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Post-synthetic modification of covalent organic frameworks with active manganese centers for electrocatalytic CO₂ reduction in water†

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The development of effective catalysts for the CO₂ reduction reaction (CO₂RR) is essential for transforming atmospheric CO₂ into valuable chemical scaffolds. While numerous catalysts have been developed for the CO₂RR, few are suitable for use in aqueous systems due to inherent design challenges. In this context, Covalent Organic Frameworks (COFs) have emerged as promising materials for the CO₂RR in water, offering potential solutions to these challenges. Thanks to their porosity, high surface area and crystalline structure, COFs are excellent hosts for single-atom catalysts (SACs), enabling the immobilization of high-value species and their utilization in heterogeneous catalytic processes. For this reason, we have explored the catalytic activity of a terpyridine–manganese complex integrated into a COF lattice, which was successfully synthesized and characterized, confirming the presence of the metal ion in the material with spectroscopic techniques such as XPS and EDS. This new material has proved to be an active heterogeneous catalyst for the CO₂RR in water as solvent, achieving a faradaic yield of 42% for CO at 300 mV overpotential and 16% for formate when 600 mV was applied. Furthermore, an *ab initio* theoretical study was performed to provide a plausible mechanism of the CO₂RR to elucidate the CO evolution pathway.

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Introduction

CO₂ has become a target molecule to use as a C1 building block to synthesize high-value renewable solar fuels in the context of

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circular economy.^{1–3} The electrocatalytic reduction of CO₂ (CO₂RR) to valuable products such as liquid fuels can not only reduce CO₂ accumulation in the atmosphere but also lead to pathways toward new renewable chemicals and high energy density fuels.⁴ Careful design of catalysts for this transformation has delivered molecular complexes with extremely high turnover frequencies, and in many cases low onset potentials, for C1 products such as CO, HCOOH and CH₄.^{5,6} Despite these advances, the majority of studies involve catalysts only soluble in organic solvents – a reaction medium that also facilitates high CO₂ concentrations. In contrast, heterogeneous catalysts including metals such as Cu, Au and Ag offer high current densities and stability in water, but rational design of the catalyst sites and development of catalysts remain significant challenges.⁷ This gap underscores the need for catalyst designs that combine the efficiency and selectivity of molecular complexes with the environmental benefits and operational feasibility of heterogeneous systems.

For instance, manganese(i) tricarbonyl complexes are efficient catalysts for the CO₂RR to generate CO with good faradaic yields in protic organic media. However, these complexes generally suffer from poor solubility in water and low recyclability. The heterogenization of catalysts has been a long-standing strategy to mitigate these problems, improving the



stability of molecular catalysts that operate in the homogeneous phase.^{8–11} To develop efficient (high turnover frequency “TOF_{max}” and low overpotential “ η ”) and robust (high turnover number “TON”) [Mn(N⁺N)(CO)₃X]-based catalysts for CO₂ reduction, the immobilization in a heterogeneous fashion at a solid-state interface has been explored.^{12–15} The heterogenization of well-defined catalysts could be prepared by immobilizing them on conducting and semiconducting substrates^{16–18} and porous polymers,¹⁹ among others.²⁰ Some of these strategies to improve the stability of [Mn(N⁺N)(CO)₃X]-based catalysts are included in Fig. 1.^{13,16–19,21–24}

Recently, a number of studies have demonstrated another route using crystalline porous materials: Covalent Organic Frameworks (COFs)²⁵ and Metal–Organic Frameworks (MOFs).²⁶ These frameworks have demonstrated high levels of activity for electrocatalytic CO₂ reduction when integrated with molecular electrocatalysts.^{27–32} The molecular nature of MOF and COF precursors facilitates the incorporation of established molecular catalysts, resulting in frameworks that often outperform their molecular counterparts in terms of efficiency, stability, and durability.^{25,26}

Therefore, the use of MOFs and COFs for catalytic purposes has been extensively explored.^{33–35} The integration of molecular catalytic units into extended frameworks has often resulted in improved efficiency, stability and durability compared to the corresponding molecular analogues.

Among the potential applications of COFs as catalysts, they stand out especially because they have been shown to be excellent precursors or hosts for single-atom catalysts (SACs).^{36,37} COFs minimize the sizes of catalytic nanoparticles and, as a result, expose more reactive sites to facilitate mass/

electron transfer at the interface and to achieve high catalytic reactivity.³⁸ In this context, the potential application of COFs for the CO₂RR has begun to be explored and, as a result, porphyrin, phthalocyanine and bipyridine-based complexes have been integrated as building blocks into COFs or MOFs for the CO₂RR.^{2,29,39–52}

Combining the merits of heterogeneous and homogeneous catalysts, reticular chemistry endows the framework materials with high surface areas, porous structures, and potential for rational tailoring for optimizing CO₂RR performance.^{38,53} Such an approach has significant advantages as it provides materials with well-defined, tunable, catalytic sites in a stable heterogenized form. Recently, a new CO₂ reduction catalyst by loading single-atom centers, {*fac*-Mn(CO)₃S}, into a bipyridyl-inspired COF with 10 times higher catalytic activity than that obtained for the equivalent manganese-based molecular catalyst and access to key catalytic intermediates within a COF matrix has been published.⁵⁴

In this context, herein, we report on the synthesis, characterization and electrochemical evaluation for CO₂ reduction of a novel *fac*-[Mn(bpy)(CO)₃Br]-terpyridine type 2D COF (**Mn@Terpy-COF**). The synthesized materials have been characterized by inductively coupled plasma optical emission spectrometry (ICP-OES), thermogravimetric analysis (TGA), infrared spectroscopy (IR), carbon-13 cross-polarization magic angle spinning nuclear magnetic resonance (¹³C-CP/MAS NMR), scanning electron microscopy-energy dispersive X-ray spectroscopy (SEM-EDX), transmission electron microscopy (TEM), powder X-ray diffraction (PXRD), X-ray photoelectron spectroscopy (XPS) and X-ray absorption spectroscopy (XAS).

