



Cite this: *Energy Environ. Sci.*, 2024, 17, 2260

Molecular modulation of nickel–salophen organic frameworks enables the selective photoreduction of CO₂ at varying concentrations†

Xiaohan Yu,‡^{ab} Mingzi Sun, ‡^c Tianran Yan, ^{ab} Lin Jia, ^{ab} Mingyu Chu, ^{ab} Liang Zhang, ^{ab} Wei Huang, *^{ab} Bolong Huang *^{ce} and Yanguang Li *^{abd}

Photocatalytic CO₂ reduction to value-added chemicals is appealing but challenging, especially under dilute CO₂ conditions. Herein, we present a molecular modulation strategy for porous metal–salophen organic frameworks (M-SOFs), involving cooperative regulation of the catalytically active metal centers and their local coordination environments for selective photocatalytic CO₂ reduction across a wide range of CO₂ concentrations. The optimal Ni-SOF shows a remarkable photocatalytic CO production rate of 16 908 μmol h⁻¹ g⁻¹ and near-unity selectivity under a pure CO₂ atmosphere, along with excellent structural stability. More impressively, it largely preserves the catalytic activity and selectivity even when exposed to dilute CO₂ (5–20 vol%). Both experimental and theoretical analyses support that the specific Ni–N₂O₂ coordination environment in the Ni-SOF endows it with strong CO₂ binding capacity. This, coupled with nanoporous skeletons, enhances local CO₂ enrichment and facilitates its subsequent conversion at the catalytic centers, thereby leading to superior photocatalytic performances at various CO₂ concentrations.

Received 30th November 2023,
Accepted 5th February 2024

DOI: 10.1039/d3ee04121b

rsc.li/ees

Broader context

Solar-driven CO₂ conversion has been considered as an ideal approach for reducing atmospheric CO₂ concentration while producing value-added chemicals and fuels. Over the past decades, tremendous progress has been made in the development of high-performance photocatalysts. Unfortunately, most of them are only operable under high-purity CO₂ conditions. Direct photoreduction of dilute CO₂ is appealing, given that industrial exhaust gases typically contain low concentrations of CO₂; however, it remains a formidable challenge. The key obstacles lie in the inadequate CO₂ capturing ability of the photocatalysts and unfavourable interactions between catalytic sites and CO₂ molecules. Here, we demonstrate the unique capability of metal–salophen organic frameworks – with properly engineered metal centers and coordination environments – to facilitate the local enrichment and subsequent conversion of CO₂. The optimal sample shows exceptional photocatalytic performance and near-unity CO₂-to-CO selectivity across a wide range of CO₂ concentrations.

Introduction

Excessive CO₂ emissions from fossil fuel combustion have been recognized as the dominant cause of global warming and climate change, posing a significant threat to our economic and environmental sustainability.¹ Several strategies have been proposed to reduce the atmospheric CO₂ concentration, either through physical sequestration or chemical conversion.^{2,3} Among them, solar-driven CO₂ reduction to valuable chemicals (e.g., CO, CH₄, CH₃OH, and HCOOH) offers a particularly attractive approach by transforming solar energy into chemical energy while neutralizing CO₂ emissions.^{4,5} Nevertheless, current investigations are primarily based on pure CO₂ feedstocks. Given that a major source of CO₂ emissions is industrial exhaust gases which contain relatively low CO₂ concentrations typically in the range of 5–20%, the direct conversion of low-concentration CO₂ is more practically relevant and highly

^a Institute of Functional Nano & Soft Materials (FUNSOM), Soochow University, Suzhou, 215123, China. E-mail: weihuang@suda.edu.cn, yanguang@suda.edu.cn

^b Jiangsu Key Laboratory for Advanced Negative Carbon Technologies, Soochow University, Suzhou 215123, China

^c Department of Applied Biology and Chemical Technology, The Hong Kong Polytechnic University, Hung Hom, Kowloon, Hong Kong SAR 999077, China. E-mail: bhuang@polyu.edu.hk

^d Macao Institute of Materials Science and Engineering (MIMSE), MUST-SUDA Joint Research Center for Advanced Functional Materials, Macau University of Science and Technology, Taipa 999078, Macau SAR, China

^e Research Centre for Carbon-Strategic Catalysis, The Hong Kong Polytechnic University, Hung Hom, Kowloon, Hong Kong SAR 999077, China

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d3ee04121b>

‡ These two authors contribute equally.

desirable.⁶ Unfortunately, limited attempts so far have shown that diluted CO₂ concentrations usually compromise the activity and/or selectivity.⁷ This is mainly attributed to inefficient CO₂ capture and unfavorable binding interactions between catalytic sites and substrates.⁸ As a result, the rational design of photocatalysts with optimized catalytic centers and CO₂ binding capacity is crucial.

In nature, enzymes can catalyze complicated redox reactions with high activity and specificity towards target products.⁹ Their active sites are typically composed of metal cations bound to the amino acid residues of the proteins containing nitrogen (N), sulfur (S), and oxygen (O) atoms. For instance, galactose oxidase is a well-known mononuclear oxidoreductase, in which the Cu center is ligated by two pyrrolic N atoms and two phenolic O atoms.¹⁰ Such a Cu–N₂O₂ site can efficiently catalyze the two-electron oxidation of primary alcohols to corresponding aldehydes. Inspired by these biocatalysts, great efforts have been devoted to exploiting artificial enzymes in order to mimic the functions of natural molecules.^{11–14} Salen or salophen compounds are a type of Schiff-base ligands with planar tetradeятate N₂O₂ sites, capable of accommodating various transition metal ions.¹⁵ By judiciously varying metal catalytic centers and ancillary organic ligands, it is possible to fine-tune their catalytic performances. Benefiting from their structural diversity and easy accessibility, metal–salen/salophen complexes have been widely used as enzyme mimics for different catalytic reactions.^{16,17} Nevertheless, their potential in photocatalytic CO₂ reduction has not been explored. This may be partly attributed to the unsatisfactory long-term stability of molecular salen/salophen catalysts during photocatalytic reactions as a result of their aggregation or degradation.¹⁸

Immobilizing molecular catalysts onto suitable porous hosts offers a potential solution to the above issue. Previous investigations have demonstrated that porous skeletons could enhance the robustness of catalytic sites, and facilitate the enrichment of CO₂ molecules in their vicinity.^{19,20} This is of great significance for durable and efficient CO₂ conversion, particularly under diluted CO₂ concentrations.

To this end, here we synthesize a new type of metal–salophen organic framework (M-SOF, M = Co, Ni, Cu, and Zn) by covalently incorporating metal–salophen nodes into porous architectures for the efficient and durable photoreduction of CO₂ at varying concentrations. By tuning the predesigned organic precursors and metal species, the activity and CO selectivity of the resulting catalysts could be readily modulated. Of particular note is that the Ni-SOF containing Ni–N₂O₂ centers is the most photocatalytically active; across a wide range of CO₂ concentrations (5–100 vol%), it exhibits an extremely high photocatalytic activity of up to 16 908 μmol h^{−1} g^{−1} and near-unity selectivity for CO production. Experimental and theoretical studies show that the outstanding catalytic activity of the Ni-SOF originates from its porous frameworks and favorable binding towards CO₂ and reaction intermediates.

