



Cite this: *J. Mater. Chem. A*, 2024, **12**, 31547

## Low CO<sub>2</sub> mass transfer promotes methanol and formaldehyde electrosynthesis on cobalt phthalocyanine†

Jie Zhang,<sup>a</sup> Thi Ha My Pham,<sup>b</sup> Shibo Xi,<sup>c</sup> Liping Zhong,<sup>b</sup> David Liem,<sup>a</sup> Futian You,<sup>a</sup> Ben Rowley,<sup>d</sup> Ramesha Ganganahalli,<sup>e</sup> Federico Calle-Vallejo,<sup>b</sup> Boon Siang Yeo,<sup>a</sup>

Cobalt phthalocyanine supported on carbon nanotubes (CoPcCNT) usually catalyzes the electroreduction of CO<sub>2</sub> (CO<sub>2</sub>RR) to CO, although several reports have also indicated methanol formation. Herein, by analyzing the effects of CoPc loading and CO<sub>2</sub> partial pressure on the CO<sub>2</sub>RR, we show that a lower rate of CO<sub>2</sub> mass transfer to each CoPc favors methanol formation, while a higher rate of CO<sub>2</sub> mass transfer favors CO evolution. The ratio of the production rates of methanol and CO is related to the average CO<sub>2</sub> mass transfer rate by a power function with a negative exponent. Hence, methanol can only be formed when the supply of CO<sub>2</sub> feed is low. This mass transfer effect is supported by supplementary experiments and computational modelling, which show that CO binding to CoPc is weaker than that of CO<sub>2</sub>, in agreement with previous studies. Consequently, \*CO may only be reduced to methanol when the supply of CO<sub>2</sub> is low and the dwelling time of CO is long. We further provide a quantitative guideline for the design of methanol-selective catalysts. At  $-0.86$  V vs. RHE, enhanced CO mass transfer boosts the CORR to methanol with a faradaic efficiency up to 70% at a total current density of  $-19$  mA cm<sup>-2</sup>. The production of formaldehyde, a reaction intermediate from CO reduction to methanol, is also boosted with a faradaic efficiency of up to 7%. We pinpoint CoPc containing Co(I) as the active CO<sub>2</sub>RR site.

Received 22nd May 2024  
Accepted 19th October 2024  
DOI: 10.1039/d4ta03531c  
[rsc.li/materials-a](http://rsc.li/materials-a)

## Introduction

The electrochemical reduction of CO<sub>2</sub> or CO (CO<sub>2</sub>RR or CORR) using renewable electricity is a promising method for producing

<sup>a</sup>Department of Chemistry, Faculty of Science, National University of Singapore, 3 Science Drive 3, Singapore 117543, Singapore. E-mail: chmyeos@nus.edu.sg

<sup>b</sup>Laboratory of Materials for Renewable Energy (LMER), Institute of Chemical Sciences and Engineering (ISIC), Basic Science Faculty (SB), École Polytechnique Fédérale de Lausanne (EPFL) Valais/Wallis, Energypolis, Rue de l'Industrie 17, Sion CH-1951, Switzerland

<sup>c</sup>Institute of Sustainability for Chemicals, Energy and Environment (ISCE2), Agency for Science, Technology and Research (A\*STAR), 1 Pesek Road Jurong Island, Singapore 627833, Republic of Singapore

<sup>d</sup>Shell Global Solutions International B.V., Energy Transition Campus Amsterdam, Grasweg 31, 1031 HW, Amsterdam, The Netherlands

<sup>e</sup>Shell India Markets Private Ltd Plot No. 7, Bengaluru Hardware Park, Mahadeva, Kodigehalli, Bangalore, North, 562149, India

<sup>f</sup>Nano-Bio Spectroscopy Group and European Theoretical Spectroscopy Facility (ETSF), Department of Advanced Materials and Polymers: Physics, Chemistry and Technology, University of the Basque Country UPV/EHU, Avenida Tolosa 72, San Sebastián 20018, Spain. E-mail: federico.calle@ehu.es

<sup>g</sup>IKERBASQUE, Basque Foundation for Science, Plaza de Euskadi 5, Bilbao 48009, Spain

