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Designing molecular and two-dimensional metallocporphyrin catalysts for the electrochemical CO_2 reduction reaction[†]

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The electrochemical CO_2 reduction reaction (e CO_2 R) is an important route toward the sustainable conversion of CO_2 to value-added chemicals. However, developing efficient catalysts with high selectivity and stability remains challenging. Metallocporphyrins (M-PORs) represent an attractive class of molecular catalysts because their structural framework offers a unique combination of tunability of the peripheral ligands, flexibility of the metal centre, and versatility of the oxidation state of the metal. These properties can be exploited to tailor the catalytic properties of M-PORs for the e CO_2 R. Here, we present a comprehensive computational study using density functional theory to systematically explore M-POR catalysts with varying metal centers (Ni, Fe, Cu, Co), oxidation states, and anchoring ligands, aimed at enhancing the selective production of the C_1 products (carbon monoxide and formic acid). Thermodynamic and electrochemical stability analyses revealed neutral M-PORs to be significantly more stable than their charged counterparts, providing crucial guidelines for catalyst design. A mechanistic analysis of reaction pathways—proton-coupled electron transfer (PCET) versus sequential proton and electron transfer (PT-ET)—identified PCET as highly favourable, with predominant selectivity towards formic acid. This study identifies Fe-POR as the one showing superior catalytic performance. Importantly, integrating these optimal molecular catalysts into two-dimensional (2D) carbonaceous frameworks led to further enhancement of catalytic performance, identifying 2D Fe-POR as a highly promising material for selective C_1 product formation, thus providing a rational framework for designing effective molecular-to-framework electrocatalysts for the e CO_2 R.

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1. Introduction

Transforming carbon dioxide (CO_2) into valuable chemicals, fuels, and materials holds substantial societal and scientific significance. This process has the potential to mitigate the effects of climate change and advance the use of sustainable energy.^{1,2} Electrochemical CO_2 reduction (CO_2 R) is considered a promising method thanks to its technical sophistication, maturity, and economic viability, which is why it has gained significant attention from both academic and industrial sectors.^{3–5} CO_2 R can produce high-energy and high-value one-carbon (C_1) products such as carbon monoxide (CO), methane (CH_4), formic acid (HCOOH), and methanol (CH_3OH), and

multi-carbon (C_2+) products like ethylene (CH_2CH_2), ethanol ($\text{CH}_3\text{CH}_2\text{OH}$), acetic acid (CH_3COOH), and n -propanol ($\text{CH}_3\text{CH}_2\text{CH}_2\text{OH}$).^{6–8} These molecules can then be used as feedstock for the synthesis of long-chain hydrocarbon fuels.^{9–11} CO_2 R into C_2+ products stands as the pinnacle achievement in electrochemical synthesis.^{12,13} Still, the endeavour to develop efficient catalysts that are capable of high product selectivity remains one of the critical technological hurdles for effective CO_2 R.¹⁴

Metallocporphyrin (M-POR) has attracted significant interest.¹⁵ The structure of this molecular catalyst consists of a central metal atom surrounded by a cyclic carbon–nitrogen ring composed of four modified pyrrole subunits. The M-POR structural framework offers a unique combination of tunability of the peripheral ligands, flexibility of the metal centre, and versatility of the oxidation state of the metal, which can be exploited to tailor the catalytic properties of M-PORs for CO_2 R¹⁶ and other electrocatalytic reactions such as the oxygen evolution reaction.¹⁷ The synthesis of M-PORs using Earth-abundant elements such as iron^{18,19} or nickel²⁰ reduces the synthetic cost of M-PORs compared to rare and precious metals, such as gold²¹ and silver,²² and thus

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provides a more sustainable alternative to develop a long-term ecologically and economically sustainable process.^{23–25}

In M-POR complexes, the metal centre can also adopt various oxidation states, specifically neutral [M-POR]⁰, single reduced [M-POR][−], or doubly reduced [M-POR]^{2−}, which affects the competition between CO₂R and the hydrogen evolution reaction (HER, H⁺ + e[−] → 1/2H₂), at the cathode.²⁶ A density functional theory (DFT) study by Masood *et al.* found that lowering the nucleophilicity of the metal centre enhances CO₂ binding relative to proton binding, promoting selectivity for CO₂R.²⁷ Specifically, [Fe-POR][−] and [Co-POR][−] showed lower electron density at the metal centre, making them less nucleophilic and more selective for CO₂ adsorption and CO formation. In contrast, nucleophilic centres, such as in [RhP]⁰ and [IrP]⁰, favour proton binding, leading to the HER. Fe-POR shows a strong potential in CO₂R applications. In the literature, one of the early studies on the iron tetraphenyl porphyrin complex FeTPP was carried out in the absence of proton donors using NEt₄ClO₄/DMF electrolyte.²⁸ These aprotic conditions showed a moderate yield for CO, which was attributed to the [Fe(TPP)]^{2−} species. The catalyst experienced rapid deactivation during the reaction, primarily due to chemical changes in the ligand, such as carboxylation and hydrogenation. These transformations negatively impacted the catalytic performance, resulting in low FE for CO production. Additionally, the stability of the catalyst was significantly compromised, further limiting its effectiveness in the reaction. In the 1990s, it was discovered that adding mono- (Li⁺, Na⁺) or divalent Lewis acids (Mg²⁺, Ca²⁺, Ba²⁺) to the electrolyte improved the activity and stability of FeTPP catalysts.^{29,30} The enhancement followed the order Mg²⁺ ≈ Ca²⁺ > Ba²⁺ > Li⁺ > Na⁺. Long-term electrolysis revealed CO as the major product and formate as a by-product, with the product ratio dependent on the Lewis acid used.

Cyclic voltammetry studies showed that in the presence of CO₂ and a Lewis acid like Mg²⁺, a significant catalytic current appeared at the potential of the [Fe(TPP)]^{2−} species. This indicated that catalysis was initiated during the final electron transfer step. The metal center of [Fe(TPP)]^{2−} underwent nucleophilic attack by CO₂, forming an intermediate Fe(II)-CO₂ adduct. Mg²⁺ facilitated the reaction by forming an ion pair with negatively charged oxygen atoms in the adduct, which pulled an electron pair from Fe(0) to CO₂, weakening C-O bonds and promoting CO release. This demonstrates that Mg²⁺ acts as an efficient Lewis acid, with its ion-pairing ability accelerating the catalytic process. A more significant enhancement of catalytic performance was found with the addition of Brønsted acids (HA). CV performed under CO₂ in the presence of Brønsted acids (HA), yielding an increased catalytic current at the potential of [Fe(TPP)]^{2−} species like that observed when Lewis acid was present.

Additional DFT studies by Cove *et al.* and Corbin *et al.* emphasised the importance of oxidation state: reduced forms [M-POR][−] and [M-POR]^{2−} showed higher faradaic efficiency (FE) for CO formation compared to neutral [M-POR]⁰, which had weak CO₂-to-formate conversion.^{31,32} This highlights that

fine-tuning the oxidation state of M-POR catalysts is critical for enhancing CO₂ selectivity and minimising undesired HER activity.^{32–35}

Following on from our previous computational screening of 110 M-PORs for their ability to activate the CO₂ molecule,³⁶ this study now focuses on the most promising catalysts (Fig. 1) and considers their ability to electrochemically convert the CO₂ molecule to CO and HCOOH. Two-dimensional (2D) models of carbonaceous frameworks made from M-POR were also considered to explore for CO₂R. The objective is to determine what factors could enhance the selectivity towards these C₁ products. This work focuses on C₁ products, such as CO and HCOOH, because atomically dispersed electrocatalysts like M-POR lack the adsorption sites necessary for C-C coupling, which leads to C₂ intermediates.³⁴ Consequently, these catalysts are more likely to facilitate the formation of only C₁ products. Moreover, low-temperature CO₂ electrolysis is very effective in producing high yields of C₁ products, making their production advantageous from a techno-economic evaluation perspective.³² Moreover, C₁ products offer several advantages, including easy storage and transportation, and they serve as valuable chemical intermediates and renewable energy carriers. For example, HCOOH can be used as a fuel in direct formic acid fuel cells,^{32,35,37} or CO could be subsequently used as a feedstock for the synthesis of alkanes through the classic Fisher-Tropsch process.³⁸ Moreover, recent advancements in “two-step” tandem electrolysis,^{39,40} where CO₂ is initially converted to CO in a CO₂ electrolyser and then CO is further converted to other products in a CO electrolyser, rely on the assumption that the first CO₂ electrolysis cell can convert CO₂ to CO with high selectivity. This necessitates the development of high-performing and sustainable CO₂-to-CO electrocatalysts.