Results and discussion

Fig. 1a schematically illustrates the synthetic procedure of Ni-SOFs *via* a metal-template-assisted Schiff-base reaction between

salicylaldehyde and 1,3,5-tris(4-aminophenyl)triazine (TAPT) in *N,N*-dimethylformamide (DMF). During the synthesis, salicylaldehyde and nickel acetate were first mixed in DMF at room temperature to form a soluble O-coordinated Ni intermediate (Ni–O₄), as evidenced by the change in solution color from light green to yellow.²¹ The subsequent addition of TAPT initiated the condensation reaction with the aldehyde functionalities of Ni–O₄ at elevated temperatures and ultimately gave rise to an extended coordination polymer with Ni–N₂O₂ nodes. It is worth noting that such a stepwise synthetic procedure is crucial in order to achieve the desired metal–N₂O₂ coordination. TAPT is employed as the building block because of its nitrogen-rich composition, which would result in nitrogen-rich porous skeletons to facilitate CO₂ capture. Other samples with different metal centers (Co-SOFs, Cu-SOFs, and Zn-SOFs) were synthesized by adding corresponding metal salts under otherwise identical conditions. To investigate the effect of coordination environments on photocatalysis, N-coordinated Ni–N₄, and N/S-coordinated Ni–N₂S₂ were also synthesized by replacing salicylaldehyde with 2-pyridinecarboxaldehyde and 2-mercaptopbenzaldehyde, respectively.

The chemical structures of our samples were first interrogated by Fourier transform infrared (FT-IR) and X-ray photoelectron spectroscopy (XPS). For the sake of clarity, the following discussion will be focused on Ni-SOF as a representative unless otherwise specified. From its FT-IR spectrum, the successful condensation between salicylaldehyde and TAPT is evidenced by the disappearance of the –NH₂ stretching vibrations at 3209 and 3321 cm^{−1} as well as the emergence of a new peak at 1621 cm^{−1} attributed to –CH=≡N (Fig. S1, ESI†).²² The presence of two peaks at 1361 and 1508 cm^{−1}, characteristic of triazine, also signifies the incorporation of TAPT within the coordination networks.²³ Moreover, two weak bands corresponding to Ni–O and Ni–N are observed at 589 and 458 cm^{−1}, respectively.²⁴ They suggest that the coordination nodes in Ni-SOF are formed through the interactions between Ni cations and surrounding imine/hydroxyl groups. Fig. S2 (ESI†) summarizes the XPS results of Ni-SOFs. The N 1s spectrum of the Ni-SOF could be deconvoluted into three peaks assigned to pyridinic N in the triazine unit (398.0 eV), C=≡N (398.8 eV) and Ni–N (399.8 eV).²² Its O 1s spectrum shows two peaks at 528.3 and 531.6 eV, corresponding to Ni–O and C–O, respectively.²⁵ The Ni center in the Ni-SOF is determined to be in the divalent state based on its Ni 2p spectrum.²⁶

The electronic state and coordination environment of the Ni-SOF were further examined by synchrotron X-ray absorption spectroscopy (XAS). Fig. 1b illustrates its X-ray absorption near-edge structure (XANES) spectrum at the Ni K-edge. The white-line peak of Ni-SOFs is similar to that of NiO, corroborating the divalent state of Ni centers in Ni-SOFs. The corresponding Fourier transform extended X-ray absorption fine structure (EXAFS) spectrum of Ni-SOFs exhibits a prominent peak at 1.60 Å, which is a value between the bonding length of Ni–N in nickel phthalocyanine (NiPc) (1.46 Å) and that of Ni–O in NiO (1.65 Å) (Fig. 1c). This again supports the mixed Ni–N/O first-shell coordination.²⁷ No Ni–Ni scattering is observed,

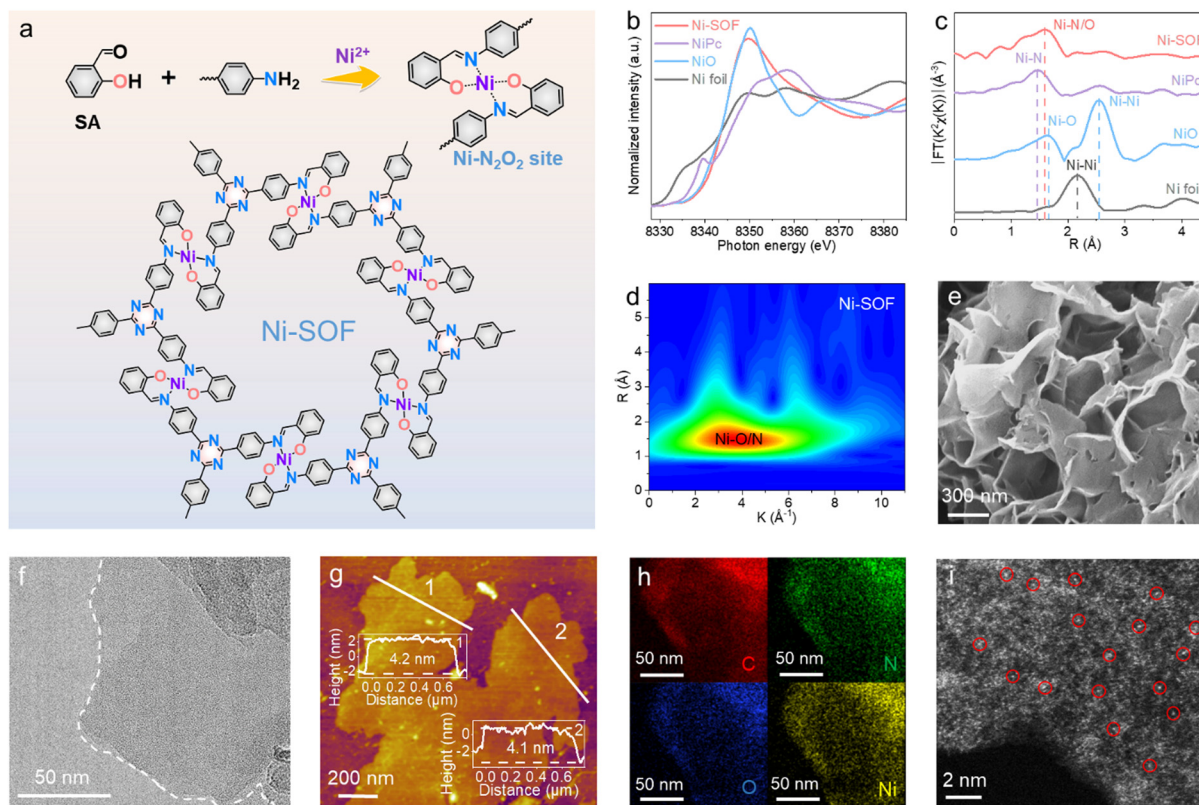


Fig. 1 Synthesis and characterizations of Ni-SOFs. (a) Schematic illustration of the synthetic procedure of Ni-SOFs. (b) XANES and (c) EXAFS spectra of Ni-SOFs, NiPc, NiO and Ni foil. (d) Wavelet transform of the EXAFS spectra of Ni-SOFs. (e) SEM, (f) TEM, (g) AFM, (h) EDS elemental mapping and (i) HAADF-STEM images of Ni-SOFs.

confirming the atomic dispersion of Ni^{2+} cations in Ni-SOFs. Further evidence is obtained from the wavelet transform (WT) of the EXAFS spectra (Fig. 1d and Fig. S3, ESI[†]). The maximum WT position of Ni-SOFs apparently differs from those of Ni-N in NiPc and Ni-O in NiO, but partially overlaps with them.