Results and discussion

Synthesis and characterization of Terpy-COF

A COF doped with terpyridine moieties, **Terpy-COF**, has been prepared by covalently linking azidoterypyridine **1** with a COF endowed with reactive alkyne moieties, [HC≡C]_{0.17}-TPB-DMTP-COF,⁵⁵ via a copper catalyzed [3 + 2] cycloaddition reaction (Scheme 1). The structural characterization of the new COF has been elucidated and compared with that of the precursor COF endowed with alkyne moieties. The completeness of the reaction between the azides and the alkyne was confirmed by Fourier transform infrared (FT-IR) spectroscopy (Fig. S5†) and Solid-state Cross-Polarization Magic Angle Spinning Carbon-13 Nuclear Magnetic Resonance (¹³C-CP/MAS-NMR) (Fig. 2a). Thus, the diagnostic $\nu(\text{C}=\text{N})$ stretching band at 1618 cm^{–1} remains almost unaltered in the FT-IR spectra after the cycloaddition reaction.⁵⁶ On the other hand, the ¹³C-CP/MAS-NMR spectrum evidences the efficient transformation as revealed by the disappearance of the signals corresponding to the alkyne group at 79 and 72 ppm (Fig. 2a). The remaining aliphatic signal at 54 ppm is attributed to the methoxy groups of the COF skeleton. The introduction of the terpyridine fragment was also confirmed by analyzing the aromatic region. First, quaternary pyridine C=N carbons were localized together with the rest of the C=N carbons of the COF in the signal at 154 ppm, which appears broader than in the starting material [HC≡C]_{0.17}-TPB-

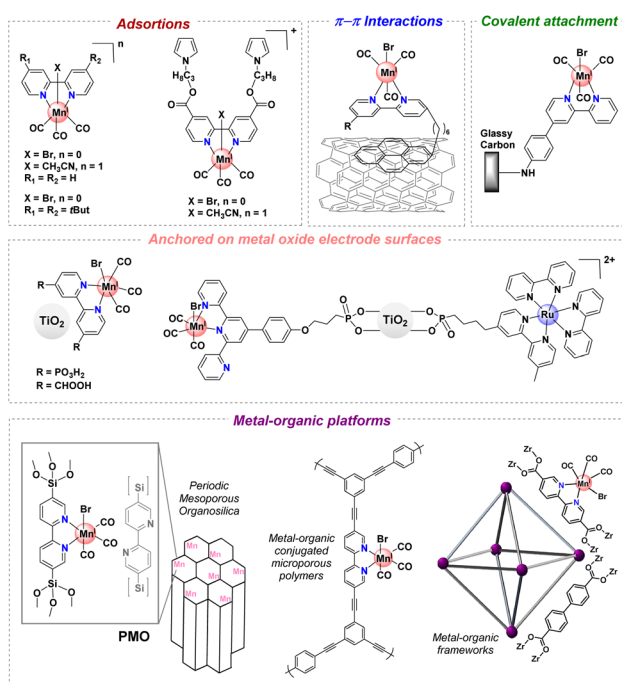
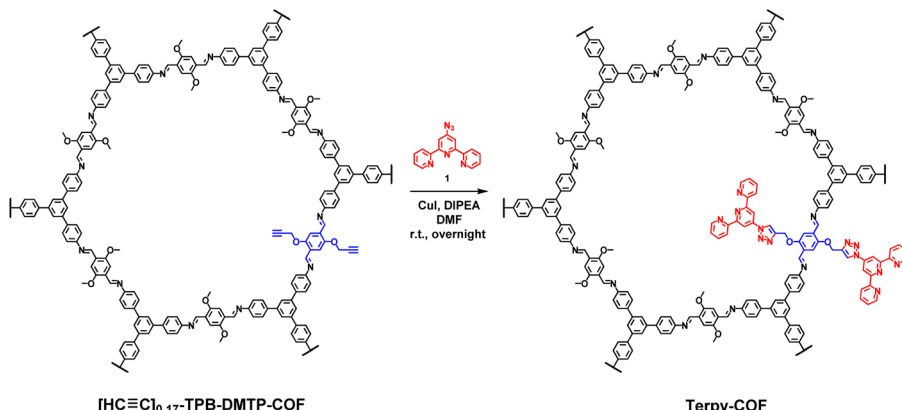


Fig. 1 Selected immobilization strategies employed in manganese tricarbonyl complexes containing polypyridyl moieties.



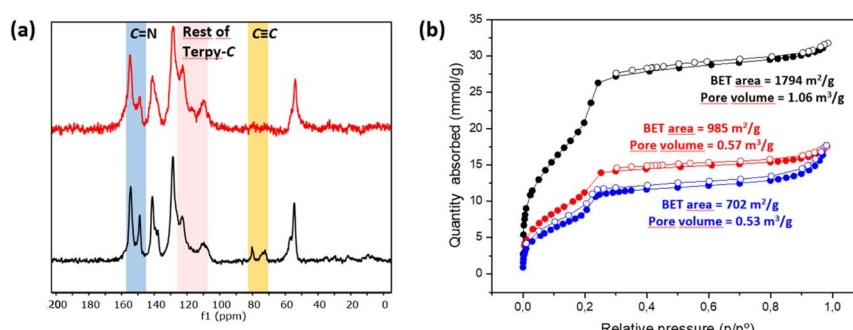
Scheme 1 Synthesis of **Terpy-COF** from $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$.