2. Computational details

2.1 Density functional theory calculations

Depending on the purpose, several DFT codes were used in this study. For the molecular M-PORs, the NWChem software (version 6.6)³⁷ was first used to conduct the full geometry optimisations of the M-PORs in Fig. 1 and the Gaussian16 code⁴¹ was then used to conduct frequency analysis and consider the effect of aqueous environment with the implicit solvation model conductor-like screening model (COSMO).^{42,43} The B3LYP functional^{44,45} combined with the LANL2DZ basis set⁴⁶ was employed for calculations using both NWChem and Gaussian. This level of theory was selected based on the benchmarking done in our work on CO₂ activation.³⁶ For the 2D M-POR carbonaceous frameworks, the CrystalMaker software⁴⁷ was used to generate 2D models from the optimized structures of the M-PORs. These models were initially subjected to a variable cell optimization at the PBE-D3 level of theory, where the *c*-axis (perpendicular to the 2D M-POR) was fixed at 20 Å to minimise the interaction between neighbouring cells, and the lengths of the *a*-axis and *b*-axis were allowed to



[M-POR_x]^{-2,-1,0}

Where M = Ni, Fe, Co, Cu, And x = M-POR system

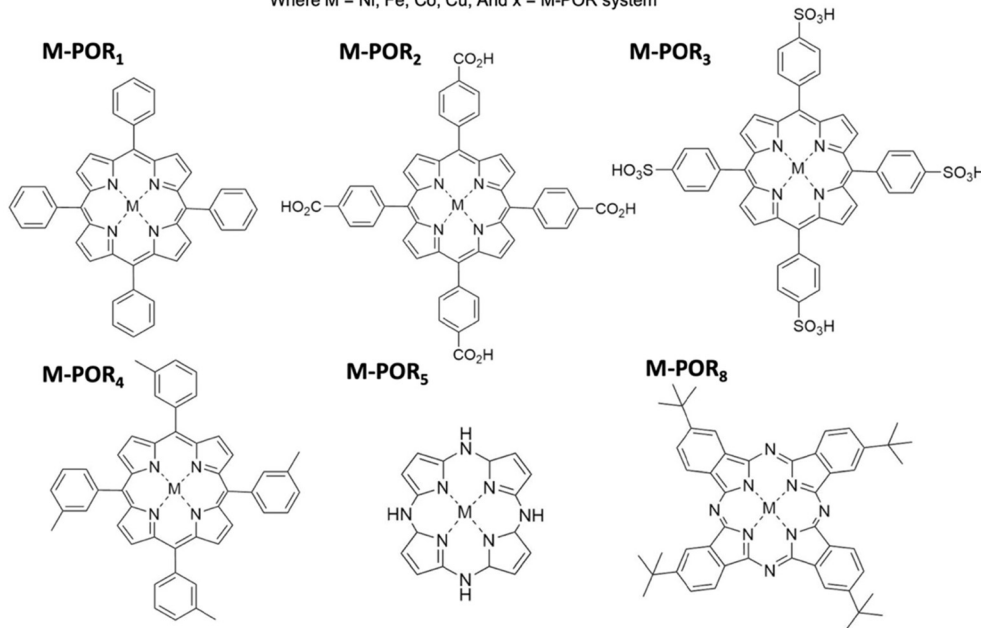


Fig. 1 The metalloporphyrins (M-PORs) with varying metal (M), porphyrin (POR), and oxidation states of the metal (+2, +1, 0). The M-PORs considered are [Ni-POR₁]⁻, [Ni-POR₃]⁻, [Fe-POR₁]⁰, [Fe-POR₂]²⁻, [Fe-POR₃]⁰, [Fe-POR₄]⁰, [Fe-POR₄]²⁻, [Fe-POR₅]⁰, [Fe-POR₈]⁰, [Cu-POR₅]²⁻, [Co-POR₂]²⁻, [Co-POR₄]⁻, and [Co-POR₅]⁰.

relax. Once the unit cell was optimized, the internal coordinates were re-optimized using the hybrid PBE0 exchange correlation functional.⁴⁸ These calculations were conducted using VASP (version 6.4.0).^{49,50} VASP was also used to compare the electron localisation function (ELF) of the 1D (molecular) and 2D M-POR systems. The cut-off energy of the plane waves was set to 500 eV, with a $3 \times 3 \times 3$ k -point mesh used for geometry optimisations and a $15 \times 15 \times 1$ k -point mesh used for the static calculation and for the analysis of the electrons and orbitals.⁵¹ Structural relaxations were performed until the maximum residual force on each atom was less than 0.05 eV \AA^{-1} . Bader charge analysis was carried out with the core charges included.⁵² The d band centre ($e(d)$) and projected density of states (PDOS) was calculated using VASPKIT.⁵³

2.2 Thermodynamic and electrochemical stability

The thermodynamic stability of the M-POR and 2D-M-POR catalysts was estimated by computing the binding energy, cohesive energy, and formation energy.^{34,54} The binding energy (E_b) was determined using the following equation:

$$E_b = E_{\text{M-POR}} - E_{\text{M}} - E_{\text{POR}} \quad (1)$$

where $E_{\text{M-POR}}$, E_{M} , and E_{POR} are the energies of the metalloporphyrin, metal centre, and the porphyrin framework without the metal centre, respectively. The cohesive energy (E_c) was computed using the following equation:

$$E_c = \frac{E_{\text{M(bulk)}}}{N} - E_{\text{M}} \quad (2)$$

where $E_{\text{M(bulk)}}$ is the energy of the bulk unit cell containing N atoms and E_{M} is the energy of the isolated metal atom in vacuum. The formation energy (E_f) was computed according to:

$$E_f = E_{\text{M-POR}} - E_{\text{POR}} - \frac{E_{\text{M(bulk)}}}{N} \quad (3)$$

where $E_{\text{M-POR}}$ is the energy of M-POR, E_{POR} is the energy of the porphyrin framework without the metal centre, $E_{\text{M(bulk)}}$ is the energy of the bulk unit cell containing N atoms and E_{M} is the energy of the isolated metal atom in vacuum.

The electrochemical stability of the M-POR catalysts was evaluated by computing the dissolution potential (U_{diss}) of the metal according to the following expression:⁵⁵

$$U_{\text{diss}}(\text{M}) = U_{\text{diss}}^0(\text{M, bulk}) - \frac{\left(E_{\text{M-POR}} - \frac{E_{\text{M(bulk)}}}{N} - E_{\text{POR}} \right)}{ne} \quad (4)$$

where $U_{\text{diss}}^0(\text{M, bulk})$ is the standard dissolution potential of the bulk metal and n is the number of electrons involved in the dissolution process. In this work, the values of $U_{\text{diss}}^0(\text{M, bulk})$ were taken from Guo *et al.* and n was set to 2.⁵⁶

2.3 Free energies of reaction

The solution-phase Gibbs free energies (G) of the intermediates involved in the electrochemical CO_2 reduction



to CO and HCOOH were calculated according to the expression:

$$G = E_e + \delta G_{\text{VRT}} + \Delta G_{\text{solv}} \quad (5)$$

where the first term is the single point electronic energy, the second term is the frequency correction computed within the harmonic potential approximation at standard conditions ($T = 298.15$ K and $p = 1$ atm), and the third term is the solvation correction computed using the COSMO model. The calculation of the frequencies was also used to verify the minimum character and, consequently, the stability of the isolated catalysts and intermediates. The free energies of reaction for the concerted proton-coupled electron transfer (PCET) steps involved in the CO₂R were studied based on Nørskov's computational hydrogen electrode (CHE) model.⁵⁷ This model provides an efficient approach to study PCET in electrocatalysis without treating solvated protons explicitly and is widely used in computational electrocatalysis, including the CO₂R on molecular and 2D single metal atomic catalysts.^{34,58} In this technique, zero voltage was defined based on the potential energy (μ) of components involved in the reversible hydrogen electrode at all pH, T , and p . Therefore, $\mu(\text{H}^+) + \mu(e^-) = 1/2\mu(\text{H}_2)$ at a potential of 0 V. The limiting potential U_L , an important parameter for evaluating the catalytic activity, was obtained according to the formula:

$$U_L = -\Delta G_{\text{max}}/ne \quad (6)$$

where ΔG_{max} is the Gibbs free energy change of the rate-determining step. In the CHE model, the effect of pH and electrode potential U is considered according to:

$$\Delta G = \Delta E_e + \Delta \delta G_{\text{VRT}} + \Delta \Delta G_{\text{solv}} + \Delta G_{\text{pH}} + \Delta G_U \quad (7)$$

where the fourth term is the free energy correction due to the difference in proton concentration, $\Delta G_{\text{pH}} = 2.303 \times k_B \times T \times \text{pH}$, and the last term is the free energy correction due to the difference in electrode potential, $\Delta G_U = -neU$, where n is the number of electrons transferred in the PCET step ($n = 1$), e is the electronic charge, and U is the applied potential. In this study, both pH and U were set to zero.