† Electronic supplementary information (ESI) available: Materials and methods, synthetic details, supplementary figures, tables, and equations related to material characterization, pH simulation, computational modeling, and ICP, Raman, TEM, and EDS studies. See DOI: <https://doi.org/10.1039/d4ta03531c>

green liquid fuels and chemical feedstocks.<sup>1–3</sup> Extensive efforts have been dedicated in recent years to understanding the CO<sub>(2)</sub>RR process, with the aim of improving its selectivity and activity to value-added, highly reduced products, such as methanol, ethylene, and ethanol.<sup>4,5</sup> Among these products, methanol has received relatively less attention, probably due to the lack of robust catalysts and clear routes for its electrocatalytic production.<sup>6,7</sup> Indeed, while many materials have been reported to catalyze the CO<sub>2</sub>RR to methanol, only a few of them have been proven reproducible.<sup>6,8</sup> Recently, the groups of Wang and Robert reported that cobalt phthalocyanine (CoPc) exhibits catalytic activity for the CO<sub>2</sub>RR to methanol.<sup>9–13</sup> Yet, interestingly, alongside these reports, numerous other research groups have observed, using the same catalyst, that CO is the sole CO<sub>2</sub>RR product.<sup>14–17</sup> Compared with CO, methanol has a higher volumetric energy density, can be easily stored and used directly as a fuel.<sup>18</sup> Therefore, it is of commercial and scientific interest to understand in detail the mechanism of the CO<sub>2</sub>RR to methanol using CoPc catalysts.

Wang and coworkers first identified the molecular dispersion of CoPc on a carbon nanotube support (CoPcCNT) as a factor to incline the selectivity of the CO<sub>2</sub>RR toward methanol.<sup>9,19</sup> However, other studies using similar catalysts reported CO as the only product.<sup>20–22</sup> This suggests that the molecular dispersion of CoPc may not govern methanol selectivity alone

and other factors still need to be elucidated. Along these lines, Wang's group observed that the loading of CoPc on carbon nanotubes impacts the CO<sub>2</sub>RR and CORR selectivity to methanol.<sup>11,19</sup> They also found that such selectivity depends on CO partial pressure.<sup>23</sup> The curvature of the carbon nanotube support and the spin state of the cobalt center have further been associated with methanol production.<sup>24,25</sup> More recently, Ren *et al.* showed that blending 10% CO<sub>2</sub> into a CO feed gas can inhibit the CORR to methanol and attributed this inhibition to the stronger binding of CO<sub>2</sub> on CoPc, as compared to CO on CoPc.<sup>13</sup> McCrory and coworkers further revealed, on the basis of microkinetic analyses, that CO<sub>2</sub> binds to CoPc with an effective equilibrium constant (11.1 atm<sup>-1</sup>) that is thrice that of CO binding (3.4 atm<sup>-1</sup>).<sup>26</sup>

Herein, we show that the average CO<sub>2</sub> mass transfer rate to each CoPc active site is the factor that determines how CoPc loading and CO<sub>2</sub> partial pressure affect the competing reduction of CO<sub>2</sub> to methanol and CO. We further reveal a power function ( $R = 0.97\nu^{-0.78}$ ) relating the ratio ( $R$ ) of the production rates of methanol and CO to the average CO<sub>2</sub> mass transfer rate ( $\nu$ ). Computational modelling indicates that this relation is caused by the weaker adsorption of CO compared to CO<sub>2</sub> on CoPc, such that only when the CO<sub>2</sub> supply falls short, can the adsorbed \*CO be further reduced to methanol. Further *in situ* Raman and cyclic voltammetry analyses show different behaviors of CoPc in CO<sub>2</sub>- and CO-saturated electrolytes, which are likely caused by the stronger adsorption of CO<sub>2</sub> compared to CO on CoPc. The computational modeling together with the experimental results implies that the dwelling time of \*CO on CoPc is critical in determining either its further conversion to methanol or its elution as a product. Additionally, DFT modelling based on the Sabatier principle provides a novel quantitative guideline for the design of methanol-selective catalysts, namely  $-0.2 \text{ eV} < \Delta G_{\text{CO}} < 0.2 \text{ eV}$ . Finally, we show that an increase in CO mass transfer to each CoPc molecule boosts the faradaic efficiency (FE) of methanol up to 70% and the FE of formaldehyde up to 7%. Our findings highlight the importance of CO<sub>2</sub> mass transfer in tuning the product selectivity of the CO<sub>2</sub>RR on CoPcCNT to either C<sub>1</sub> oxygenates or CO.