DMTP-COF (Fig. S3†). The introduction of the pyridine fragment was not only evidenced by the widening of the signal corresponding to $\text{C}=\text{N}$, but also by the appearance of a new signal (observed as a shoulder) at 117 ppm and enhancement of the relative intensities of signals at 122 and 109 ppm (Fig. S3†), which corresponds to chemical shifts of terpyridine $\text{C}^3(5')$, $\text{C}^3(3'')$ and $\text{C}^5(5'')$ carbons (Fig. S2 and S4†). The chemical shifts of the rest of the anisochronous nuclei correspond well with those expected for the structure of the COF core, those of the triazole rings and that of the terpyridine system. It should be highlighted that neither the precursor $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$ nor the final cycloaddition product exhibited FTIR or ^{13}C -CP-MAS-NMR signals of the starting materials corresponding to aldehyde or amine functionalities. The introduction of the terpyridine fragment into the pore resulted, as expected, in a decrease of the Brunauer–Emmett–Teller (BET) surface area of **Terpy-COF** ($985 \text{ m}^2 \text{ g}^{-1}$) in comparison with that of the starting material $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$ ($1794 \text{ m}^2 \text{ g}^{-1}$) (Fig. 2b). N_2 sorption isotherms at 77 K also revealed a reduction of pore volume (from $1.06 \text{ cm}^3 \text{ g}^{-1}$ to $0.57 \text{ cm}^3 \text{ g}^{-1}$ at 0.95 p/p^0) and pore size distribution, calculated by NLDFT (Fig. S7†). These data, together with the IV-type isotherm obtained, evidence the mesoporous nature of **Terpy-COF**.⁵⁵

Morphological characterization of **Terpy-COF** and $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$ has been carried out by using

Scanning Electron Microscopy (SEM). SEM micrographs revealed that the mild conditions of the copper catalyzed click reaction did not significantly alter the morphology of the COF and therefore $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$ and **Terpy-COF** exhibit almost identical rodlike morphologies (Fig. S8 and S9†).

X-ray powder diffraction (PXRD) analysis of **Terpy-COF** revealed a high degree of crystallinity, with six distinctive diffraction peaks which almost replicate those previously observed for $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$ with an eclipsed AA stacking (Fig. 3a).⁵⁵ The most intense diffraction maximum, at 2.90° , had an excellent FWHM value of 0.2303° and corresponds to the (100) facet.⁵⁷ The other peaks appeared at 4.95° , 5.70° , 7.54° , 9.83° and 25.20° and can be assigned to (110) , (200) , (210) , (220) and (001) reflections according to that previously reported for $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$.⁵⁵ With the aim to confirm the macromolecular crystal structure of **Terpy-COF**, we have conducted an extensive set of DFT-based calculations using a combination of Gaussian 16 C.01 (ref. 58) and QUANTUM ESPRESSO⁵⁹ software packages to determine the crystalline structures of **TPB-DMTP-COF** and **Terpy-COF** in this study (further computational details in the ESI†). The simulations revealed purely hexagonal symmetry for all crystal structures, with monolayer lattice parameters ranging from 37.57 to 37.72 Å (Fig. 3). We studied the preferred stacking configuration for **TPB-DMTP-COF** and **Terpy-COF**. In both cases, the AA

Fig. 2 Comparative (a) ^{13}C CP/MAS NMR patterns of $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$ (black) and **Terpy-COF** (red), and (b) N_2 (77 K) sorption isotherms of $[\text{HC}\equiv\text{C}]_{0.17}\text{-TPB-DMTP-COF}$ (black), **Terpy-COF** (red) and $\text{Mn}@\text{Terpy-COF}$ (blue).

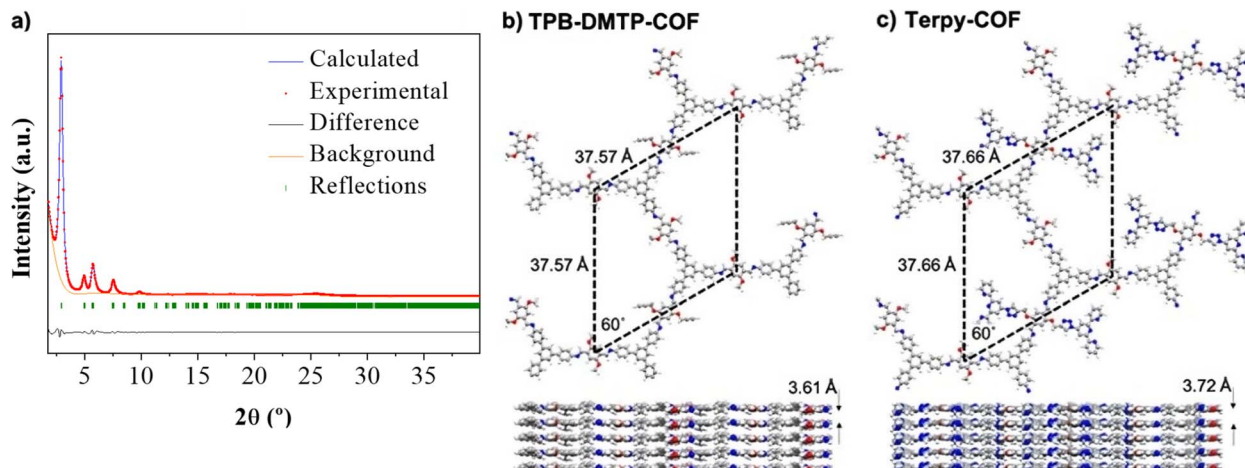


Fig. 3 (a) Comparison between the experimental (red) and the theoretically simulated (blue) diffractograms obtained from the resulting optimized structures for $[\text{HC}\equiv\text{Cl}]_{0.17}$ -TPB-DMTP-COF and Terpy-COF (practically identical). Experimental diffractogram has been refined (Rietveld refinement) with parameters $R_{\text{wp}} = 4.54\%$ and $R^2 = 3.08\%$. (b) and (c) Top and side pictorial views of the structures, indicating the resulting optimized unit cell in all cases, for (b) $[\text{HC}\equiv\text{Cl}]_{0.17}$ -TPB-DMTP-COF and (c) Terpy-COF. The preferential interlaminar stacking for $[\text{HC}\equiv\text{Cl}]_{0.17}$ -TPB-DMTP-COF and Terpy-COF is the eclipsed configuration, showing a very similar interlayer distance.