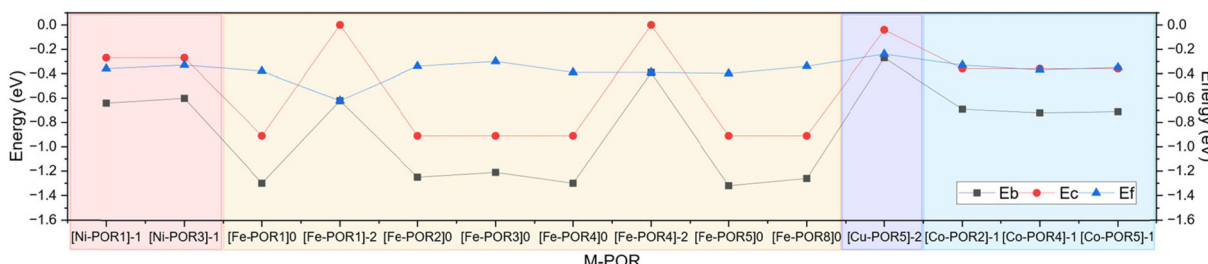


Fig. 2 Thermodynamic stability of the M-PORs. Values of the cohesive energy (E_c) of the transition metal (M) bulk, binding energy between M and POR (E_b), and formation energy (E_f) of M-POR. Pink, yellow, purple, and blue zones correspond to the Ni-POR, Fe-POR, Cu-POR, and Co-POR catalysts, respectively.

- i. $\text{CO}_2 + * \rightarrow * \text{CO}_2$
- ii. $* \text{CO}_2 + \text{H}^+ + \text{e}^- \rightarrow * \text{COOH}$
- iii. $* \text{COOH} + \text{H}^+ + \text{e}^- \rightarrow * \text{CO} + \text{H}_2\text{O}$
- iv. $* \text{CO} \rightarrow \text{CO} + *$



- i. $* \text{CO}_2 + \text{H}^+ + \text{e}^- \rightarrow * \text{OCHO}$
- ii. $* \text{OCHO} + \text{H}^+ + \text{e}^- \rightarrow * \text{HCOOH}$
- iii. $* \text{HCOOH} \rightarrow \text{HCOOH} + *$



Scheme 1 Elementary steps for the two proton/two electron ($2\text{H}^+/2\text{e}^-$) electrochemical CO₂ conversion to CO and HCOOH according to the PCET mechanism. The (*) is used to show adsorbed species.

3. Results and discussion

3.1 Stability and structural features of M-PORs

As reported in Fig. 2 and Table S1 in the ESI,[†] the formation energies (E_f) of metal coordination in the M-PORs are all highly negative, indicating the thermodynamic stability of these complexes. However, the formation energies of the M-PORs vary with the metal centre and oxidation state, with the most stable system being $[\text{Fe-POR}_1]^{2-}$ (-0.62 eV). Also, the neutral $[\text{M-POR}]^0$ catalysts are thermodynamically more stable (lower E_b) than the doubly reduced $[\text{M-POR}]^{2-}$. According to the values of the dissolution potential of the M-PORs (Table S2 in the ESI,[†] all of the catalysts have values of U_{diss} between -0.3 and 0.5 V, which are higher than the experimentally applied potentials for the CO₂R to CO ($E^0 = -0.52$ V) and CHCOOH ($E^0 = -0.61$ V) and of most products of the CO₂R.⁵⁹ This suggests that these active sites are all generally stable against dissolution in the CO₂R and have good electrochemical stability. Consequently, the reaction energetics of the CO₂R on the M-POR catalysts can be examined.

3.2 Mechanisms of reduction reactions on M-PORs

3.2.1 Electrochemical CO₂ reduction to carbon monoxide and formic acid. This section presents calculations of the electrochemical conversion of CO₂ to CO and HCOOH. Two reaction mechanisms are considered: proton-coupled



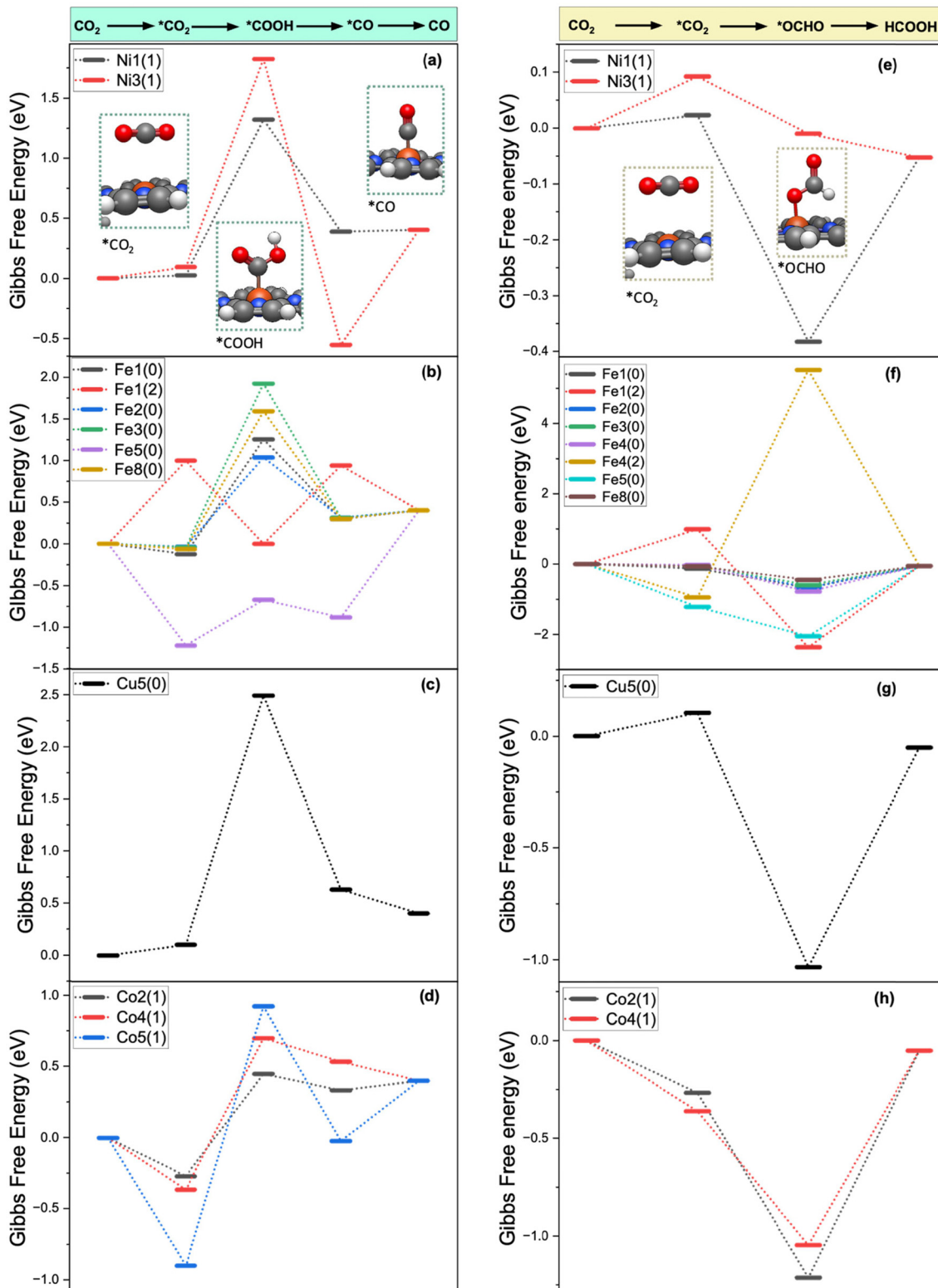
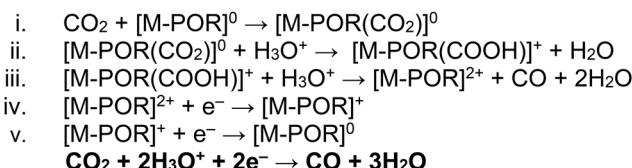