To fully unveil the steric structure of $\text{Ni-N}_2\text{O}_2$ nodes in Ni-SOFs, a model molecule (denoted as Ni-salophen) was synthesized as its mimic and analyzed by single-crystal X-ray diffraction (XRD) measurement (Fig. S4 and Table S1, ESI[†]). It is shown to possess a four-coordinated Ni center with the first coordination sphere consisting of two phenolic O atoms and two imine N atoms. The bond length of Ni-O is calculated to be 1.83 Å, slightly shorter than that of Ni-N (1.91 Å) as a result of the higher electronegativity of O atoms. Two pairs of *trans*-coordinated O and N atoms around the central Ni atom form a nearly planar rhomboid geometry owing to the small differences in bond lengths and bond angles. It is believed that such an open metal center with exposed axial coordination is advantageous for CO_2 binding and activation.²⁸

Scanning electron microscopy (SEM) and transmission electron microscopy (TEM) imaging show that the Ni-SOF consists of crumpled and stacked nanosheets (Fig. 1e and f). Individual nanosheets are estimated to have a size ranging from 200 nm to 1 μm . From atomic force microscopy (AFM) height profiles at multiple locations, the nanosheet thickness is measured to be approximately 4 nm (Fig. 1g). Such an ultrathin nanosheet

geometry is desirable for catalytic applications as it promotes the exposure of inner active sites and maximizes atom utilization. Energy dispersive X-ray spectroscopy (EDS) mapping under scanning transmission electron microscopy (STEM) reveals the uniform distribution of C, N, O and Ni on Ni-SOF nanosheets (Fig. 1h). The aberration-corrected high-angle annular dark-field STEM (HAADF-STEM) image shows discrete bright spots (some marked with red circles) corresponding to heavy Ni atoms, corroborating the atomic dispersion of Ni sites throughout the examined region (Fig. 1i). Moreover, the Ni content in Ni-SOFs is found to be 10.5 wt% as analyzed by inductively coupled plasma mass spectrometry (ICP-MS), which is in good agreement with the theoretical value (10.2 wt%). N_2 isotherm sorption measurements show that the Ni-SOF has a specific surface area of 116 $\text{m}^2 \text{ g}^{-1}$ (Fig. S5, ESI[†]). In addition to Ni-SOFs, the structures of other samples with different metal sites and coordination environments were also characterized, and results are summarized in the ESI[†] (Fig. S6–S8 and Table S2).

We evaluated the photocatalytic performance of our samples under pure CO_2 . The reactions were conducted in a mixed solution of acetonitrile and water under visible light irradiation ($\lambda > 420 \text{ nm}$) using $[\text{Ru}(\text{bpy})_3]\text{Cl}_2$ as the photosensitizer and triisopropanolamine (TIPA) as the hole scavenger. Gaseous products (e.g., CO and H_2) were analyzed and quantified by gas chromatography (GC) based on their calibration curves (Fig. S9, ESI[†]). We explored different solvent ratios, catalyst

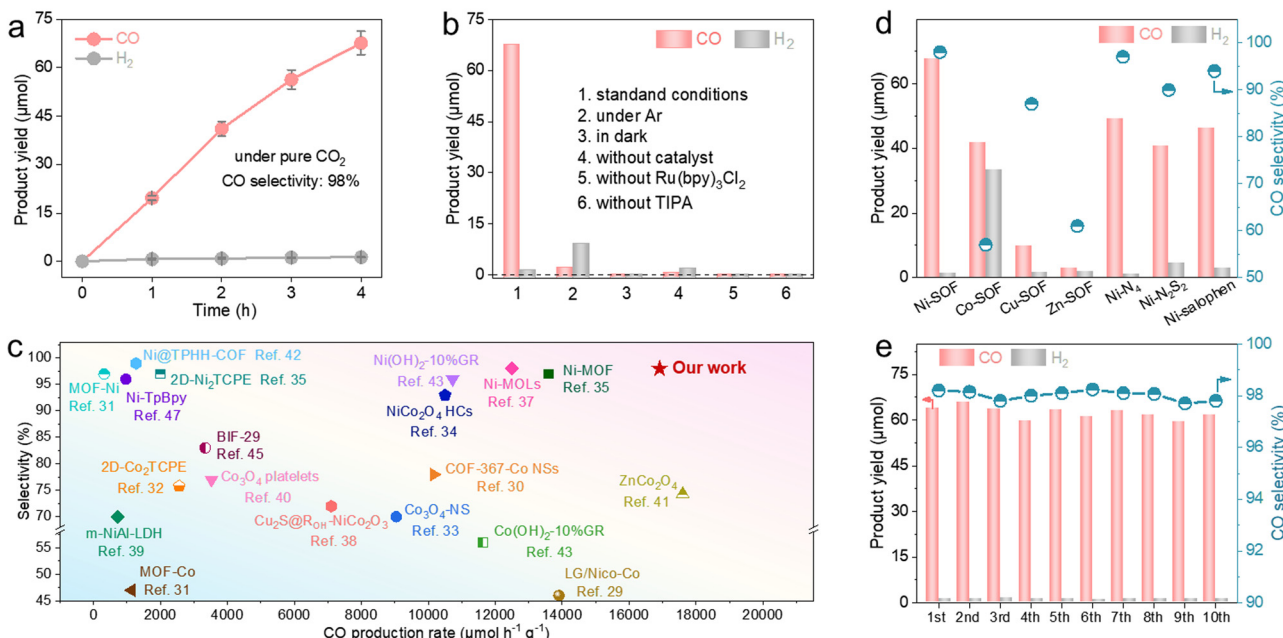


Fig. 2 Photocatalytic performance of Ni-SOFs under pure CO₂. (a) Time-dependent CO and H₂ production on Ni-SOFs. (b) Control experiments conducted under different conditions. (c) Performance comparison of Ni-SOFs with other organic or inorganic catalysts in terms of the CO production rate and selectivity under pure CO₂. (d) CO production and selectivity of different samples. (e) Cycling tests of Ni-SOFs.

concentrations, and hole scavengers (Fig. S10, ESI[†]). Under the optimized conditions, the Ni-SOF exhibits an almost linear accumulation of CO over time and yields a total of 67.6 μmol of CO after 4 h irradiation (Fig. 2a). This corresponds to an average mass-specific CO evolution rate of 16 908 μmol h⁻¹ g⁻¹. Its maximum apparent quantum efficiency (AQE) toward CO production is 2.8% at 450 nm (Fig. S11, ESI[†]). The isotope experiment using ¹³CO₂ provides evidence that CO stems from CO₂ photoreduction rather than other organic substances or the catalyst itself (Fig. S12, ESI[†]). Control experiments also show that negligible CO is produced in the absence of light, CO₂, Ni-SOF, photosensitizer or hole scavenger under otherwise identical conditions (Fig. 2b). In addition to CO, only a trace amount of H₂ (1.3 μmol) is measured on Ni-SOFs, and no liquid products (such as formic acid or methanol) are detected in the reaction solution by nuclear magnetic resonance (NMR) spectroscopy (Fig. S13, ESI[†]). As a result, a high CO selectivity of 98% is achieved for our catalyst. It is worth highlighting that the great activity and selectivity measured for our Ni-SOF places it at the top of other organic competitors including covalent organic frameworks (COFs) and metal–organic frameworks (MOFs) as well as most inorganic counterparts (Fig. 2c and Table S3, ESI[†]). In particular, its mass-specific CO production rate is about 17.5 and 8.5 times larger than those of the state-of-the-art COF catalyst (Ni-TpBpy) and MOF catalyst (2D-Ni₂TCPE), respectively.^{32,47} We are aware that higher mass-specific rates have been reported for some transition metal oxides, but their selectivity for carbonaceous products was generally lower than 80%.⁴¹