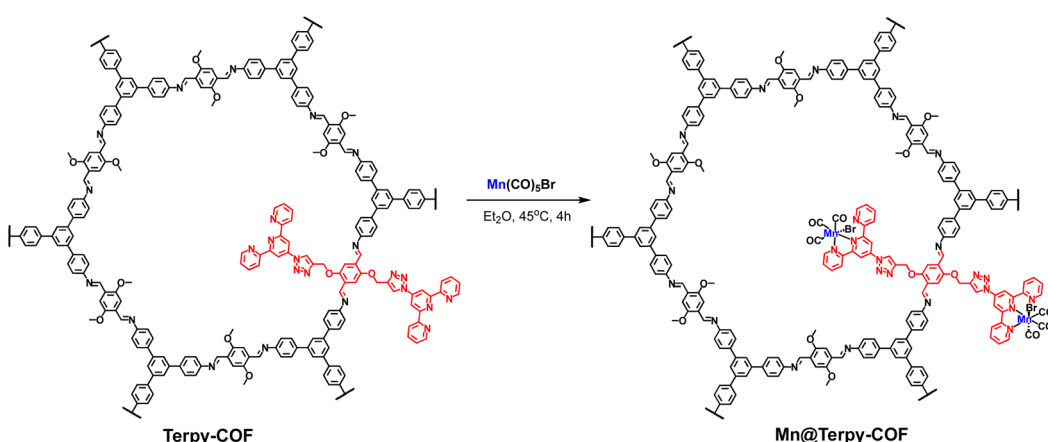
eclipsed configuration was found to be preferential, with very similar interlayer distances of 3.61 and 3.72 Å (Fig. 3). The theoretical diffractograms based on optimized crystal structures (Fig. 3c, practically identical for both) showed excellent agreement with experimental ones. The diffraction peaks in the simulated PXRD pattern are located at 2.803° , 4.821° , 5.605° , 7.287° , 13.677° and 24.776° in excellent agreement with the experimental evidence, as shown by Rietveld refinement (Fig. 3a). The simulated PXRD pattern even reproduced high-angle features, confirming the validity of structures obtained from simultaneous structure + cell DFT geometrical optimization.

Synthesis of Mn@Terpy-COF

The post-functionalization of **Terpy-COF** was performed by a direct reaction with $\text{Mn}(\text{CO})_5\text{Br}$ at 45°C in diethyl ether for 4 h, forming **Mn@Terpy-COF** as a brown powder (Scheme 2).

The amount of Mn measured by inductively coupled plasma-optical emission spectroscopy (ICP-OES) results in an inclusion of 38 mg of Mn per gram of **Terpy-COF**, which accounted for $\sim 75\%$ of the COF terpyridines coordinated to Mn after de metalation.

The incorporation of $\{\text{fac-}\text{Mn}(\text{CO})_3\text{Br}\}$ fragments into **Terpy-COF** was confirmed by ATR-FT-IR spectroscopy, showing the presence of two CO stretching frequencies at 2020 cm^{-1} and a shoulder at 1919 cm^{-1} (Fig. 4). The frequency at 2020 cm^{-1} matches the $\nu_{(\text{CO})}$ of the symmetric CO vibration of the facial tricarbonyl moiety and the unresolved broad band at 1919 cm^{-1} corresponds to the symmetric and asymmetric equatorial CO vibration. This pattern is analogous to that shown by the molecular $\{\text{fac-}\text{Mn}^{(\text{I})}(\kappa^2\text{-terp})(\text{CO})_3\text{Br}\}$ complex reported by Kubiak and col.⁶⁰ UV-vis spectroscopy (Fig. S6†), thermogravimetric analysis (Fig. S14†), PXRD (Fig. S15†) and porosity and surface area measurements (Fig. 2 and S7†) were also conducted



Scheme 2 Synthesis of **Mn@Terpy-COF** from **Terpy-COF**.

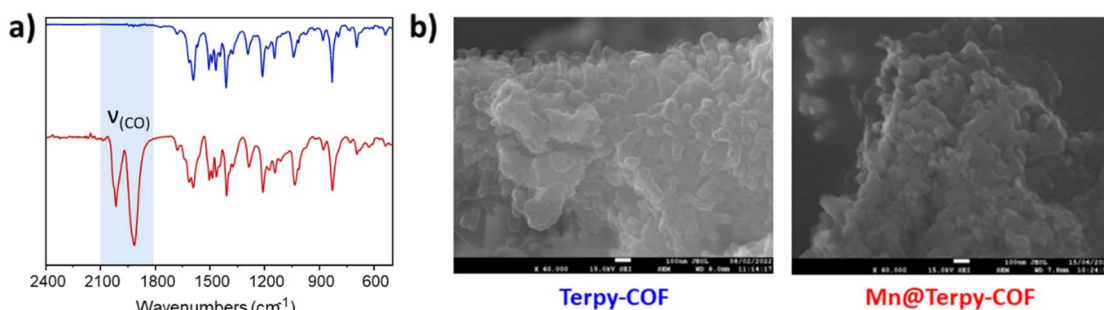


Fig. 4 (a) ATR spectra $\nu_{(\text{CO})}$ at 2025 and 1915 (sh) and (b) SEM of Terpy-COF (blue) and Mn@Terpy-COF (red).

to characterize the material. Powder X-ray diffraction analysis confirmed that the crystallinity of the COF was preserved after the metalation process (Fig. S15[†]). As anticipated, the porosity and surface area decreased following the incorporation of the metal within the pore structure, yielding a BET surface area of $702 \text{ m}^2 \text{ g}^{-1}$ (Fig. 2) and a pore volume of $0.53 \text{ cm}^3 \text{ g}^{-1}$ at a relative pressure of $0.95 (p/p^0)$, (Fig. S7[†]). These observations align with the expected reduction in porosity and pore volume upon introduction of Mn and CO moieties into the pore network, confirming the impact of these modifications on the material's structural properties.