Fig. 3 Gibbs free energy diagram for the CO_2R to CO according to the PCET mechanism on (a) Ni-POR, (b) Fe-POR, (c) Cu-POR, and (d) Co-POR. Gibbs free energy diagram for the CO_2R to HCOOH according to the PCET mechanism on (e) Ni-POR, (f) Fe-POR, (g) Cu-POR, and (h) Co-POR. Calculations conducted at the B3LYP/LANL2DZ/COSMO level of theory. Values in eV.

electron transfer (PCET) and separated proton transfer-electron transfer (PT-ET). In the PCET, the transfer of a

proton (H^+) and an electron (e^-) occurs simultaneously. In contrast, in the PT-ET these steps occur sequentially.



Scheme 2 Elementary steps in the PT-ET mechanism of CO_2 -to-CO conversion with the initial CO_2 adsorption taking place with the metal in the neutral $[\text{M-POR}]^0$ oxidation state.

Proton-coupled electron transfer. The mechanism for PCET to generate CO and HCOOH is shown in Scheme 1.⁶⁰ After the adsorption of CO_2 to the catalyst ($\text{CO}_2 \rightarrow * \text{CO}_2$), the first PCET step leads to two possible intermediates, $* \text{OCHO}$ or $* \text{OCOH}$, depending on the atom coordinated to the catalyst, O or C, respectively. Depending on the intermediate, the second PCET will then generate HCOOH or CO. The free energy of reactions (ΔG) of the elementary steps leading to the formation of HCOOH and CO were computed using the CHE model. For HCOOH, these steps are the PCET to convert adsorbed CO_2^* to the O-coordinated intermediate ($\text{CO}_2^* + \text{H}^+ + \text{e}^- \rightarrow * \text{OCHO}$, $\Delta G_{* \text{OCHO}}$) and the next PCET converts adsorbed $* \text{OCHO}$ to gas-phase formic acid ($* \text{OCHO} + \text{H}^+ + \text{e}^- \rightarrow \text{HCOOH}$, ΔG_{HCOOH}). For CO, the elementary steps are the PCET to convert adsorbed $* \text{CO}_2$ to the C-coordinated intermediate ($* \text{CO}_2 + \text{H}^+ + \text{e}^- \rightarrow * \text{COOH}$, $\Delta G_{* \text{COOH}}$), and the next PCET converts adsorbed $* \text{COOH}$ to adsorbed carbon monoxide ($* \text{COOH} + \text{H}^+ + \text{e}^- \rightarrow * \text{CO} + \text{H}_2\text{O}$, $\Delta G_{* \text{CO}}$), which can then be released from the catalyst surface to form gas-phase CO (ΔG_{CO}).

The Gibbs free energy diagrams of the CO_2 R for the PCET pathways leading to the formation of CO and HCOOH on Ni-

POR ($[\text{Ni-POR}_{1,3}]^-$), Fe-POR ($[\text{Fe-POR}]^{0,2-}$, $[\text{Fe-POR}_2]^0$, $[\text{Fe-POR}_3]^0$, $[\text{Fe-POR}_4]^{0,2-}$, $[\text{Fe-POR}_5]^0$, $[\text{Fe-POR}_8]^0$), Cu-POR ($[\text{Cu-POR}_5]^{2-}$), and Co-POR ($[\text{Co-POR}_{2,4,5}]^-$) are reported in Fig. 3. The stability of the intermediates ($* \text{OCHO}$ and $* \text{CO}$) controlling the formation of CO is significantly affected by the porphyrin ligands, metal centres, and oxidation state. The potential limiting step (ΔG_{PLS}), identified as the step with the highest positive ΔG , serves as an indicator of the catalytic performance of an M-POR. Specifically, a high ΔG_{PLS} signifies poor catalytic performance. The Gibbs free energy diagram for the CO_2 R pathways to CO and HCOOH on each M-POR is discussed in detail below.

Ni-POR. According to Fig. 3(a) and (e), the adsorption of CO_2 is more favourable on $[\text{Ni-POR}_1]^-$ ($\text{DG}_{\text{CO}_2^*} = 0.02 \text{ eV}$) than on $[\text{Ni-POR}_3]^-$ ($\text{DG}_{\text{CO}_2^*} = 0.09 \text{ eV}$). This can be justified by the larger orbital splitting on $[\text{Ni-POR}_1]^-$ (11.1 eV) than on $[\text{Ni-POR}_3]^-$ (2.32 eV), as a larger orbital splitting leads to a less favourable adsorption (see Fig. S3(a) and (c) in the ESI†), thus making it more difficult to activate CO_2 . The formation of $* \text{OCHO}$ is most favourable on $[\text{Ni-POR}_1]^-$ because of the lower free energy of the OCHO^* intermediate ($\Delta G_{* \text{OCHO}} = -0.41 \text{ eV}$), which suggests that the CO_2 R is more selective towards HCOOH on this system. Similarly, the formation of OCHO^* is most favourable on $[\text{Ni-POR}_3]^-$ ($\Delta G_{* \text{OCHO}} = -0.10 \text{ eV}$). The formation of $* \text{CO}$ is more favourable on $[\text{Ni-POR}_3]^-$ with $\Delta G_{* \text{CO}} = -0.55 \text{ eV}$ than on $[\text{Ni-POR}_1]^-$. To summarise, Ni-POR systems favour the formation of HCOOH over the formation of CO.

Fe-POR. For the Fe-POR catalysts in Fig. 3(b) and (f), the most favourable system for the formation of $* \text{CO}$ is $[\text{Fe-POR}_5]^0$ ($\text{DG}_{* \text{CO}_2} = -1.22 \text{ eV}$), while $[\text{Fe-POR}_1]^{2-}$ has the least favourable CO_2 adsorption ($\text{DG}_{* \text{CO}_2} = 1.0 \text{ eV}$). All Fe-POR systems favour

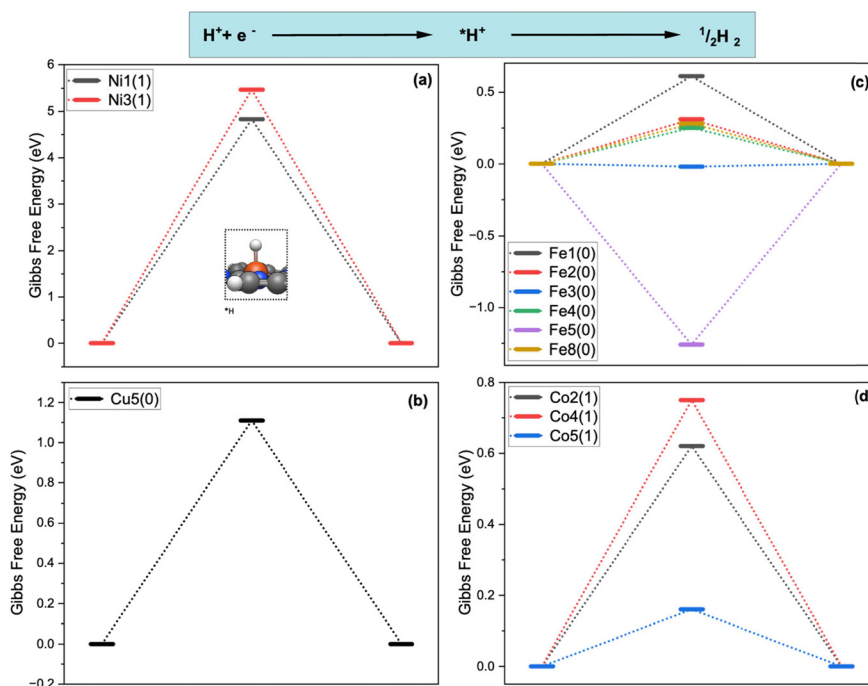


Fig. 4 Gibbs free energy diagram for the HER on (a) Ni-POR, (b) Fe-POR, (c) Cu-POR, and (d) Co-POR. Calculations conducted at the B3LYP/LANL2DZ/COSMO level of theory. Values in eV.