The catalytic performances of coordinated metal centers are greatly impacted by not only the metal identities but also their

local chemical environments (Fig. 2d). Under identical reaction conditions, the Co-SOF shows a significantly decreased CO production rate of 10 157.5 μmol h⁻¹ g⁻¹ and a higher H₂ evolution activity of 8322.5 μmol h⁻¹ g⁻¹, leading to a low CO selectivity of 57%. Even though the Cu-SOF demonstrates a good CO selectivity (87%), its CO production rate is 7 times lower than that of Ni-SOFs. The Zn-SOF exhibits the lowest CO production rate of 695 μmol h⁻¹ g⁻¹ and a moderate CO selectivity of 61%. Moreover, the substitution of O atoms in the first coordination sphere of Ni-SOFs with less electronegative N or S (Ni-N₄ and Ni-N₂S₂, respectively) leads to noticeable decreases in both the activity and selectivity, and their measured activities are found to correlate with the electronegativity of coordinating elements. The above results suggest the critical roles of metal centers and their coordination environments during CO₂ photoreduction. The Ni-SOF featuring Ni-N₂O₂ active sites represents the optimal combination to enable active and selective CO₂ photoreduction to CO. In addition, despite having identical Ni-N₂O₂ catalytic sites, the Ni-SOF is more active than Ni-salophen, presumably due to its porous microstructure which enriches local CO₂ concentration.⁴⁸

Stability is another important performance metric of photocatalysts. Here, we carried out long-term photocatalysis as shown in Fig. S14 (ESI[†]). The CO production rate of the Ni-SOF gradually decreases and reaches a plateau after 5 h. However, its activity can be completely restored when fresh [Ru(bpy)₃]²⁺ is replenished. This observation suggests that the performance decay is due to the degradation of the photosensitizer rather than the Ni-SOF.⁴⁹ In contrast, Ni-salophen experiences an irreversible activity loss after about 5 h. It underlines the evident advantage of metal-salophen organic

frameworks with improved structural robustness and catalytic stability. The excellent stability of Ni-SOFs is also supported by the cycling experiment showing no noticeable performance loss after a total of 40 h illumination (Fig. 2e). In addition, the ICP-MS analysis of the reaction filtrate after the cycling test reveals a negligible Ni concentration (0.264 ppm), thus ruling out the possibility of Ni leaching out of Ni-SOFs during photocatalysis. Spectroscopic and microscopic analyses of the recovered Ni-SOF disclose no discernable structural changes (Fig. S15, ESI[†]).

Effective charge transfer from the photosensitizer to the catalytic sites is a prerequisite for initiating redox reactions. To probe the charge transfer behavior between $[\text{Ru}(\text{bpy})_3]^{2+}$ and Ni-SOFs, steady-state and time-resolved photoluminescence (PL) spectroscopy studies were conducted. As shown in Fig. S16 (ESI[†]), the steady-state PL peak intensity of $[\text{Ru}(\text{bpy})_3]^{2+}$ gradually decreases as the Ni-SOF concentration increases, while the TIPA concentration has no significant effect on the

PL intensity. Corresponding Stern–Volmer plots show the much higher quenching efficiency of Ni-SOFs than TIPA. These results reveal that the electron transfer from the photosensitizer to the catalyst is the initial step of the photocatalytic cycle.³² The time-resolved PL spectrum of $[\text{Ru}(\text{bpy})_3]^{2+}$ follows a single exponential decay with an average PL lifetime of 237 ns, consistent with literature results (Fig. 3a).⁵⁰ In the presence of Ni-SOFs, a double-exponential decay is observed with fitted lifetimes of 0.49 ns and 225 ns (Fig. 3b). The former is attributed to the charge transfer between $[\text{Ru}(\text{bpy})_3]^{2+}$ and Ni-SOFs. Compared to those measured in the presence of other M-SOFs (M = Co, Cu, and Zn), the shorter lifetime observed with Ni-SOFs at the early stage suggests more efficient electron transfer (Fig. S17, ESI[†]).⁵¹ Moreover, electron paramagnetic resonance (EPR) spectroscopy was conducted (Fig. 3c). Upon irradiation, a sharp EPR signal ascribed to Ni^{+} is observed at $g = 2.06$,⁵² while it becomes significantly attenuated upon CO_2