An extensive X-ray photoemission spectroscopy (XPS) characterization study was also performed to confirm the structure assigned to **Terpy-COF** and **Mn@Terpy-COF** (Fig. S16–19[†]). Analyzing first the starting material **Terpy-COF**, the N 1s emission peak was clearly identified and showed different components, so it was deconvoluted for a more detailed analysis. The two main components, labeled A and B, were identified at 398.5 and 399.9 eV, and an additional minor component, labeled C, was observed at 401.5 eV (Fig. 5). The main component,

corresponding to a binding energy of 398.5 eV, was attributed to $-\text{N}=\text{}$ of the imine and pyridine groups. The peaks detected at 399.9 and 401.5 eV were assigned to imidazole nitrogen atoms. Specifically, the lower binding energy signal was assigned to N, labeled 2 and 3, while the less intense crest was assigned to the nitrogen directly attached to the terpyridine fragment, labeled 1 (Fig. 5). These relative positions of binding energy and intensity percentages agree with our predictions and with data from the literature. The addition of manganese to the COF caused a noticeable change in the N 1s peak area percentages and a slight displacement of the component with the highest binding energy (C) as can be seen in Fig. 5. The observed percentage change of the peak is related to the shift of the binding energy of some of the terpyridine nitrogen atoms from the initial imine region (398.5 eV) to the imidazole 2 and 3 zones at 399.9 eV, where the signal is broader and has a higher relative intensity. This result is consistent with what is expected for the structure of **Mn@Terpy-COF** because the nitrogens coordinated with manganese have given charge to the metal ion and appear more oxidized. On the other hand, the signal originating from

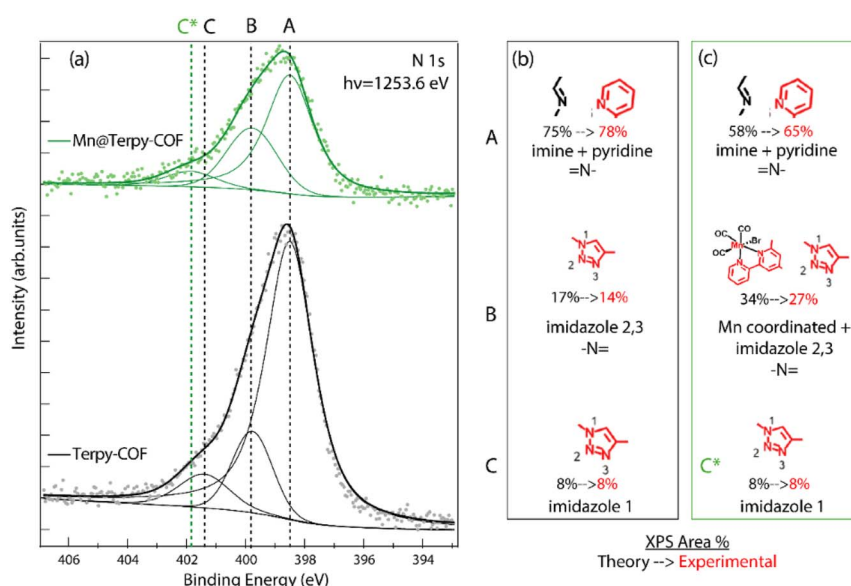


Fig. 5 (a) XPS spectra of the 1 N 1s peak region of Terpy-COF (black) and Mn@Terpy-COF (green). Dots correspond to experimental points and continuous lines to the deconvolution in different components and the background. Vertical dotted lines indicate the binding energy positions of the components. Panels (b) and (c) show the origin of each N 1s component. Data obtained with a Mg K α (1253.6 eV) photon source.



Mn 2p in the **Mn@Terpy-COF** XPS spectrum shows two components (Fig. S17†). The major component (642 eV, 67%) is assigned to manganese coordinated to terpyridine. The minor component (646 eV, 33%) could be related to manganese in a different environment, but we discard this interpretation considering the high value of the binding energy. Instead, we attribute it to a shake-up satellite observed in other Mn coordination complexes about 4 eV higher than the main Mn 2p photoelectron line.⁶¹ In both compounds, **Terpy-COF** and **Mn@Terpy-COF**, the C 1s signals are similar (Fig. S18†), with the major component attributed to aromatic sp^2 carbons (A label). Minor components were also identified at higher binding energies and were assigned to sp^2 carbon directly bonded to nitrogen and oxygen (B label).

Electrocatalytic CO_2 reduction in water with **Mn@Terpy-COF**

Preparation and characterization of the **Mn@Terpy-COF|NT thin film.** In order to study the electrocatalytic activity of **Mn@Terpy-COF**, a film of the material was deposited on the working electrode (glassy carbon). The immobilization was achieved by mixing the catalyst powder with Multi-Walled Carbon Nanotubes (NT) in CH_3CN and Nafion (5% water) (1 mg : 1 mg : 100 μL : 100 μL) by sonication (15 minutes) to obtain the **Mn@Terpy-COF|NT|ink** catalyst ink. 10 μL of the dispersion were added dropwise onto the electrode surface and allowed to dry in the dark and under ambient conditions to yield **Mn@Terpy-COF|NT** (Fig. S13) (see ESI for details†).

To gain insight into the COF matrix and the electronic structure and coordination environment of the Mn centers in the catalyst ink, an XPS study was performed. The XPS analysis of **Mn@Terpy-COF|NT|ink** suspension and **Mn@Terpy-COF|NT** showed, as previously described, C 1s signals but with new significantly shifted components, assigned to the carbon atoms bonded to fluorine atoms present in the Nafion polymer used (component C in Fig. S19†). In addition, a new small component of C sp^2 was observed in the **Mn@Terpy-COF|NT** XPS spectrum, in a position similar to the benzene carbon atoms of the previously described **Terpy-COF** and **Mn@Terpy-COF** (label A* in Fig. S19†). The similarity in the binding energy position of

the previous A component and this new A* component associated with the carbon nanotubes makes it difficult to deconvolute this into two components, but the increase in the C 1s A/B component ratio is sufficient to conclude that a new component is present. In all these new materials, the signals of N 1s were barely noticeable and the signal to noise ratio was low (not shown). In the case of Mn, the strong signal of fluorine (F 1s and KLL Auger F) is superimposed on the Mn 2p peaks (Fig. S16†), making it difficult to observe and analyze the Mn state.

Catalytic activity of **Mn@Terpy-COF|NT for CO_2 reduction in water.** To evaluate the electrochemical behavior and the catalytic CO_2 reduction performances of **Mn@Terpy-COF|NT** ($[\text{Mn}]_{\text{total}} = 850 \text{ nmol cm}^{-2}$), cyclic voltammetry (CV) and controlled-potential electrolysis (CPE) measurements were carried out in $\text{K}_2\text{SO}_4/\text{K}_2\text{B}_4\text{O}_7$ electrolyte under Ar (pH 8.2) and CO_2 (pH 7.6) atmospheres, respectively.