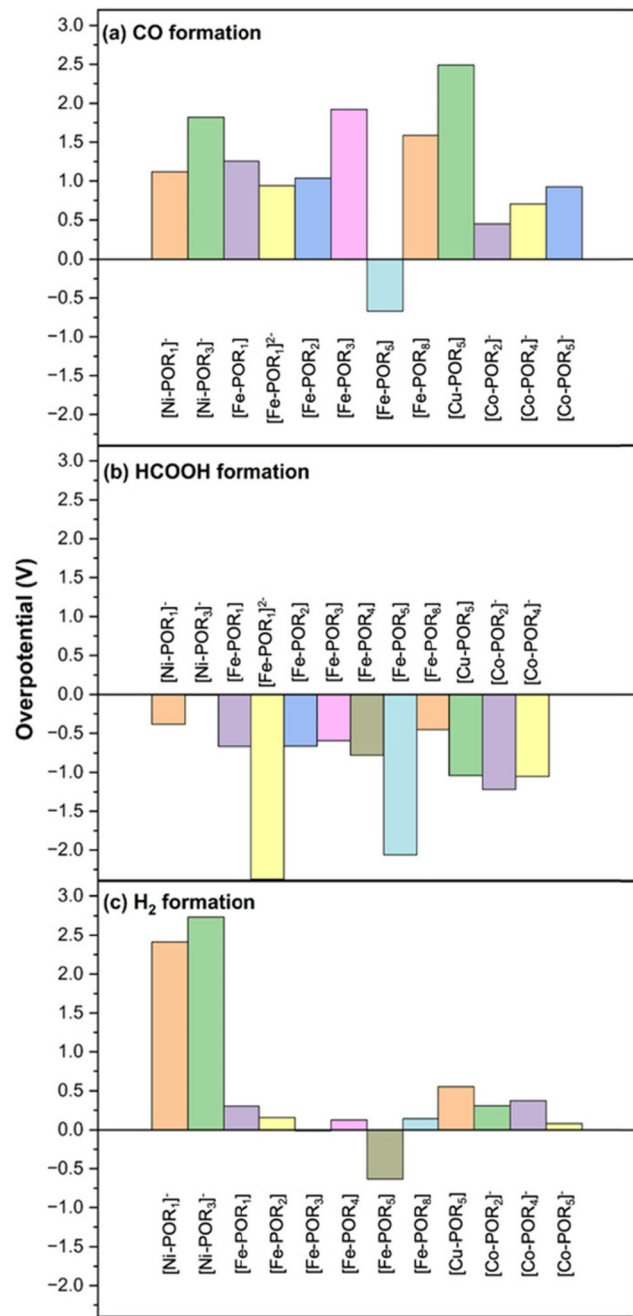


Fig. 5 Overpotential for the electrocatalytic formation of (a) CO, (b) HCOOH, and (c) H₂ on M-POR. Values in V.

the formation of *OCHO over *COOH except for [Fe-POR₅]⁰, which favours *COOH by 0.04 eV. The overall trend from the most favourable to the least favourable Fe catalyst based on the DG_{PLS} of the reaction for HCOOH formation is [Fe-POR₄]²⁻ > [Fe-POR₈]⁰ > [Fe-POR₃]⁰ > [Fe-POR₂]⁰ > [Fe-POR₁]⁰ > [Fe-POR₄]⁰ > [Fe-POR₅]⁰. This highlights the significance of the effect of moving from d⁸ to d⁶ electron configuration on the CO₂R: [Fe-POR₁]²⁻ (d⁸) is more reactive than [Fe-POR₁]⁰ (d⁶). These complexes are all high spin. Moreover, ligands with alkyl benzyl ([Fe-POR₄]²⁻ and [Fe-POR₈]⁰ in Fig. 1) are more reactive than those with carboxyl benzyl groups ([Fe-POR₂]⁰).

However, the M-POR [Fe-POR₅]⁰ is shown to be the least reactive despite the less steric hindrance, which indicates the importance of the ligands in M-POR catalysts in affecting their catalytic behaviour. Insights into the d-orbital splitting were obtained by computing the PDOS of the d-orbitals of the metal centre (Fig. S3 in the ESI†) to determine the energy difference between the d-orbitals with maximum PDOS (Fig. S4†). The orbitals presented for the PDOS of M-POR-CO and M-POR-HCOOH (Fig. S3†) show strong overlap and, thus, strong bonding. Based on the d-band values, Fe systems exhibit superior catalytic properties compared to other M-POR systems. The d-band centre is calculated in VASP by analysing the PDOS of the metal's d-orbitals in the system. The d-band centre is defined as the energy of the centre of mass of the d-states, often measured relative to the Fermi level.⁶¹ It correlates to catalytic performance because a d-band centre closer to the Fermi level typically enhances adsorption of reaction intermediates, thereby improving catalytic activity. If the d-band centre is too high or low, it may lead to too weak or too strong adsorption, respectively, which can reduce efficiency in catalytic reactions. The [Fe-POR₃-CO]⁰ system has a d-band center value of -0.30 eV, positioned close to the Fermi level (Fig. S4(k)†). According to previous reports,^{62,63} the closer the d-band centre is to the Fermi level, the higher the electron occupation of the bonding state is, and the stronger the adsorption strength of the adsorbed molecules on the catalyst surface is because of the increase of orbital overlap and enhancement of electron donation and back-donation between the metal centre and the adsorbed CO₂.⁶⁴ This electronic effect stabilizes the CO₂ adsorption intermediate by forming stronger chemical bonds and lowering the energy of the intermediate state. Consequently, the stabilization provided by the optimal d-band positioning directly contributes to facilitating the CO₂ adsorption step, identified as the rate-determining step in our catalytic mechanism. The [Fe-POR₄]²⁻ system, with a d-band centre value of -0.32 eV (Fig. S4(p)†), demonstrates the best catalytic performance for HCOOH formation. Applying a potential of -1 V makes CO₂R to CO on [Fe-POR₁]⁰ more favourable than at 0 V, as ΔG decreases from -0.39 eV (at 0 V) to -1.39 eV (at -1 V; Fig. S8†). This is consistent with Lu's findings, which show that higher applied potentials increase the FE for CO formation.⁶⁵ The applied potential also favours the HER (-0.39 eV), although it is not as favoured as CO formation.

Cu-POR. [Cu-POR₅]⁰ in Fig. 3 was the only Cu-POR system that activates CO₂ in our previous study.³⁶ This seems feasible as all Cu-PORs are d¹⁰, making them very stable. However, in the neutral state, they are in d⁹ and can donate an electron to CO₂ (Table S2 in the ESI†). In Fig. 3(c), the pathways to both HCOOH and CO formation have very stable intermediates which lead to high DG_{PLS}. The [Cu-POR₅]²⁻ system shows to be consistent for CO and HCOOH formation with d-band centre values of -2.54 eV and -2.20 eV, respectively (Fig. S4(u) and (v)†), which suggests poor binding of substrates to the Cu centre. In a square planar d⁹ configuration, Cu(II) metal has fewer available low-energy