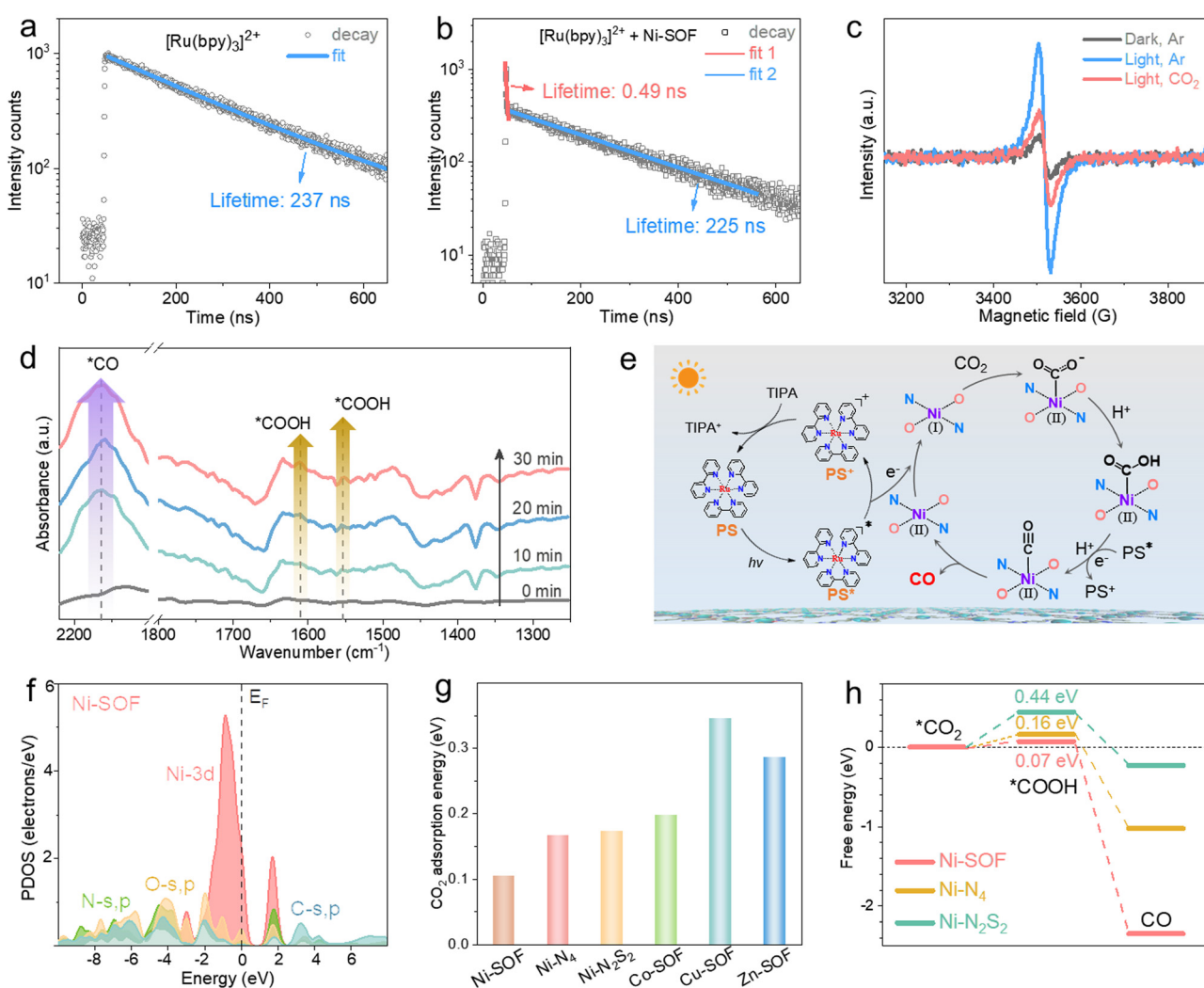


Fig. 3 Charge transfer kinetics and possible reaction mechanism. (a) and (b) Time-resolved PL spectra of (a) $[\text{Ru}(\text{bpy})_3]^{2+}$ and (b) $[\text{Ru}(\text{bpy})_3]^{2+} + \text{Ni-SOF}$ in acetonitrile. (c) EPR spectra of Ni-SOFs under different conditions. (d) Operando DRIFTS spectra of Ni-SOFs under irradiation for different durations of time. (e) Schematic illustration of the proposed catalytic cycle. (f) PDOS spectra of Ni-SOFs. (g) CO_2 adsorption energy on different catalysts. (h) Calculated free energy of CO_2 photoreduction on Ni sites with different coordination environments.

introduction. Considering that Ni^{2+} is EPR-silent, this observation indicates that Ni^{2+} sites can readily accept electrons from the excited photosensitizer, and subsequently transfer them to CO_2 to initiate the reduction reaction. The CO_2 reduction process was then be tracked by *in situ* diffuse reflectance infrared Fourier transform spectroscopy (DRIFTS). Fig. 3d clearly shows the vibration bands from intermediates within a 30-minute irradiation period. The two peaks at 1611 and 1552 cm^{-1} are characteristic of the $^{*}\text{COOH}$ intermediate, and their peak intensity increases as the irradiation continues.⁵³ In the meantime, a pronounced $^{*}\text{CO}$ adsorption peak at 2163 cm^{-1} is also observed, indicating a high $^{*}\text{CO}$ surface coverage under irradiation.⁵⁴

Based on the above spectroscopic results, a possible mechanism for CO_2 photoreduction is proposed and illustrated in Fig. 3e. Upon visible light irradiation, $[\text{Ru}(\text{bpy})_3]^{2+}$ (PS) is promoted to its excited state (PS^*). Subsequently, photoexcited electrons are transferred to the Ni^{2+} sites of Ni-SOFs within a

short time scale, generating catalytically active Ni^+ centers. The oxidized photosensitizer (PS^+) is then reduced back to its ground state by accepting electrons from TIPA. Electrons located on the Ni^+ centers are further transferred to absorbed CO_2 , initiating the reduction reaction which involves the sequential formation of $^{*}\text{COOH}$ and $^{*}\text{CO}$ intermediates. At the end, $^{*}\text{CO}$ desorbs from the catalyst surface, thus stopping the reaction cycle.

To gain more insights into the catalytic mechanism, density functional theory (DFT) calculations were performed. The electronic distribution diagrams of Ni-SOFs, Ni- N_4 , and Ni- N_2S_2 near the Fermi level (E_F) are depicted in Fig. S18 (ESI[†]). The maximum bonding orbital distribution is located closer to Ni in Ni-SOFs compared to those in Ni- N_4 and Ni- N_2S_2 , suggesting the electron-rich feature of Ni sites in Ni-SOFs. The electronic modulation of metal centers by the coordination environment is evident from the projected partial density of states (PDOS) analysis. As shown in Fig. 3f, the Ni-3d orbitals of Ni-SOFs