We started the characterization by performing CV of **Mn@Terpy-COF|NT** under Ar and CO_2 atmospheres in $\text{K}_2\text{SO}_4/\text{K}_2\text{B}_4\text{O}_7$ (Fig. 6a). Under Ar (pH 8.2), CV showed an irreversible cathodic wave, suggesting a catalytic hydrogen evolution reaction (HER) that was confirmed by the detection of H_2 after control potential electrolysis (see below). When the solution was saturated with CO_2 (pH 7.4), the onset of the catalytic wave shifted 360 mV to more positive potentials and the catalytic current increased 3.5-fold ($j_{\text{CO}_2}/j_{\text{Ar}}$ at -1.3 V).

To estimate the neat CO_2 reduction activity of **Mn@Terpy-COF|NT**, CPE measurements of the thin film on carbon paper (CP) electrode as a current collector were recorded in aqueous $\text{K}_2\text{SO}_4/\text{K}_2\text{B}_4\text{O}_7$ under CO_2 (pH 7.4) (see the Experimental section for details). CPE measurements were carried out to benchmark the CO_2 reduction activity selectivity and the stability of the catalyst **Mn@Terpy-COF|NT** (Fig. 6b and c). The generated products were analyzed by gas chromatography and $^1\text{H-NMR}$. CPE measurements at different applied potentials (from -1.1 to -1.8 V vs. SCE) of **Mn@Terpy-COF|NT** ($[\text{Mn}]_{\text{total}} = 850 \text{ nmol cm}^{-2}$) showed current densities from -0.06 to -17 mA cm^{-2} . The generated products were quantified (nmol cm^{-2}) at different overpotentials (Fig. 6c), and the selectivity of the CO_2 reduction reaction shifted to H_2 with the increase of the overpotential. This can be attributed to the protonation of the Mn

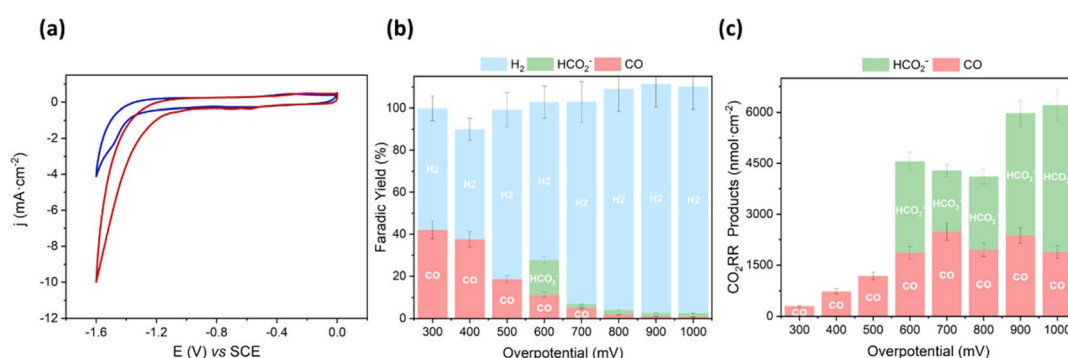


Fig. 6 (a) CV on GC of **Mn@Terpy-COF|NT** ($[\text{Mn}]_{\text{total}} = 850 \text{ nmol cm}^{-2}$) in water ($\text{K}_2\text{SO}_4/\text{K}_2\text{B}_4\text{O}_7$) under Ar (blue, pH 8.2) and CO_2 (red, pH 7.4) at 10 mV s^{-1} . (b) faradaic yield and (c) generated CO_2 RR products for CPEs on Carbon Paper (CP) of **Mn@Terpy-COF|NT** ($[\text{Mn}]_{\text{total}} = 850 \text{ nmol cm}^{-2}$) at different overpotentials (η). Generated products: HCOO^- (green), CO (pink) and H_2 (blue).



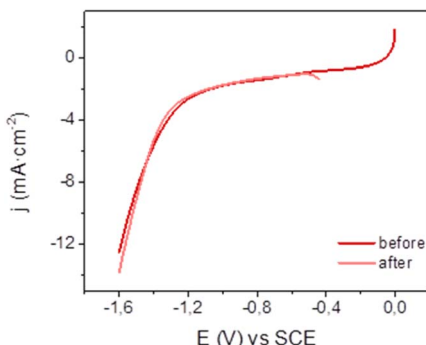


Fig. 7 Linear sweep voltammetry of **Mn@Terpy-COF|NT** ($[\text{Mn}]_{\text{total}} = 850 \mu\text{mol cm}^{-2}$) on CP at 10 mV s^{-1} before and after CPE at -1.3 V vs. SCE in water $\text{K}_2\text{SO}_4/\text{K}_2\text{B}_4\text{O}_7$ under CO_2 .

centers forming the Mn-H species, which further protonates to produce H_2 , and to the exacerbation of the background reaction of H_2 formation by the carbon nanotubes as previously proposed.⁵⁴ The formation of CO and HCO_2^- was observed from a low η of 300 mV (Fig. 6b). At $\eta = 600 \text{ mV}$, **Mn@Terpy-COF|NT** generated 2000 and $2700 \text{ nmol cm}^{-2}$ of CO and HCO_2^- , respectively. Although the CO obtained was insufficient to be detectable by online GC-MS, CPE experiments with $^{13}\text{CO}_2$ indicated that formate comes from CO_2 (Fig. S21†).

Mn@Terpy-COF|NT electrodes showed good stability under catalytic conditions. A superimposition of linear sweep voltammetry (LSV) traces recorded before and after CPE indicates negligible catalyst decomposition (Fig. 7). In general, no significant changes were observed by XPS and SEM analysis before and after electrocatalysis in water under CO_2 . EDX showed the homogeneous distribution of Mn before and after catalysis, further highlighting the robustness of **Mn@Terpy-COF|NT**.