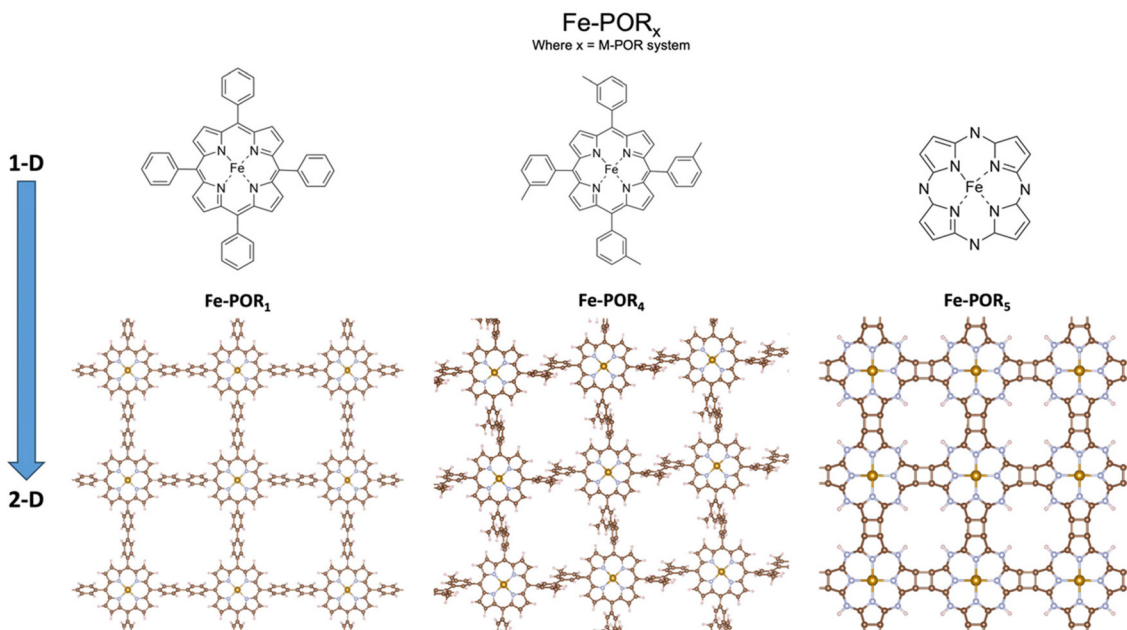


Fig. 6 1D and 2D Fe-POR systems considered for the carbonaceous framework analysis. Structures obtained at the PBE0 level of theory using VASP.

empty orbitals to effectively participate in back-donation to CO_2 . In contrast, Fe(II) metals in a d^6 square planar configuration can engage in stronger π -backbonding, where electron density from the metal is donated into the antibonding π^* orbitals of CO_2 . This increased electron donation helps activate CO_2 by weakening the C–O bonds and bending its linear structure, facilitating reduction. Because Cu(II) in a d^9 state lacks a fully effective π -backbonding interaction, it forms weaker bonds with CO_2 , making its activation less favorable compared to a d^6 metal. This could also suggest that the substrates can easily be desorbed from Cu-POR as there is less orbital overlap between the Cu centre and the substrate orbitals, meaning less electrons shared.

Co-POR. In Fig. 3(d) and (h), the CO_2 adsorption is favourable on all Co-MORs, with free energies of adsorption being between $DG_{\text{CO}_2}^* = -0.90$ eV on $[\text{Co-POR}_5]^0$ and $DG_{\text{CO}_2}^* = -0.27$ eV and the least favourable is on $[\text{Co-POR}_2]^0$. The $[\text{Co-POR}_2]^-$ and $[\text{Co-POR}_4]^-$ catalysts favour the formation of the $^*\text{OCHO}$ intermediate ($DG_{\text{OCHO}^*} = -0.95$ eV and -0.68 eV, respectively) over $^*\text{COOH}$ ($DG_{\text{COOH}^*} = 0.45$ eV and 0.70 eV, respectively). However, $[\text{Co-POR}_5]^-$ favours $^*\text{COOH}$ ($DG_{\text{COOH}^*} = 0.92$ eV) and highly disfavours OCHO^* formation. Overall, the most favourable CO_2R pathways to CO and HCOOH occur on $[\text{Co-POR}_2]^-$, where $DG = 0.67$ eV for CO formation and $DG = 0.05$ eV for HCOOH formation.

Proton transfer-electron transfer. Gottle and Koper proposed an alternative PT-ET mechanism for the electrochemical CO_2 conversion to CO on M-POR,⁶⁶ which includes the following elementary steps (Scheme 2): (i) CO_2 adsorption onto the metal centre of M-POR; (ii) protonation of the $[\text{M-POR}(\text{CO}_2)]$ intermediate to form bound formate

$[\text{M-POR}(\text{CO}_2\text{H})]$; (iii) protonation of the $[\text{M-POR}(\text{CO}_2\text{H})]$ intermediate to form bound CO, $[\text{M-POR}(\text{CO})]$; (iv) electron transfer to the $[\text{M-POR}(\text{CO})]$ intermediate to release CO; (v) electron transfer to reduce the metalloporphyrin to its initial oxidation state. In steps 2 and 3, H_3O^+ was used as the proton source. The reaction free energy diagrams for the PT-ET on Ni-POR ($[\text{Ni-POR}_1]^-$, $[\text{Ni-POR}_3]^-$), Fe-POR ($[\text{Fe-POR}_1]^0$, $[\text{Fe-POR}_1]^-$, $[\text{Fe-POR}_2]^0$, $[\text{Fe-POR}_3]^0$, $[\text{Fe-POR}_4]^0$, $[\text{Fe-POR}_4]^-$, $[\text{Fe-POR}_5]^0$, $[\text{Fe-POR}_8]^-$), Cu-POR ($[\text{Cu-POR}_5]^{2-}$), and Co-POR ($[\text{Co-POR}_2]^{2-}$, $[\text{Co-POR}_4]^-$, $[\text{Co-POR}_5]^-$) are reported in Fig. S3 of the ESI.[†] The reaction free energies for PT-ET are significantly higher than those of PCET, which makes PCET by far the most favourable pathway for the CO_2R on M-POR.

3.2.2 Hydrogen evolution reaction. In aqueous solutions, there is a continuous competition between the CO_2R and the HER.⁶⁷ Typically, the FE of the HER outperforms that of CO_2R due to its significantly lower overpotential on most M-POR catalysts.⁶⁸ The free energy of reaction for the HER on 16 M-PORs is reported in Fig. 4. Most systems exhibit a positive free energy of proton H adsorption (ΔG_{H}). However, there are exceptions. $[\text{Fe-POR}_3]^0$ and $[\text{Fe-POR}_5]^0$ show ΔG_{H} values of -0.01 eV and -0.63 eV, respectively, which favours the HER, potentially making them less suitable for CO_2R . This could be because they are both d^6 systems, and from our previous work on CO_2 activation,³⁶ M-PORs with the metal in the d^6 configuration were found to not favour both CO_2R and the HER. In contrast, $[\text{Ni-POR}_3]^-$ has the highest ΔG_{H} , 2.73 eV, and thus is not active towards the HER.⁵⁶ DFT calculations of the HER on a transition metal embedded in single vacancies of graphene reported ΔG_{H} of 1.46 eV for Ni and 0.14 eV for Fe,⁶⁹ supporting the trend that Ni-POR is less favourable to the HER than Fe-POR.