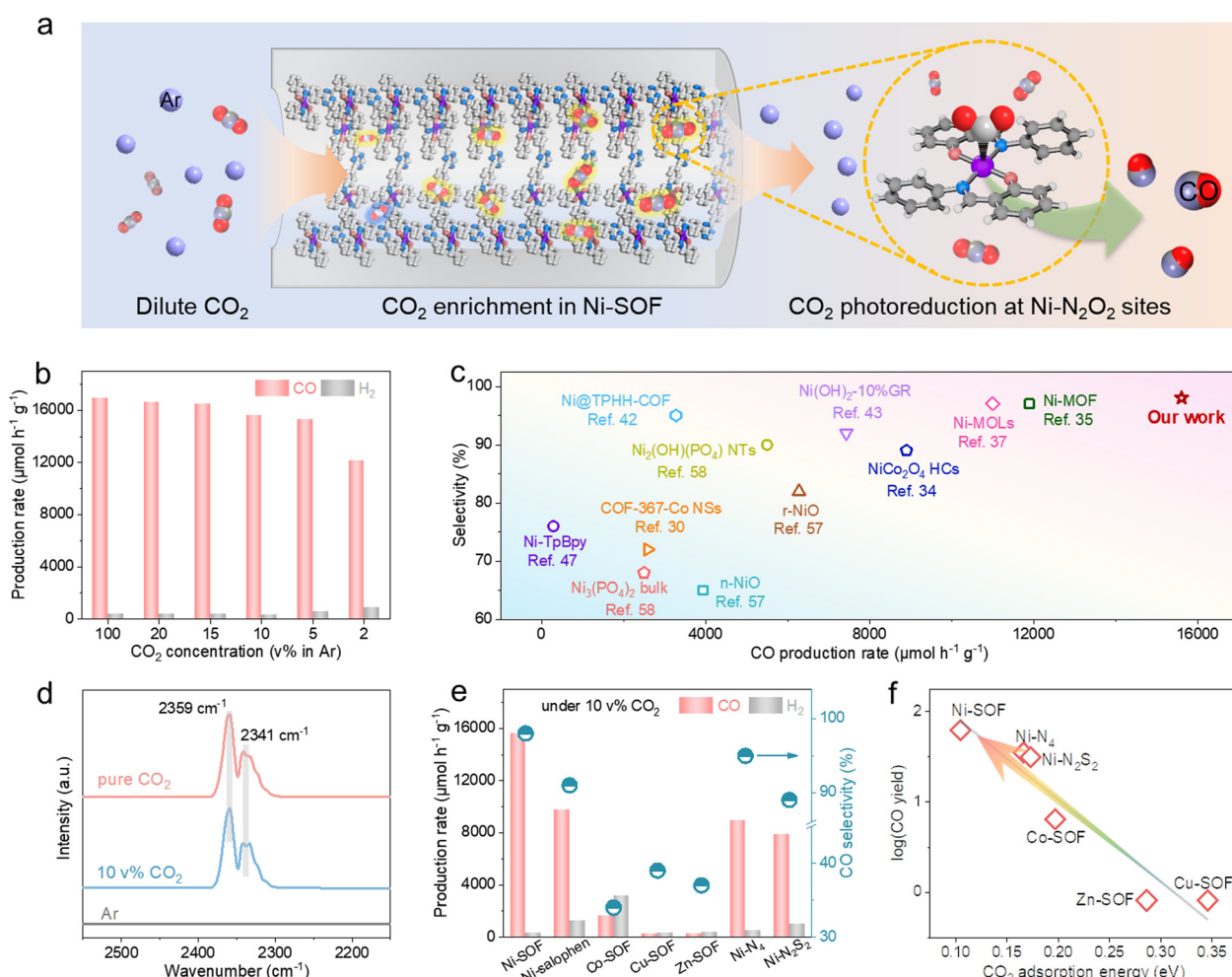


Fig. 4 Performance evaluation of the catalysts under dilute CO_2 . (a) Schematic illustration of the enrichment and photoreduction of dilute CO_2 in Ni-SOFs. (b) Photocatalytic activities of Ni-SOFs at various CO_2 concentrations. (c) Performance comparison of Ni-SOFs with other organic or inorganic catalysts in terms of the CO production rate and selectivity under 10 vol% CO_2 in Ar. (d) DRIFTS spectra of Ni-SOFs in pure Ar, pure CO_2 or 10 vol% CO_2 atmospheres. (e) CO production rate and selectivity of different catalysts under 10 vol% CO_2 . (f) Correlation between the logarithm of the CO production rate on different catalysts under 10 vol% CO_2 and their CO_2 adsorption energy.

exhibit a sharp peak near E_F at $E_V - 0.88$ eV ($E_V = 0$ eV). The s, p orbitals of O and N sites have good overlaps with the Ni-3d orbitals, indicating their strong binding interactions with central Ni atoms are necessary for the formation of a stable coordination configuration and efficient site-to-site electron transfer.⁵⁵ By comparison, the metallic 3d orbitals of other catalysts (Ni-N₄, Ni-N₂S₂, Co-N₂O₂, Cu-N₂O₂, and Zn-N₂O₂) are located at more negative positions related to their E_F with obviously decreased overlaps with the s and p orbitals of surrounding heteroatoms (Fig. S19, ESI[†]). The d-band center and optical absorption comparisons also confirm the improved catalytic activity and optical properties of Ni-SOFs to promote the photocatalytic performances of CO₂ reduction (Fig. S20, ESI[†]).

Furthermore, CO₂ adsorption energies on different catalysts were calculated and compared (Fig. 3g). The Ni-SOF has the lowest adsorption energy, suggesting its strongest binding with CO₂. This result is corroborated by the temperature programmed CO₂ desorption (TPD) measurements showing that the Ni-SOF has the largest CO₂ desorption capacity and highest desorption temperature (Fig. S21, ESI[†]). Favorable CO₂ binding ability is believed to be conducive to CO₂ capture and enrichment around the catalytic sites, thereby accelerating CO₂ conversion.⁵⁶ In addition, the free energy profiles of different catalysts during CO₂ reduction were simulated. The conversion from *CO₂ to *COOH is found to be the rate-determining step (RDS) for all the catalysts (Fig. 3h and Fig. S22, ESI[†]). Among them, the Ni-SOF shows the smallest energy barrier of 0.07 eV, indicating that the redox reaction on the Ni-SOF is the most favorable. Further conversion from *COOH to CO is exothermic and thereby spontaneous once *COOH is formed.

Encouraged by the great catalytic capacity and favorable CO₂ binding of the Ni-SOF, we moved on to assess its catalytic performance at dilute CO₂ concentrations. Fig. 4a schematically illustrates the local enrichment and subsequent photoreduction of diluted CO₂ within the porous skeletons of Ni-SOFs. As demonstrated in Fig. 4b and Fig. S23 (ESI[†]), the Ni-SOF exhibits an exceptional catalytic performance at varying CO₂ concentrations. Both the CO production rate and selectivity remain largely stable across a wide range of CO₂ concentrations from 5 to 20 vol% in Ar. At a CO₂ concentration as low as 2 vol%, the mass-specific CO production rate is measured to be 12 100 $\mu\text{mol h}^{-1} \text{g}^{-1}$, retaining 72% of the value achieved under pure CO₂ as well as a high CO selectivity of 93%. Moreover, the AQE value at 10 vol% CO₂ is 2.7% at 450 nm, which is very close to that obtained under pure CO₂ (2.8%) (Fig. S24, ESI[†]). To the best of our knowledge, such high activity and selectivity have rarely been achieved before under dilute CO₂ (Fig. 4c and Table S4, ESI[†]),^{37,43,57,58} exceeding those of other organic and inorganic candidates, even measured under pure CO₂.^{34,42} The DRIFTS spectra of Ni-SOFs under pure and diluted (10 vol%) CO₂ show two identical absorption bands at 2359 and 2341 cm^{-1} , assigned to adsorbed CO₂ with the end-on configuration (Fig. 4d).⁵⁹ Importantly, their peak intensity and integrated areas are found to be largely independent of the CO₂ concentration. This implies that Ni-SOFs can enrich local CO₂ concentration when the feedstock becomes much diluted and may rationalize

the exceptional photocatalytic activity measured under low-concentration CO₂.

The great photocatalytic performance under low-concentration CO₂ appears to be unique to Ni-SOFs. Ni-salophen presents considerably lower activity despite its similar coordination environment with Ni-SOFs (Fig. 4e). Replacing the central Ni with Co results in a decreased mass-specific activity of 2167 $\mu\text{mol h}^{-1} \text{g}^{-1}$ (more than 4 times lower than that under pure CO₂ on Co-SOFs) and low CO selectivity of 34%. More strikingly, CO₂ photoreduction on Cu-SOFs and Zn-SOFs becomes almost quenched under the same conditions. Furthermore, both Ni-N₄ and Ni-N₂S₂ exhibit about half of the activity of Ni-SOFs under 10 vol% CO₂. We note that the measured activity here is roughly correlated with the CO₂ binding energy predicted in Fig. 4f. The strongest binding affinity of Ni-SOF affords our catalyst with the most appealing activity by enriching CO₂ local concentration and facilitating its subsequent conversion.

Conclusions

In summary, we synthesized a new class of M-SOF materials *via* facile metal-assisted Schiff-base reactions for efficient photocatalytic CO₂ reduction. All the samples are composed of atomically dispersed metal sites and controlled local coordination environments, both of which could be precisely modulated *via* varying the predesigned organic moieties and metal species. The optimal sample, the Ni-SOF with Ni-N₂O₂ nodes, showed the highest CO production rate of 16 908 $\mu\text{mol h}^{-1} \text{g}^{-1}$ with an excellent selectivity up to 98%. Impressively, the Ni-SOF was able to maintain its photocatalytic performance across a wide range of CO₂ concentrations (5–20 vol% in Ar) without reducing the selectivity. Experimental and theoretical studies revealed that the superior catalytic performance of Ni-SOFs mainly originated from the favorable d-band electronic modulation of the coordinated Ni atoms and optimal binding strength with the intermediates. In particular, the strong CO₂ affinity of Ni-SOFs is identified as the key factor for the efficient conversion of diluted CO₂. Our study demonstrates a promising strategy for rationalizing the chemical and electronic configurations of metal catalytic centers toward the direct utilization of diluted CO₂ from industrial exhaust gas.