To delve into the microscopical morphologies of **Mn@Terpy-COF|NT** before and after its test as a CO_2 reduction catalyst, different SEM-EDX spectra were recorded. Deposition of the **Mn@Terpy-COF|NT/Nafion** suspension over the electrode resulted in a heterogeneous surface, where **Mn@Terpy-COF** is embedded in different parts of the carbon-based electrode matrix (Fig. S12†). This microstructure was maintained unchanged after the electrode's testing process, which is promising for a hypothetical long-time use of this dispositive as a CO_2 reductive system (Fig. S13†). In addition, XPS spectra of the electrodes after their test as catalysts were recorded, showing no significant change in the C 1s core level with respect to the starting material **Mn@Terpy-COF|NT** (Fig. S19†). However, the catalyst exhibited altered selectivity in a subsequent catalytic run, producing mainly H_2 .

Spectroelectrochemistry. SEC experiments provided further evidence of the intermediates that can be formed at different potentials in catalysis. To shed some light on the catalytic activity, we have examined the material by IR-SEC, a technique that enables redox changes to be monitored through changes in the ATR spectra as a function of potential and time. Indeed, ATR-IR-SEC (Fig. 8) has been previously employed to obtain

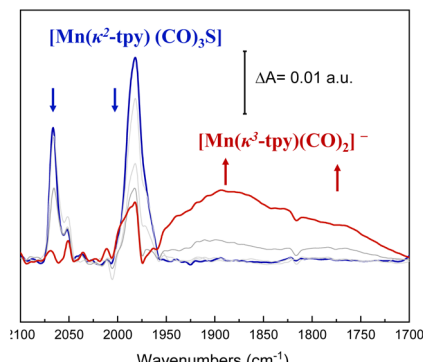


Fig. 8 ATR-SEC spectra of **Mn@Terpy-COF** in $0.2 \text{ M TBAPF}_6/\text{CH}_3\text{CN}$ under Ar saturation. Red curve describes the final spectra, whereas grey lines refer to selected intermediate spectra. Blue curve is the starting catalyst.

structural information on the catalytically relevant species of previous reported Mn single sites in **COF_{bpyMn}|NT**.⁵⁴ Considering these advantages, ATR-IR-SEC of **Mn@Terpy-COF|NT** was performed in CH_3CN (0.2 M TBAPF_6) by a stepwise decrease of potential from 0.4 V to -1.9 V vs. SCE under Ar to monitor the reduction of the precatalyst.

Before performing ATR-IR-SEC, we performed CV of **Mn@Terpy-COF|NT** in dry CH_3CN (0.2 M TBAPF_6) under Ar. The CV showed several irreversible cathodic waves in the range from -0.1 V to -1.5 V vs. SCE. Besides, during the forward scan, three reoxidation processes were observed at -1.05 , -0.5 and 0.6 V vs. SCE, respectively. These redox transformations are similar to those observed for **Mn@Terpy** molecular complexes previously reported,⁶⁰ suggesting an electronic similarity for the $\text{Mn}(\kappa^2\text{-tpy})(\text{CO})_3\text{S}$ coordination between **Mn@Terpy-COF|NT** and **Mn@Terpy** as ligands.

With these considerations, ATR-IR-SEC was carried out to gain structural insights into the electrochemical activation of the **Mn@Terpy-COF|NT** precatalyst. At open-circuit potential (E_{OCP}), the ATR-FT-IR of **Mn@Terpy-COF|NT** (Fig. 8) showed a $\nu_{(\text{CO})}$ band at 2063 cm^{-1} and a broad feature at 1982 cm^{-1} , and the IR pattern corresponds to a *fac* arrangement within the core structure of the COF, which also matched well with the pattern observed for the complex **Mn@Terpy**. This profile also agrees with the $\nu_{(\text{CO})}$ bands of several reported immobilized facial tricarbonyl manganese polypyridine complexes² and with that previously observed for **COF_{bpyMn}|NT**.⁵⁴ When the potential was stepwise decreased from 0.4 to -1.6 V (vs. SCE), the CO vibrations show an increase in intensities at lower energies with the evolution of the redox potential to more negative values that could be associated with a redox transformation. The ATR-IR-SEC spectra showed a decrease of the precatalyst signals with the concomitant increase of a series of bands at 1937 , 1982 , 1893 , 1827 and 1750 cm^{-1} indicating the formation of new species. We hypothesized that after the reduction of **Mn@Terpy-COF|NT**, two different intermediates contributed to the spectra, the dimer and the anionic species (Fig. 8), similar to the intermediates detected for the molecular **Mn@Terpy**.⁶⁰

Computed CO₂ reduction reaction (CO₂RR) mechanism. As a proof-of-concept, a large set of *ab initio* theoretical calculations were carried out to obtain a viable mechanism for the CO₂RR towards the CO evolution as the reaction product (see further details in the ESI†). Formate evolution has not been theoretically analyzed since its faradaic yield (FY) is significantly lower than that of CO for all voltages applied.

Fig. 9 shows the computed Gibbs free energy diagram for the proposed mechanism, for which all the structures were fully optimized in water solution, for externally applied voltages *vs.* SCE mimicking the voltage range explored in the experiments: *U* = 0, -1.1 and -1.8 V.

Catalyst activation. The first step (1) depicts the organometallic unit in the COF system, in which the Mn atom has an oxidation state of +1 and a six-fold coordination with a pseudo-octahedral structure. After two sequential one-electron reductions, two electrons are transferred to the Mn center for the catalyst activation resulting in a net free energy change at *U* = 0 V of +2.01 eV. The first one-electron reduction leads to barrierless Br⁻ release towards the formation of (2). After that step, (2) can be easily electro-reduced into (3), with a net free energy gain of +1.42 eV. After these two electro-reductions, the two additional electrons are located in the d_{xy}(Mn) orbitals.⁶²

CO₂ adsorption and protonation. Once the active species (3) are formed, CO₂ can anchor in the Mn vacant site with a computed low energy barrier of 0.26 eV (4), to finally form the Mn–CO₂ intermediate (5) with a net free energy change of around +0.07 eV (endergonic) at *U* = 0 V, and a bond-length Mn–C of around 3.16 Å. After this step, the reaction proceeds by the protonation of the CO₂ ligand. This process has been modelled by the explicit inclusion of a (H₂O)₅ cluster in a chair

configuration as previously reported,⁶³ which leads the water cluster to coordinate with CO₂ to yield (6). In this step a transfer of a proton is produced from the water cluster to the C–O unit to form (7), followed by the subsequent release of OH⁻(H₂O)₄ with a slight net free energy change of +0.03 eV (endergonic) at *U* = 0 V.