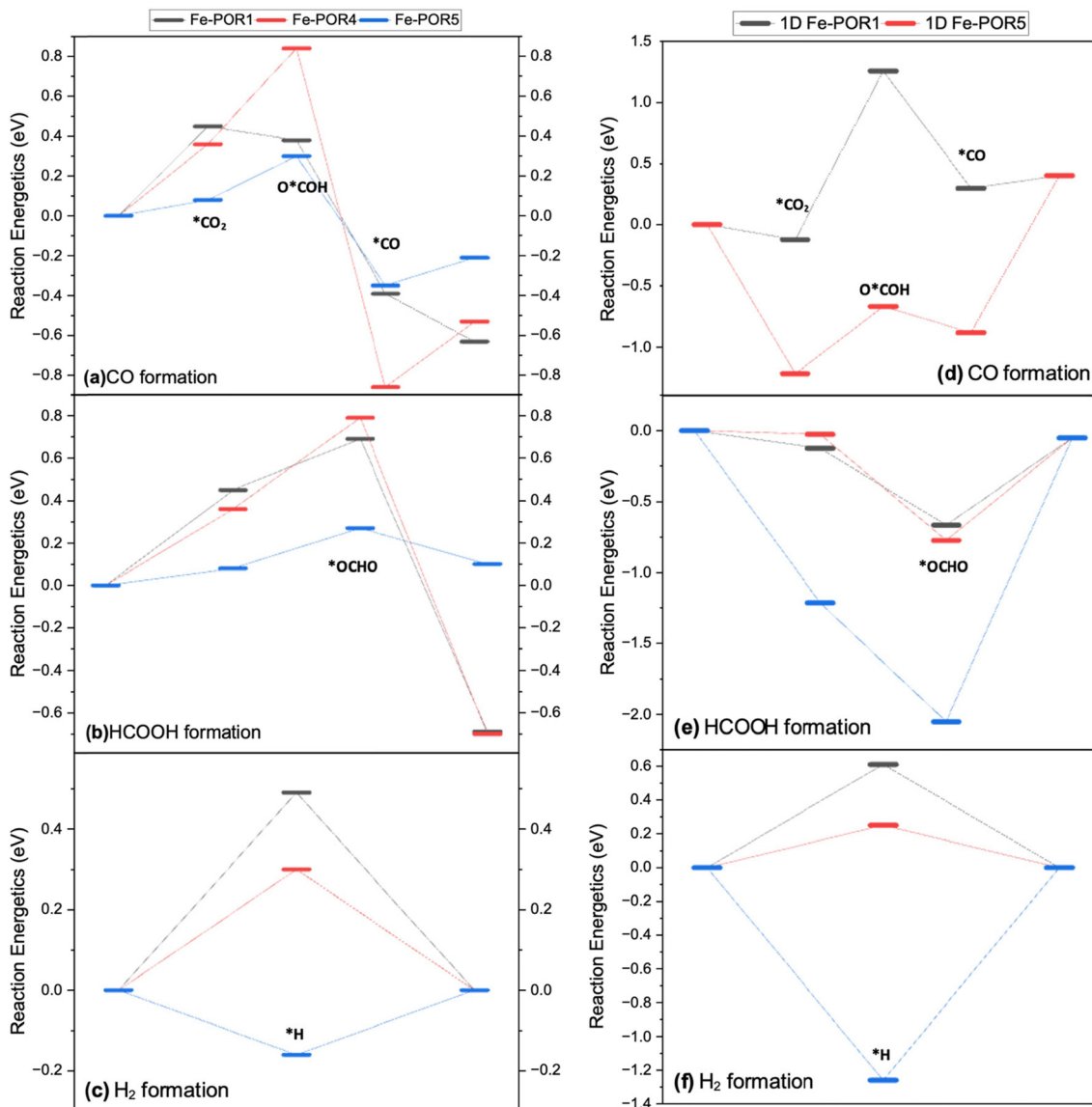


Fig. 7 Reaction energetics for the CO₂R to (a) CO and (b) HCOOH formation and (c) the HER to H₂ formation on the 2D Fe-POR_{1,4,5} catalysts and reaction energetics for the CO₂R to (d) CO and (e) HCOOH formation and (f) the HER to H₂ formation on the 1D-Fe-POR_{1,4,5} catalysts. Calculations conducted at the PBE0 level of theory. Values in eV.

3.2.3 Selectivity. In electrocatalysis, the overpotential (η) represents the extra voltage required beyond the theoretical value necessary to drive an electrochemical reaction. Computationally, the overpotential can be obtained from the difference between the equilibrium ($U_{\text{eq.}}$) and the limiting ($U_{\text{eq.}}$) potentials:

$$\eta = U_{\text{eq.}} - U_{\text{L}} \quad (8)$$

Thus, η represents the minimum applied potential required to facilitate the formation of relevant intermediates. This value reflects the thermodynamic barrier the catalyst needs to overcome. Large values of η indicate a poor catalytic performance. Therefore, for efficient CO₂R, the desired catalyst should exhibit lower

overpotentials for CO₂R compared to the HER. This ensures that CO₂R is prioritised over the competing HER process. The overpotentials for the CO₂R to CO and HCOOH and for the HER on the M-POR catalysts are summarised in Fig. 5. Almost all systems favour HCOOH formation over H₂ or CO formation. Overall, the systems favourable to HCOOH are ranked in the following order: [Fe-POR₅]⁰ > [Co-POR₅]⁻ > [Co-POR₄]⁻ > [Co-POR₂]⁻ > [Fe-POR₁]⁰ > [Fe-POR₈]⁰ > [Fe-POR₃]⁰ ≈ [Fe-POR₂]⁰ > [Fe-POR₄]⁰ > [Ni-POR₁]⁻ > [Ni-POR₃]⁻. Thus, [Fe-POR₅]⁰ is the most efficient catalyst with the lowest overpotential (-2.06 V). On the other hand, HER is generally favoured over the CO₂-to-CO conversion process, except for [Ni-POR₁]⁻ and [Ni-POR₃]⁻, which exhibit much lower overpotentials (high catalytic activity) for CO formation (η

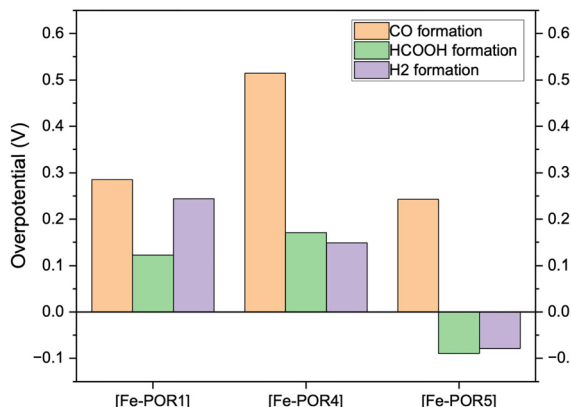


Fig. 8 Overpotentials for the H_2 , CO, and HCOOH formation on the 2D Fe-POR₅.

= 1.12 V and 1.82 V, respectively) compared to H_2 (η = 2.41 V and 2.73 V, respectively).

3.3 CO₂R and the HER on two-dimensional M-PORs

The versatility of M-PORs, coming from the variety of available metal centres and organic linkers, makes them ideal building blocks for advanced porous frameworks with unique properties for diverse catalytic applications including CO₂ reduction.¹⁶ 2D M-PORs were thus designed using the most promising molecular M-POR catalysts to investigate their activity towards CO₂R. Three Fe-PORs were selected as precursors for the generation of 2D frameworks, as shown in Fig. 6. These systems were chosen based on the efficiency of their 1D counterparts in the CO₂-to-CO or CO₂-to-HCOOH conversion process. Moreover, their potential suitability for forming stable 2D structures was evaluated by computing their cohesive energy, binding energy, formation energy, and dissolution potential (Fig. S6 in the ESI†). The 2D Fe-POR systems display good thermodynamic stability, attributed to the absence of heteroatoms in the peripheral ligands, which often introduces structural or electronic instabilities in similar complexes.^{70,71}

The free energy pathways for the CO₂R to CO and HCOOH and the HER to H_2 on the 2D and 1D Fe-POR catalysts are compared in Fig. 7. The stability of the intermediates *OCHO and *COOH, which control the formation of HCOOH and CO, respectively, is significantly affected by the POR ligands. For CO formation (Fig. 7(a)) the adsorption of CO₂ is more favourable on 2D Fe-POR₅ ($\text{DG}_{\text{CO}_2^*} = 0.08$ eV) than on 2D Fe-POR₁ and 2D Fe-POR₄ ($\text{DG}_{\text{CO}_2^*} = 0.45$ eV and 0.36, eV respectively). As all systems have the same metal centre, the metal does not influence the stability of the intermediates. The adsorption of *COOH is also most favourable on 2D Fe-POR₅ with $\text{DG}_{\text{COOH}^*} = 0.30$ eV, suggesting 2D Fe-POR₅ as the most favourable system for CO formation and this is the G_{PLS} . The strongest *CO adsorption is found with 2D Fe-POR₄ at -0.86 eV, leading to unfavourable desorption of CO. For HCOOH formation (Fig. 7(b)), 2D Fe-POR₅ shows the

most favourable CO₂ adsorption ($\text{DG}_{\text{CO}_2^*} = 0.08$ eV) and the most favourable *OCHO intermediate adsorption ($\text{DG}_{\text{OCHO}^*} = 0.27$ eV). This suggests that 2D Fe-POR₅ is the most promising candidate for HCOOH formation. 2D Fe-POR₁ and 2D Fe-POR₄ show similar reaction thermodynamics, which suggests poor favourability to HCOOH formation. Therefore, from the free energy profiles, 2D Fe-POR₅ is the most favourable for C₁ product formation. However, when looking at the 1D Fe-POR, the order of reactivity is [Fe-POR₁]⁰ > [Fe-POR₄]⁰ > [Fe-POR₅]⁰, with [Fe-POR₅]⁰ being the least favourable.