Author contributions

Y. L. and W. H. conceived the project and designed the experiments. X. Y. synthesized the catalysts and conducted the structure analysis and photocatalytic tests. B. H. and M. S. performed the theoretical calculations. T. Y. and L. Z. conducted the X-ray absorption measurements. L. J. carried out TEM characterization studies. M. C. performed TPD measurements. X. Y., M. S., W. H., B. H., and Y. L. co-wrote the paper. All authors discussed the results and commented on the manuscript.

Conflicts of interest

There are no conflicts to declare.

Acknowledgements

We acknowledge the financial support received from the National Key R&D Program of China (2021YFA1501101), the National Natural Science Foundation of China (22002100, U2002213, and 52161160331), the National Natural Science Foundation of China/Research Grant Council of Hong Kong Joint Research Scheme (N_PolyU502/21), the National Natural Science Foundation of China/Research Grants Council of Hong Kong Collaborative Research Scheme (CRS_PolyU504/22), the Natural Science Foundation of Jiangsu Province (BK20220027), the Natural Science Foundation of the Jiangsu Higher Education Institutions of China (20KJA430002), the Science and Technology Development Fund Macau SAR (0077/2021/A2), the Collaborative Innovation Center of Suzhou Nano Science and Technology, and the funding for Projects of Strategic Importance of The Hong Kong Polytechnic University (Project Code: 1-ZE2V), the Shenzhen Fundamental Research Scheme-General Program (JCYJ20220531090807017), the Natural Science Foundation of Guangdong Province (2023A1515012219) and Departmental General Research Fund of The Hong Kong Polytechnic University (Project Code: ZVUL). We thank the Shanghai Synchrotron Radiation Facility (beamline 11B) for the allocation of synchrotron beamtime, Dr Jie Xu for HAADF-STEM characterization and Chunpeng Wu for DRFITS measurements. B. H. is also grateful for the support provided by the Research Centre for Carbon-Strategic Catalysis (RC-CSC), the Research Institute for Smart Energy (RISE), and the Research Institute for Intelligent Wearable Systems (RI-IWEAR) of the Hong Kong Polytechnic University.

Notes and references

- 1 D. Campbell-Lendrum, T. Neville, C. Schweizer and M. Neira, *Nat. Med.*, 2023, **29**, 1631–1638.
- 2 C. J. Nielsen, H. Herrmann and C. Weller, *Chem. Soc. Rev.*, 2012, **41**, 6684–6704.
- 3 J. Wu, Y. Huang, W. Ye and Y. Li, *Adv. Sci.*, 2017, **4**, 1700194.
- 4 E. Gong, S. Ali, C. B. Hiragond, H. S. Kim, N. S. Powar, D. Kim, H. Kim and S.-I. In, *Energy Environ. Sci.*, 2022, **15**, 880–937.
- 5 W. Huang, W. Luo and Y. Li, *Mater. Today*, 2020, **40**, 160–172.
- 6 T. Nakajima, Y. Tamaki, K. Ueno, E. Kato, T. Nishikawa, K. Ohkubo, Y. Yamazaki, T. Morimoto and O. Ishitani, *J. Am. Chem. Soc.*, 2016, **138**, 13818–13821.
- 7 Y. Yamazaki, M. Miyaji and O. Ishitani, *J. Am. Chem. Soc.*, 2022, **144**, 6640–6660.
- 8 X. Wu, Y. Li, G. Zhang, H. Chen, J. Li, K. Wang, Y. Pan, Y. Zhao, Y. Sun and Y. Xie, *J. Am. Chem. Soc.*, 2019, **141**, 5267–5274.
- 9 S. Bierbaumer, M. Nattermann, L. Schulz, R. Zschoche, T. J. Erb, C. K. Winkler, M. Tinzl and S. M. Glueck, *Chem. Rev.*, 2023, **123**, 5702–5754.
- 10 A. John, M. M. Shaikh and P. Ghosh, *Dalton Trans.*, 2008, 2815–2824.
- 11 H. Huang, X. Jing, J. Deng, C. Meng and C. Duan, *J. Am. Chem. Soc.*, 2023, **145**, 2170–2182.
- 12 M. Vázquez-González, Z. Zhou, Y. Biniuri, B. Willner and I. Willner, *Biochemistry*, 2021, **60**, 956–965.
- 13 A. A. Shtensteinman, *Catalysts*, 2023, **13**, 415.
- 14 Z. Wang, P. Yeary, X. Feng and W. Lin, *J. Am. Chem. Soc.*, 2023, **145**, 8647–8655.
- 15 J. Zhang, L. Xu and W.-Y. Wong, *Coord. Chem. Rev.*, 2018, **355**, 180–198.
- 16 C. J. Whiteoak, G. Salassa and A. W. Kleij, *Chem. Soc. Rev.*, 2012, **41**, 622–631.
- 17 S. Signorella, C. Palopoli and G. Ledesma, *Coord. Chem. Rev.*, 2018, **365**, 75–102.
- 18 C.-B. Li, Y. Chu, J. He, J. Xie, J. Liu, N. Wang and J. Tang, *ChemCatChem*, 2019, **11**, 6324–6331.
- 19 V. Mouarrawis, R. Plessius, J. I. van der Vlugt and J. N. H. Reek, *Front. Chem.*, 2018, **6**, 623.
- 20 M. Lu, M. Zhang, J. Liu, Y. Chen, J.-P. Liao, M.-Y. Yang, Y.-P. Cai, S.-L. Li and Y.-Q. Lan, *Angew. Chem., Int. Ed.*, 2022, **61**, e202200003.
- 21 N. Zhao, K. Cai and H. He, *Dalton Trans.*, 2020, **49**, 11467–11479.
- 22 H. Zhang, Z. Lin, P. Kidkhunthod and J. Guo, *Angew. Chem., Int. Ed.*, 2023, **62**, e202217527.
- 23 L. Chen, L. Wang, Y. Wan, Y. Zhang, Z. Qi, X. Wu and H. Xu, *Adv. Mater.*, 2020, **32**, 1904433.
- 24 S. Rai, A. Bajpai and S. Lokhandwala, *J. Polym.*, 2013, **2013**, 278576.
- 25 P. Thangasamy, S. Shanmuganathan and V. Subramanian, *Nanoscale Adv.*, 2020, **2**, 2073–2079.
- 26 J. Wang, W. Zhu, F. Meng, G. Bai, Q. Zhang and X. Lan, *ACS Catal.*, 2023, **13**, 4316–4329.
- 27 H. B. Yang, S.-F. Hung, S. Liu, K. Yuan, S. Miao, L. Zhang, X. Huang, H.-Y. Wang, W. Cai, R. Chen, J. Gao, X. Yang, W. Chen, Y. Huang, H. M. Chen, C. M. Li, T. Zhang and B. Liu, *Nat. Energy*, 2018, **3**, 140–147.
- 28 C. A. Trickett, A. Helal, B. A. Al-Maythalony, Z. H. Yamani, K. E. Cordova and O. M. Yaghi, *Nat. Rev. Mater.*, 2017, **2**, 17045.
- 29 G. Yang, S. Li, N. Li, P. Zhang, C. Su, L. Gong, B. Chen, C. Qu, D. Qi, T. Wang and J. Jiang, *Angew. Chem., Int. Ed.*, 2022, **61**, e202205585.
- 30 W. Liu, X. Li, C. Wang, H. Pan, W. Liu, K. Wang, Q. Zeng, R. Wang and J. Jiang, *J. Am. Chem. Soc.*, 2019, **141**, 17431–17440.
- 31 X.-K. Wang, J. Liu, L. Zhang, L.-Z. Dong, S.-L. Li, Y.-H. Kan, D.-S. Li and Y.-Q. Lan, *ACS Catal.*, 2019, **9**, 1726–1732.
- 32 H.-L. Zheng, S.-L. Huang, M.-B. Luo, Q. Wei, E.-X. Chen, L. He and Q. Lin, *Angew. Chem., Int. Ed.*, 2020, **59**, 23588–23592.
- 33 W. Chen, B. Han, C. Tian, X. Liu, S. Liang, H. Deng and Z. Lin, *Appl. Catal., B*, 2019, **244**, 996–1003.
- 34 B. Han, J. Song, S. Liang, W. Chen, H. Deng, X. Ou, Y.-J. Xu and Z. Lin, *Appl. Catal., B*, 2020, **260**, 118208.
- 35 B. Han, X. Ou, Z. Zhong, S. Liang, X. Yan, H. Deng and Z. Lin, *Appl. Catal., B*, 2021, **283**, 119594.