## Results and discussion

### Synthesis of molecularly dispersed CoPc on carbon nanotubes

We first anchored CoPc onto the surfaces of multiwalled carbon nanotubes (CNTs). The resulting materials are denoted as CoPcCNT\_x%, where x% represents the loading of CoPc on the CNTs in weight percentages, from 0.2 to 7.0% (Fig. S6†). As the molecular dispersion of CoPc on CNTs is critical for the production of methanol,<sup>9</sup> we characterized the CoPcCNT\_x% extensively to confirm its successful synthesis. CoPcCNT\_7.0%, which has the highest CoPc loading, is likely to contain CoPc aggregates, if these were formed.<sup>11</sup> Hence, we used it here as a representative example for physical characterization.

High resolution transmission electron microscopy (HRTEM) images show that CoPcCNT\_7.0% and pristine CNTs have no discernible morphological differences (Fig. 1a and S7†). The inter-shell spacing of pristine CNTs was 0.35 nm (Fig. S7a†),

which remained stable after calcination (Fig. S7b†) and when the CoPc was loaded onto the CNTs (Fig. 1a). No other phases indicative of CoPc were observed in the HRTEM images. Similarly, the fast Fourier transform patterns of the HRTEM images (Fig. S8†) only show the (002) and (100) crystalline planes of the CNTs, with no crystalline domains corresponding to CoPc.

Elemental mapping using energy dispersive spectroscopy (EDX), however, reveals a homogeneous distribution of cobalt and nitrogen atoms on the CNTs (Fig. 1b-d and S9†). Additionally, the Co K-edge X-ray absorption near edge structure (XANES) spectrum of CoPcCNT\_7.0% exhibits a decrease in the intensity of its 1s → 4p<sub>z</sub> transition peak at 7716 eV and a red shift at the rising edge of its 1s → 4p<sub>x,y</sub> transition at around 7723 eV, compared to that of bulk CoPc crystals (Fig. 1e).<sup>27,28</sup> These changes indicate a distortion of the  $D_{4h}$  planar symmetry of CoPc, due to interactions between CoPc and the CNTs.<sup>27,29</sup>

There are also differences between the Raman spectra of CoPcCNT\_7.0% and bulk CoPc crystals (Fig. 1f and S10†). Specifically, the spectrum of CoPcCNT\_7.0% shows smaller intensity ratios for several peaks against the peak of the pyrrole C=C stretching at 1537 cm<sup>-1</sup>. These peaks include those of the Co-N stretching at 240 cm<sup>-1</sup>, isoindole deformation at 589 cm<sup>-1</sup>, and macrocycle deformation at 750 cm<sup>-1</sup>.<sup>30</sup> We further observed a red shift in the bands of the pyrrole C=C stretching at 1537 cm<sup>-1</sup> and the isoindole deformation at 482 and 195 cm<sup>-1</sup>. The N 1s X-ray photoelectron spectrum (XPS) of CoPcCNT\_7.0% also exhibits a shift towards higher binding energy and an increase in the intensity of the shakeup satellites (S<sub>A</sub>) compared to that of the bulk CoPc crystals (Fig. 1g). The aforementioned changes in the Raman bands and N 1s XPS spectrum notably involve the nitrogen atoms of CoPc, which indicates that CoPc is likely to interact with the CNT surface *via* its nitrogen atoms.<sup>31-33</sup> All in all, our characterization results indicate that CoPc is molecularly anchored to the CNT surface.

### Co(i)Pc centers are catalytically active

The CO<sub>2</sub>RR activity of CoPcCNT\_7.0% was evaluated in 0.1 M KHCO<sub>3</sub> electrolyte saturated by 1 atm CO<sub>2</sub> in an H-type cell (Fig. 2a, S11 and Table S1†). Each constant-potential electrolysis process was conducted for 15 min. H<sub>2</sub> and CO products were quantified by online gas chromatography (GC), while methanol was identified by <sup>1</sup>H nuclear magnetic resonance spectroscopy and quantified by headspace GC (Fig. S11a†). Formaldehyde (CH<sub>2</sub>O) was quantified by ultraviolet-visible spectroscopy. The product distribution as a function of applied potential is consistent with that in Wang *et al.*'s work.<sup>9</sup>

