₄ (CoX/C₃N₄ in Section 3.3).¹⁵⁷ Under the same experimental condition, Co²⁺@C-C₃N₄ demonstrated ~14 times higher activity than Co²⁺@C₃N₄ in CO₂ reduction. Therefore, C doping likely also affected the coordination environment of Co²⁺ sites in C-C₃N₄. In our study, the Co-N₂₊₂ coordination at C₃N₄ edge sites is also the most probable structure for Co²⁺@C-C₃N₄. A simplified model of C doping was proposed to explore possible additional effects of C doping. In this model, the presence of C dopant near the cobalt centers (Fig. 19, circled in red) results in shorter Co-N bond length and stronger Co-N binding energy. In addition to enhanced light absorption and charge separation in C-doped C₃N₄, the stronger Co-N binding upon C doping likely contributes to the improved catalytic properties of Co²⁺@C-C₃N₄.

It is much desired to obtain unified structure–property relationships for different catalytic materials.¹⁶² In our research, the Co-N₄ unit is crucial for the CO₂-reduction activity of both the heterogenized Co-cyclam and the Co SAC on C₃N₄. The calculated Co-N bond length in Co²⁺@C-C₃N₄ (1.96–1.97 Å) is very close to that in Co-cyclam (1.97–2.02 Å).¹⁵⁷ It is possible that insights learned from studying such SACs could serve as inspiration for the design and development of innovative molecular catalysts, and *vice versa*.

5. Summary and outlook

Heterogenized molecular catalysts and SACs contain well-defined surface catalytic sites. They have demonstrated exciting activities in various applications, including solar CO₂ reduction. Our recent research has focused on (i) covalently attaching two well-known molecular catalysts, Re-bpy and Co-cyclam, on different surfaces, and (ii) understanding the structures of Co SACs on C₃N₄. We have utilized catalyst synthesis, spectroscopic characterization, and computation modeling to interrogate the structures of well-defined surface catalytic sites and understand factors affecting their structures and photocatalytic activities.

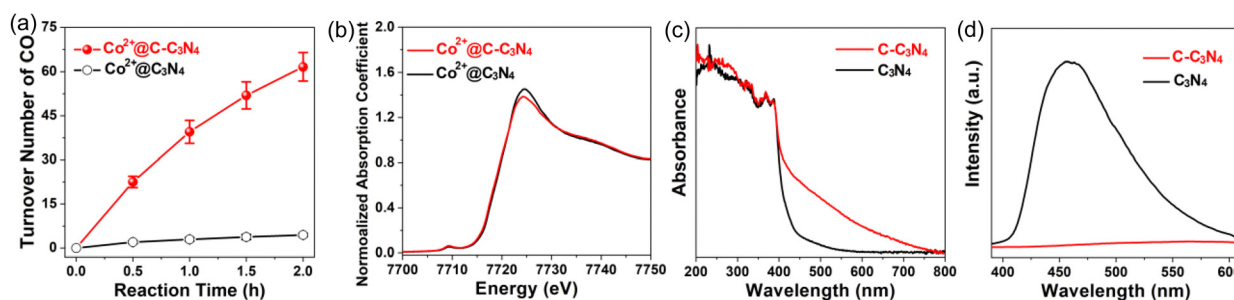


Fig. 18 Pristine C₃N₄ versus C-doped C₃N₄: (a) CO production in photocatalytic CO₂ reduction using Co SACs, (b) normalized Co K-edge XANES spectra of Co SACs and reference samples (CoO and Co₃O₄); (c) optical absorption, and (d) photoluminescence spectra. Adapted with permission from *J. Phys. Chem. C*, 2022, **126**, 8596–8604. Copyright 2022 American Chemical Society.



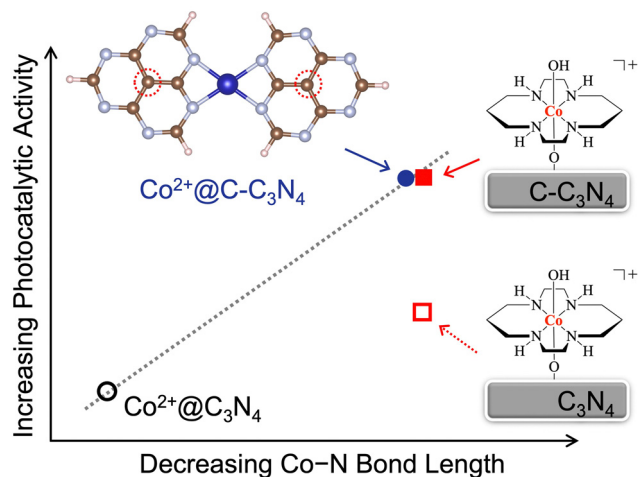


Fig. 19 A proposed correlation between observed photocatalytic activity and calculated Co–N bond length of $\text{Co}^{2+}@\text{C}_3\text{N}_4$, $\text{Co}^{2+}@\text{C-C}_3\text{N}_4$, $\text{CoX}/\text{C}_3\text{N}_4$, and $\text{CoX}/\text{C-C}_3\text{N}_4$. In the model of $\text{Co}^{2+}@\text{C-C}_3\text{N}_4$, two of the framework N atoms are replaced with C atoms (circled in red). Color scheme: Co (blue), N (silver), C (gold), and H (light brown).