36 K. Song, S. Liang, X. Zhong, M. Wang, X. Mo, X. Lei and Z. Lin, *Appl. Catal., B*, 2022, **309**, 121232.

37 B. Han, X. Ou, Z. Deng, Y. Song, C. Tian, H. Deng, Y.-J. Xu and Z. Lin, *Angew. Chem., Int. Ed.*, 2018, **57**, 16811–16815.

38 L. Li, X. Dai, D.-L. Chen, Y. Zeng, Y. Hu and X. W. Lou, *Angew. Chem., Int. Ed.*, 2022, **61**, e202205839.

39 L. Tan, S.-M. Xu, Z. Wang, Y. Xu, X. Wang, X. Hao, S. Bai, C. Ning, Y. Wang, W. Zhang, Y. K. Jo, S.-J. Hwang, X. Cao, X. Zheng, H. Yan, Y. Zhao, H. Duan and Y.-F. Song, *Angew. Chem., Int. Ed.*, 2019, **58**, 11860–11867.

40 C. Gao, Q. Meng, K. Zhao, H. Yin, D. Wang, J. Guo, S. Zhao, L. Chang, M. He, Q. Li, H. Zhao, X. Huang, Y. Gao and Z. Tang, *Adv. Mater.*, 2016, **28**, 6485–6490.

41 S. Wang, Z. Ding and X. Wang, *Chem. Commun.*, 2015, **51**, 1517–1519.

42 M. Dong, J. Zhou, J. Zhong, H.-T. Li, C.-Y. Sun, Y.-D. Han, J.-N. Kou, Z.-H. Kang, X.-L. Wang and Z.-M. Su, *Adv. Funct. Mater.*, 2022, **32**, 2110136.

43 K.-Q. Lu, Y.-H. Li, F. Zhang, M.-Y. Qi, X. Chen, Z.-R. Tang, Y. M. A. Yamada, M. Anpo, M. Conte and Y.-J. Xu, *Nat. Commun.*, 2020, **11**, 5181.

44 J.-W. Wang, L.-Z. Qiao, H.-D. Nie, H.-H. Huang, Y. Li, S. Yao, M. Liu, Z.-M. Zhang, Z.-H. Kang and T.-B. Lu, *Nat. Commun.*, 2021, **12**, 813.

45 H.-X. Zhang, Q.-L. Hong, J. Li, F. Wang, X. Huang, S. Chen, W. Tu, D. Yu, R. Xu, T. Zhou and J. Zhang, *Angew. Chem., Int. Ed.*, 2019, **58**, 11752–11756.

46 K. Niu, Y. Xu, H. Wang, R. Ye, H. L. Xin, F. Lin, C. Tian, Y. Lum, K. C. Bustillo, M. M. Doepp, M. T. M. Koper, J. Ager, R. Xu and H. Zheng, *Sci. Adv.*, 2017, **3**, e1700921.

47 W. Zhong, R. Sa, L. Li, Y. He, L. Li, J. Bi, Z. Zhuang, Y. Yu and Z. Zou, *J. Am. Chem. Soc.*, 2019, **141**, 7615–7621.

48 W. Zhou, Q.-W. Deng, G.-Q. Ren, L. Sun, L. Yang, Y.-M. Li, D. Zhai, Y.-H. Zhou and W.-Q. Deng, *Nat. Commun.*, 2020, **11**, 4481.

49 J. Van Houten and R. J. Watts, *J. Am. Chem. Soc.*, 1976, **98**, 4853–4858.

50 Y. Su, Z. Song, W. Zhu, Q. Mu, X. Yuan, Y. Lian, H. Cheng, Z. Deng, M. Chen, W. Yin and Y. Peng, *ACS Catal.*, 2021, **11**, 345–354.

51 C. Gao, S. Chen, Y. Wang, J. Wang, X. Zheng, J. Zhu, L. Song, W. Zhang and Y. Xiong, *Adv. Mater.*, 2018, **30**, 1704624.

52 M. Hartmann, N. Azuma and L. Kevan, *J. Phys. Chem. C*, 1995, **99**, 10988–10994.

53 W. Ma, S. Xie, T. Liu, Q. Fan, J. Ye, F. Sun, Z. Jiang, Q. Zhang, J. Cheng and Y. Wang, *Nat. Catal.*, 2020, **3**, 478–487.

54 G. Qian, W. Lyu, X. Zhao, J. Zhou, R. Fang, F. Wang and Y. Li, *Angew. Chem., Int. Ed.*, 2022, **61**, e202210576.

55 S. Jiao, X. Fu and H. Huang, *Adv. Funct. Mater.*, 2022, **32**, 2107651.

56 Y. Wang, N.-Y. Huang, J.-Q. Shen, P.-Q. Liao, X.-M. Chen and J.-P. Zhang, *J. Am. Chem. Soc.*, 2018, **140**, 38–41.

57 W. Chen, X. Liu, B. Han, S. Liang, H. Deng and Z. Lin, *Nano Res.*, 2021, **14**, 730–737.

58 S. Liang, X. Liu, Z. Zhong, B. Han, X. Zhong, W. Chen, K. Song, H. Deng and Z. Lin, *Nano Res.*, 2021, **14**, 2558–2567.

59 P. D. C. Dietzel, R. E. Johnsen, H. Fjellvag, S. Bordiga, E. Groppo, S. Chavan and R. Blom, *Chem. Commun.*, 2008, 5125–5127.