From  $-0.4$  to  $-0.8$  V *vs.* the reversible hydrogen electrode (RHE; all potentials will hereafter be referenced to the RHE), CO was the sole CO<sub>2</sub>RR product. Methanol was observed with a FE of 7% at  $-0.9$  V. CO<sub>2</sub> as the carbon source for methanol was verified with <sup>13</sup>CO<sub>2</sub>RR experiments (Fig. S11b†). As the potential changed from  $-0.9$  to  $-1.1$  V, the FE of methanol increased to 35%, accompanied by an increase in the FE of H<sub>2</sub> from 6 to 33% and a decrease in the FE of CO from 82 to 32%. Formaldehyde was also detected with a FE of <4% (Fig. S11c and d†). This product has been recently identified as an intermediate of CO



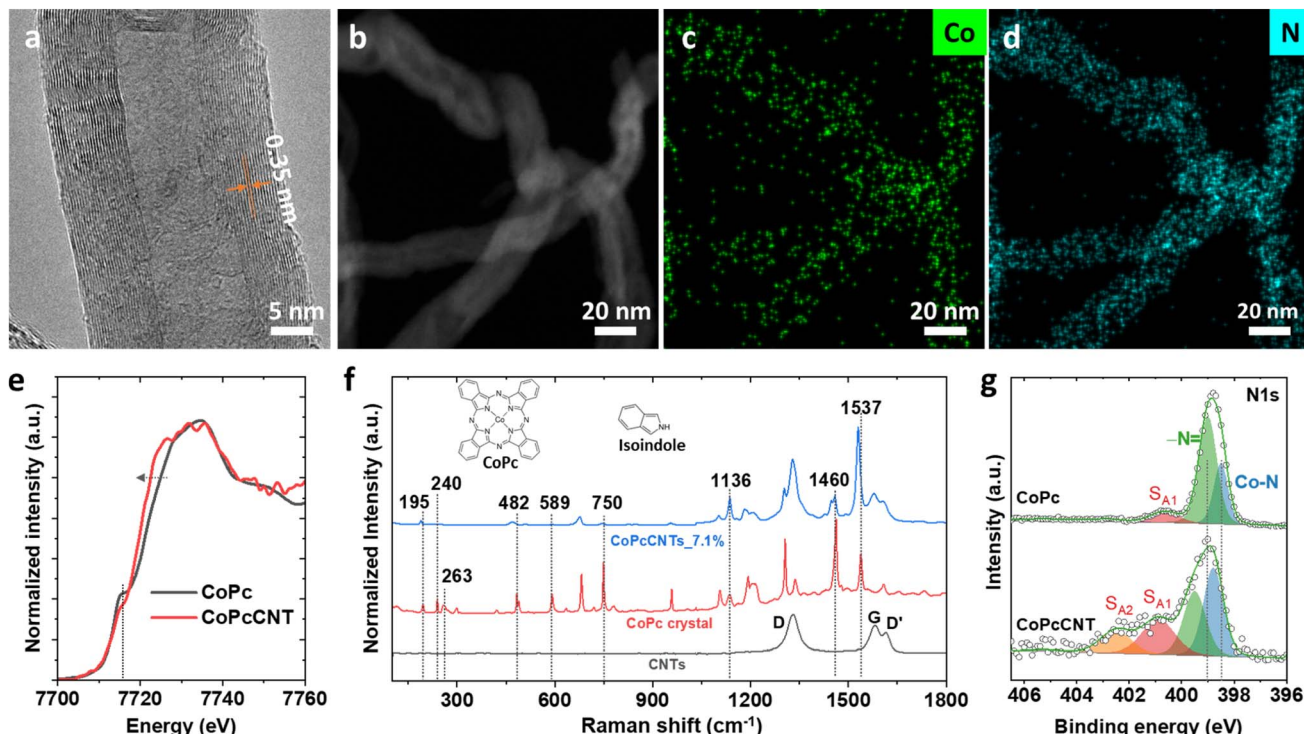


Fig. 1 (a) HRTEM image, (b) scanning transmission electron microscopy image and the (c and d) corresponding elemental maps of Co and N of as-synthesized CoPcCNT\_7.0%. (e) Co K-edge XANES spectra of CoPcCNT\_7.0% and bulk CoPc crystals. (f) Raman spectra of CoPcCNT\_7.0%, bulk CoPc crystals, and CNTs. (g) N 1s XPS spectra of bulk CoPc crystals and as-synthesized CoPcCNT\_7.0%.

reduction to methanol.<sup>12</sup> Overall, a minimum overpotential of ~0.8–0.9 V is needed to reduce CO<sub>2</sub> to methanol and formaldehyde, below which only CO is produced.