CO release and catalyst regeneration. An explicit (H₂O)₅ water cluster can coordinate again with (7) to exergonically form (8) ($\Delta G = -0.23$ eV), capturing an OH⁻ by subsequent release of OH⁻(H₂O)₅ with a slight net free energy loss of -0.15 eV, leading to the positively charged Mn(CO)₃CO⁺ unit (10) by surmounting an associated energy barrier of 0.41 eV (9). From this step, the CO release occurs spontaneously by a synergistic electro-reduction with an associated net free energy gain of +1.18 eV at *U* = 0 V to yield (11). Finally, in order to regenerate the catalyst, (11) can be electro-reduced ($\Delta G = +1.16$ eV) to regenerate active species (3).

Fig. 9 reveals that at *U* = 0 V *vs.* SCE, the mechanism is far from being thermodynamically viable, particularly due to the sub-reactions involving the one-electron reductions. Starting from the *U* = 0 V profile, we have computed the Gibbs free energy paths at *U* = -1.1 and -1.8 V *vs.* SCE (see Fig. 9), which correspond to voltage range experimentally explored. Interestingly, at *U* = -1.1 V the entire reaction becomes exergonic by approximately -0.11 eV. The reaction is limited by step 9, which involves C–O bond cleavage followed by OH⁻ abstraction, with a net energy barrier of 0.71 eV. This result in excellent agreement with the experimental evidence. Finally, at *U* = -1.8 V the whole reaction is clearly exergonic by <-3 eV. For this case, once again, the C–O bond cleavage and subsequent OH⁻ abstraction is the reaction limiting step, since it is not linked to the

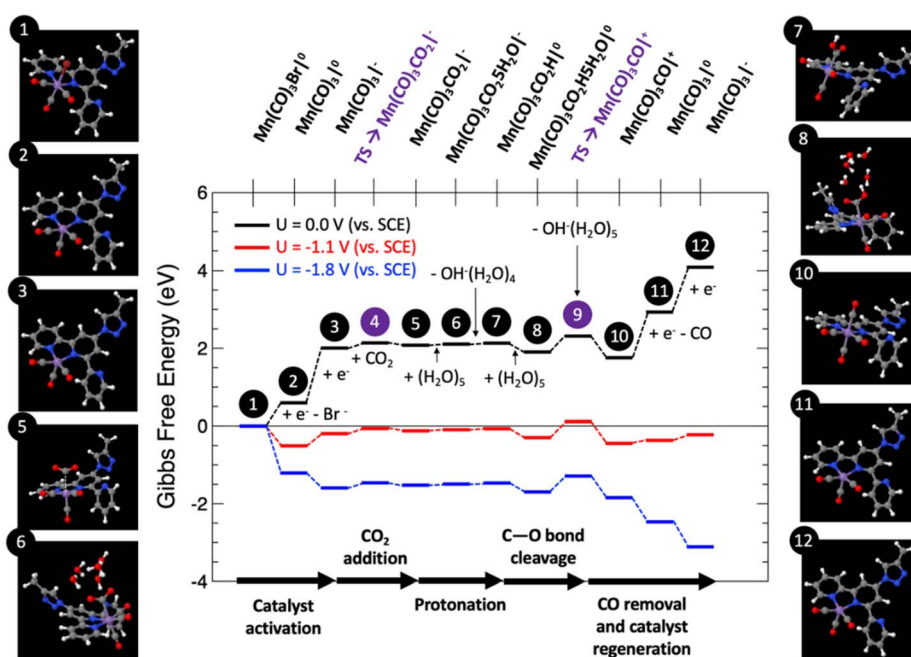


Fig. 9 Computed Gibbs free energy diagram (in eV) for the CO₂RR towards CO evolution at different applied voltages of *U* = 0 (black path), -1.1 (red path) and -1.8 V (blue path) *vs.* SCE. Calculated geometries of all the intermediates are shown numbered according to the reaction path steps, indicating each sub-reaction of the mechanistic path.



externally applied potential, yielding a net barrier of 0.41 eV, 0.3 eV lower than for the $U = -1.1$ V case, thus justifying the CO production increase. Thus, the atomistic mechanism theoretically proposed clearly predicts the activation of the CO₂ reduction/CO evolution channel for the voltage window experimentally explored.

Conclusions

A novel **Mn@Terpy-COF** has been synthesized from **[HC≡C]_{0.17}-TPB-DMTP-COF** following a pathway that involves the introduction of a terpyridine fragment into the macromolecular material by a CuAAC reaction, followed by the complexation of the ligand with manganese. This new **Mn@Terpy-COF** has been shown to be an active catalyst for the electrochemical reduction of CO₂ in water. The spectroscopic characterization of **Mn@Terpy-COF** by IR shows that the Mn center has an a fac coordination environment, homologous to the molecular complex **Mn@Terpy** and the **COF_{bpyMn}**. XPS analysis also evidences the formation of **Mn@Terpy-COF**. Positive shifts in the onset potentials and increases in current are observed in the CV of the frameworks when the atmosphere is changed from Ar to CO₂. **Mn@Terpy-COF** reduces CO₂ to CO with a FY of 42% at an overpotential of only 300 mV. In addition, a FY of 16% was observed for formate production at an overpotential of 600 mV.

Data availability

The data supporting this article have been included as part of the ESI.†

Author contributions

Conceptualization: J. L. S., F. Z., and J. L.-F. investigation: E. G., G. C. D. B., M. V., S. R., M. M. F., and E. S. methodology: E. G., G. C. D. B., S. R., J. I. M., and E. S. writing – original draft: E. G., G. C. D. B., M. M. F., J. I. M., and E. S. writing – review and editing: E. G., M. M. F., E. S., F. Z., J. I. M. E. G. M., J. L.-F., and J. L. S. visualization: all the authors. Project administration and supervision: J. I. M., E. G. M., F. Z., J. L.-F., and J. L. S.

Conflicts of interest

There are no conflicts to declare.

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