In particular, Re-bpy was covalently grafted onto SiO_2 surfaces *via* amide linkages. Derivatization of the bpy ligand with electron-withdrawing amide groups significantly lowered the activity of Re-bpy in photocatalytic CO_2 reduction. Such derivatization at different positions of the bpy ligand resulted in electron density being localized in different parts of Re-bpy upon photoreduction and subsequently different photocatalytic activity. The presence of surface water was found to have a strong impact on the complexation and even photoactivity of surface Re-bpy.

Similar effects of ligand derivatization and surface characteristics were observed in heterogenization of Co-cyclam on different surfaces. In the presence of *p*-terphenyl as a molecular photosensitizer, Co-cyclam grafted onto SiO_2 *via* one of the coordinating N atoms showed poor activity towards CO_2 reduction. Under the same experimental conditions, Co-cyclam directly deposited on SiO_2 *via* an oxo bridge on the Co center demonstrated excellent activity. This oxo bridge strategy was employed to deposit Co-cyclam on TiO_2 , which functioned as a light absorber in photocatalysis, eliminating the need for *p*-terphenyl as a molecular photosensitizer. The presence of surface OH groups on SiO_2 and TiO_2 was found to be key for the effective surface deposition of Co-cyclam. Building on these results, visible-light CO_2 reduction was achieved using Co-cyclam deposited C_3N_4 , which has a narrower bandgap than TiO_2 and can be activated under visible-light irradiation.

Inspired by the activity of Co-cyclam on C_3N_4 and prior work on SACs by others, we discovered a Co SAC featuring Co^{2+} coordinated to framework N atoms on C_3N_4 . The Co SAC demonstrated excellent activity in visible-light CO_2 reduction. Based on spectroscopic and modeling studies, a Co-N_{2+2} coordination at C_3N_4 edge sites was suggested to be the most probable structure. Doping C_3N_4 with an appropriate amount of carbon significantly enhanced the photocatalytic activity of

the Co SAC. A model of carbon doping was also proposed, which involves the substitution of N atoms near the Co centers with C atoms.

In our research, spectroscopic techniques, including infrared, UV-visible, electron paramagnetic resonance, and X-ray absorption spectroscopies, were extensively utilized to understand the structures and properties of heterogenized molecular catalysts and SACs. Computational modeling further provided important insights regarding the structures of these well-defined surface catalytic sites at the molecular level. Our research has been largely inspired by exciting research results from other groups, some of which are briefly discussed in this article. Needless to say, many questions remain unanswered as we continue to investigate these photocatalytic materials.

Despite the tremendous progress made so far, major challenges and ample opportunities co-exist in this field, calling for collaborative efforts by experts from different disciplines. Many photocatalysts, especially those involving molecular photosensitizers, require the use of non-renewable, sacrificial electron donors in CO_2 reduction. A sustainable photocatalytic system should use water as the sole electron donor. In this sense, photoelectrochemical synthesis, which often employ heterogenized molecular CO_2 -reduction catalysts on semiconductors with appropriate band structures, appear to be the ultimate solution to address this issue and huge progress has been made in this direction.^{163,164} Innovative design of SACs has allowed effective coupling between CO_2 reduction and water oxidation.¹⁵¹ The development of photocatalysts based on earth-abundant metals¹⁶⁵ is also important for achieving energy sustainability through solar CO_2 reduction.

In some instances, heterogenization of molecular catalysts could lead to reduced catalytic activity due to detrimental ligand derivatization, in spite of enhanced stability upon surface attachment. This was observed in our studies using Re-bpy and Co-cyclam, as described in Sections 2 and 3. Therefore, effective heterogenization strategies should aim to preserve or even improve the catalytic activity of the metal centers. Research in this area could really benefit from existing knowledge on the effects of ligand derivatization on the activity of molecular catalysts.

It remains a challenge to reveal the exact structures of well-defined surface catalytic sites at the molecular level. Our research demonstrated that combining catalyst synthesis with advanced spectroscopy and theory is a powerful approach to study heterogenized molecular catalysts and SACs. Such an approach could generate critical information beyond just catalyst structures. For example, Lian, Batista, and co-workers^{50,166,167} employed phase-sensitive sum frequency generation spectroscopy and DFT calculations to interrogate the configuration of Re-bpy on TiO_2 surfaces. The researchers investigated Re-bpy attached to TiO_2 *via* molecular linkages of various lengths, which could allow effective manipulation of the spatial orientation of Re-bpy on TiO_2 surfaces.

Furthermore, elucidating the unique advantages of heterogenized molecular catalysts and SACs in photocatalysis requires mechanistic studies using ultrafast spectroscopies and *in situ*/



operando techniques.^{168–171} As mentioned earlier, results obtained using transient absorption spectroscopy revealed underlying reasons for observed enhancement in the CO₂-reduction activity of Re-bpy upon heterogenization on TiO₂ surfaces.^{52,53} In this sense, one advantage of photocatalysts containing well-defined surface sites lies in the fact that they could utilize techniques from both homogeneous catalysis and heterogeneous catalysis.

Conflicts of interest

There are no conflicts to declare.

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