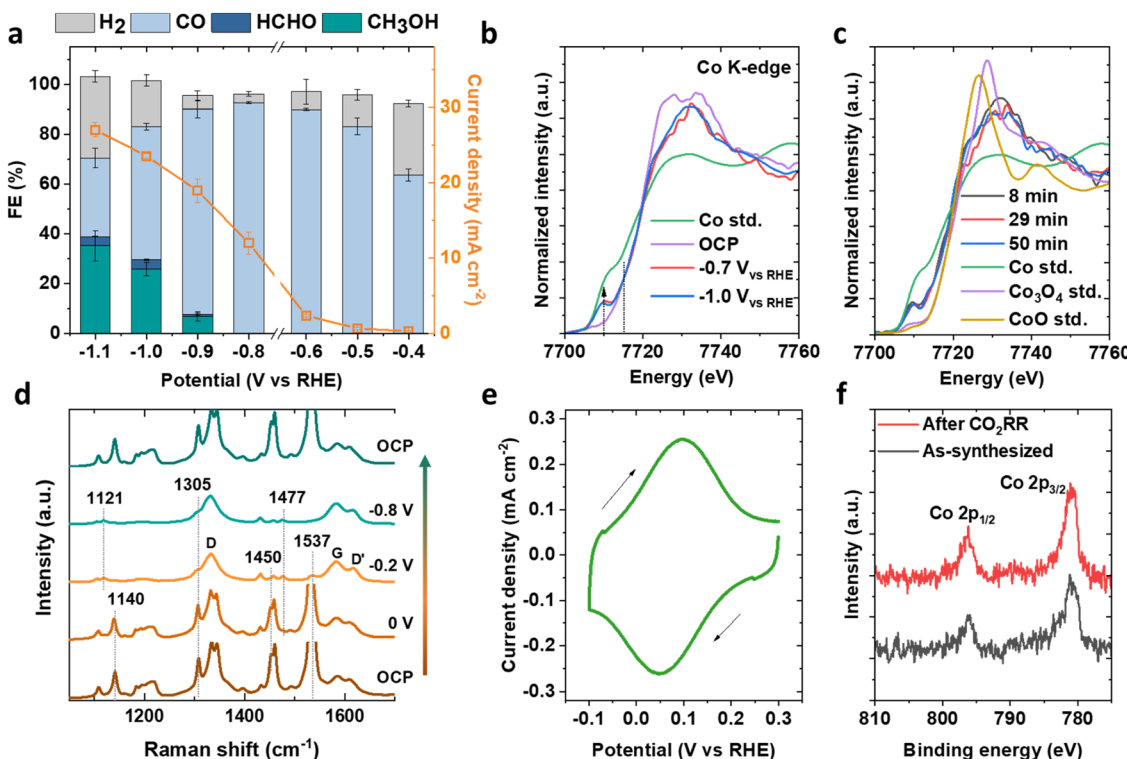
We conducted *in situ* XANES and Raman spectroscopy on CoPcCNT\_7.0% during the CO<sub>2</sub>RR. A new peak at 7710 eV in the Co K-edge XANES evolved at representative potentials of −0.7 and −1.0 V, which were the potentials at which CO and methanol were formed. By comparison with the edge feature of a metallic cobalt standard (Fig. 2b), the oxidation state of Co in CoPc appeared to be reducing from +2 towards zero.<sup>19,34</sup> However, the intensity of the newly formed peak remained fairly constant even after 50 min of electrolysis, suggesting that the metal in CoPc did not reduce to bulk metallic cobalt (Fig. 2c). Wang and coworkers recently assigned this peak to the Co 1s → 3d transition, asserting it as evidence for both the reduction of Co(II)Pc to Co(I)Pc and the binding of the CO intermediate to the Co center.<sup>19</sup>