These results suggest that 2D Fe-POR₅ is the most promising candidate for HCOOH formation. In contrast, 2D Fe-POR₁ and 2D Fe-POR₄ exhibit similar reaction thermodynamics, indicating poor favourability for HCOOH production. Interestingly, while Fe-POR₅ is the least reactive in the 1D configuration ([Fe-POR₁]⁰ > [Fe-POR₄]⁰ > [Fe-POR₅]⁰), transitioning to the 2D structure significantly enhances its catalytic activity. This trend correlates with the shift in its d-band center, which decreases from -0.64 eV (Fig. S3(q)†) in 1D to -2.14 eV (Fig. S7(e)†) in 2D, suggesting stronger adsorption and activation of CO₂.

A similar trend is observed in the HER. Among the 2D Fe-POR catalysts, Fe-POR₅ exhibits the most favorable adsorption of the H intermediate ($\text{DG}_{\text{H}^*} = -0.16$ eV), while 2D Fe-POR₁ and 2D Fe-POR₄ have positive adsorption energies, indicating lower activity. Notably, the reactivity trend for HER in the 2D configuration follows the same order as in the 1D configuration (Fig. 7(d-f)), reinforcing the relationship between electronic structure and catalytic performance.

The overpotentials for the eCO₂R and HER on the 2D M-POR are shown in Fig. 8. For Fe-POR₅, negative overpotentials were observed for both HCOOH (-0.09 V) and H_2 (-0.08 V), indicating that both reactions are highly favoured with no significant preference for one over the other. In contrast, Fe-POR₁ exhibited a distinct difference in overpotentials, with HCOOH formation being more favoured (0.12 V) compared to H_2 formation (0.24 V).

To gain deeper insight into the electronic properties influencing CO₂ activation, the electron localisation function (ELF) was computed for 2D Fe-POR_{1,4,5} catalysts. The ELF provides a measure of electron density distribution, particularly around the Fe center, helping to visualise the degree of charge accumulation and its influence on adsorption. The ELF maps for CO⁻ and OCOH⁻ adsorbed M-POR systems (Fig. 9) reveal notable differences in electron density distribution. In Fe-POR₁ (Fig. 9(a)), asymmetric electron density accumulation is observed, particularly on one side of the extended porphyrin ring. This localised electron density may guide CO₂ adsorption towards the catalytic Fe center, facilitating activation. In contrast, Fe-POR₄ exhibits a more dispersed and weakly localised electron density, which could contribute to its poorer catalytic performance by reducing effective charge transfer to the adsorbate. Finally, Fe-POR₅ displays strong regions of localized electron density near the Fe center, possibly due to

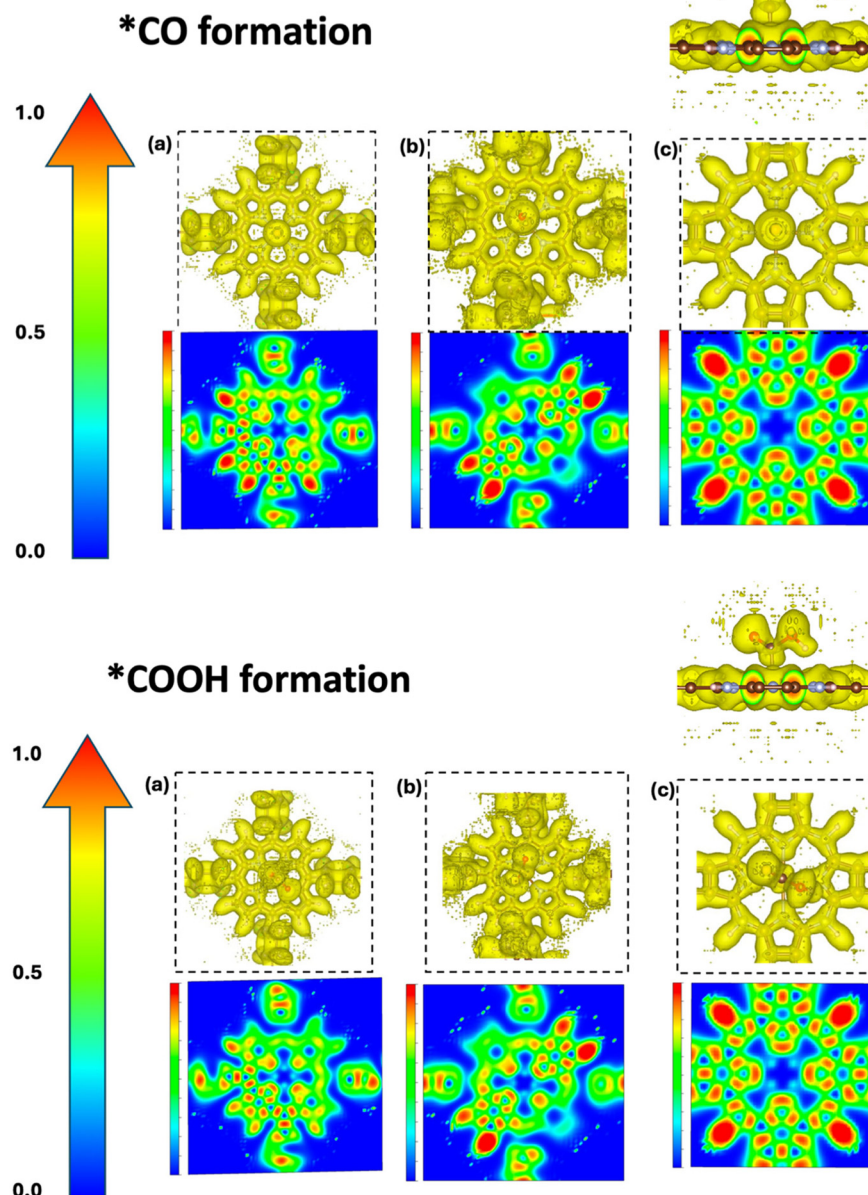


Fig. 9 ELF plots for CO (top) adsorbed on (a) 2D-Fe-POR₁, (b) 2D-Fe-POR₄, and (c) 2D-Fe-POR₅, and for OCOH (bottom) adsorbed on (a) 2D-Fe-POR₁, (b) 2D-Fe-POR₄, and (c) 2D-Fe-POR₅.

its smaller porphyrin framework restricting charge delocalisation. This localised charge distribution may enhance Fe-CO₂ interactions, leading to improved catalytic activity.

Further insights can be drawn from projected density of states (PDOS) analysis (Fig. S7†). Fe-POR₅ exhibits the smallest bandgap (-2.14 eV for CO-adsorbed and -1.91 eV for OCOH-adsorbed configurations), suggesting enhanced electronic conductivity, which may facilitate charge transfer processes during CO₂ reduction. The combination of ELF and PDOS data highlights the importance of electronic structure in governing catalytic performance, with Fe-POR₅

emerging as the most promising candidate among the studied systems.

4. Conclusions

A computational investigation of the electrocatalytic CO₂ reduction reaction on various metalloporphyrin catalysts with different metal centers (Ni, Fe, Cu, and Co), ligands, and oxidation states was conducted using DFT calculations. Thermodynamic and electrochemical stability analyses revealed that neutral systems exhibit higher stability compared to charged systems, with doubly reduced systems being the

least stable. Two possible reaction mechanisms—proton-coupled electron transfer and sequential proton transfer–electron transfer—were explored, with PCET identified as the significantly more favourable pathway, predominantly forming HCOOH rather than CO. Among the studied catalysts, Fe–POR demonstrated superior catalytic performance towards these C₁ products. Comparison of overpotentials for CO₂ reduction and the competing hydrogen evolution reaction showed that most systems preferentially produce HCOOH.

Transitioning to two-dimensional (2D) frameworks, particularly with Fe–POR systems, led to enhanced catalytic activity compared to their molecular counterparts, highlighting the potential for these frameworks in practical electrocatalysis applications. The main electronic descriptor impacting catalytic performance and product selectivity in both molecular and 2D M–POR frameworks appears to be the position of the d-band center relative to the Fermi level.

This study provides fundamental insights into the rational design and optimization of effective metalloporphyrin-based electrocatalysts, guiding future development toward efficient and selective CO₂ conversion technologies.

Data availability

The data supporting the findings of this study are openly accessible on ZENODO at <https://zenodo.org/records/11361107>, reference number 11361107. Additionally, the authors confirm that the data are included in the ESI† and can also be requested from the corresponding author, DDT.

Conflicts of interest

The authors declare no competing financial interest.

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