The reduction of CoPc, as the potential changed from 0 to −0.2 V, was also indicated by changes in its Raman spectra (Fig. 2d). Specifically, its pyrrole breathing peak shifted from 1140 to 1121 cm<sup>−1</sup>, a pyrrole C–N peak at 1450 cm<sup>−1</sup> and a pyrrole C=C stretching peak at 1537 cm<sup>−1</sup> faded, and a pyrrole ring stretching peak at 1477 cm<sup>−1</sup> appeared. These changes persisted from −0.2 to −0.8 V. The peak shift from 1140 to 1121 cm<sup>−1</sup> is induced by the reduction of Co(II)Pc to Co(I)Pc, as the Co atom is bound to the nitrogen atoms in the pyrrole ring.<sup>30,35</sup> The decrease in the intensities of the bands at 1537 and 1477 cm<sup>−1</sup> can be attributed to the change in the

interaction between CoPc and CNTs, as the increased surface charge density of CNTs can weaken the molecule–substrate interaction.<sup>36</sup> The Raman spectrum reverted to its original state upon switching the potential from −0.8 V to the open circuit potential, indicating a reversibility of the aforementioned changes. The observed reduction of CoPc is also consistent with its redox wave of Co(n)/Co(l) at 0.1 V in its cyclic voltammogram (CV, Fig. 2e).<sup>37</sup> The Co 2p XPS spectrum of CoPcCNT\_7.0% post-CO<sub>2</sub>RR closely aligns with that of the as-prepared one, notably lacking the characteristic satellite feature (around 804 eV) associated with cobalt oxides (Fig. 2f).<sup>38</sup> The Co 2p<sub>3/2</sub> binding energy of CoPcCNT\_7.0% (780.9 eV) is also higher than that of metallic cobalt (778.3 eV).<sup>39</sup> Additionally, the electron microscopy images of the catalyst post-CO<sub>2</sub>RR show no features corresponding to bulk metallic cobalt or cobalt oxide particles (Fig. S12 and S13†). Therefore, we assign the newly formed peak in the Co K-edge XANES spectrum to CoPc with Co in the +1 state, which we propose to be catalytically active for the CO<sub>2</sub>RR.

#### Low CO<sub>2</sub> mass transfer positively impacts methanol selectivity

We separately investigated the effect of CoPc loading and the CO<sub>2</sub> partial pressure on the electroreduction of CO<sub>2</sub> to methanol on CoPc, and found that both factors impact the selectivity to methanol. Further analysis based on the effect of these two factors reveals an underlying relation: the average CO<sub>2</sub> mass transfer rate ( $\nu$ ) determines the ratio ( $R$ ) of the production rates of methanol and CO. In fact, a power function ( $R = 0.97\nu^{-0.78}$ )



**Fig. 2** (a) Faradaic efficiencies of CO<sub>2</sub>RR products and the total current density as a function of the applied potential on CoPcCNT\_7.0%. (b) *In situ* XANES spectra of CoPcCNT\_7.0% at -0.7 and -1.0 V. OCP denotes open circuit potential and 'Co std.' denotes a Co metal standard. (c) Evolution of the *in situ* XANES spectra over reaction times at -1.0 V. Co std., Co<sub>3</sub>O<sub>4</sub> std., and CoO std. are respectively the Co metal, Co<sub>3</sub>O<sub>4</sub>, and CoO standards. (d) *In situ* Raman spectra of CoPcCNT\_7.0% measured at various potentials. The peak at 1537 cm<sup>-1</sup> is truncated when its signal is too intense, so as to make the comparison clearer. (e) Cyclic voltammograms of CoPcCNT\_7.0%. The scan rate is 50 mV s<sup>-1</sup>. (f) Co 2p XPS spectra of CoPcCNT\_7.0% before and after its use for the 15 min CO<sub>2</sub>RR experiment at -1.0 V. CO<sub>2</sub>-saturated 0.1 M KHCO<sub>3</sub> electrolyte was used for all the aforementioned experiments.

fits the data from these two investigations with a high correlation coefficient of 0.948.

In detail, we first evaluate CoPcCNT\_x% samples with CoPc loading from 0.2 to 7.0% for the CO<sub>2</sub>RR at -1.0 V in an H-cell. Aqueous 0.1 M KHCO<sub>3</sub> saturated with 1 atm CO<sub>2</sub> was used as the electrolyte. CoPcCNT has rather poor stability when catalyzing the CO<sub>2</sub>RR to methanol due to the possible CoPc detachment and decomposition at a negative potential.<sup>9</sup> We observed a small increase in the H<sub>2</sub> FE (such as from 4.9 to 5.8% on CoPcCNT\_1.9%) during 15 min of electrolysis, similar to what was reported in Wang's work.<sup>9</sup> Hence, to minimize the effect of CoPcCNT instability on the performance assessment, we conducted the electrolysis for only 15 min (Fig. S14†).

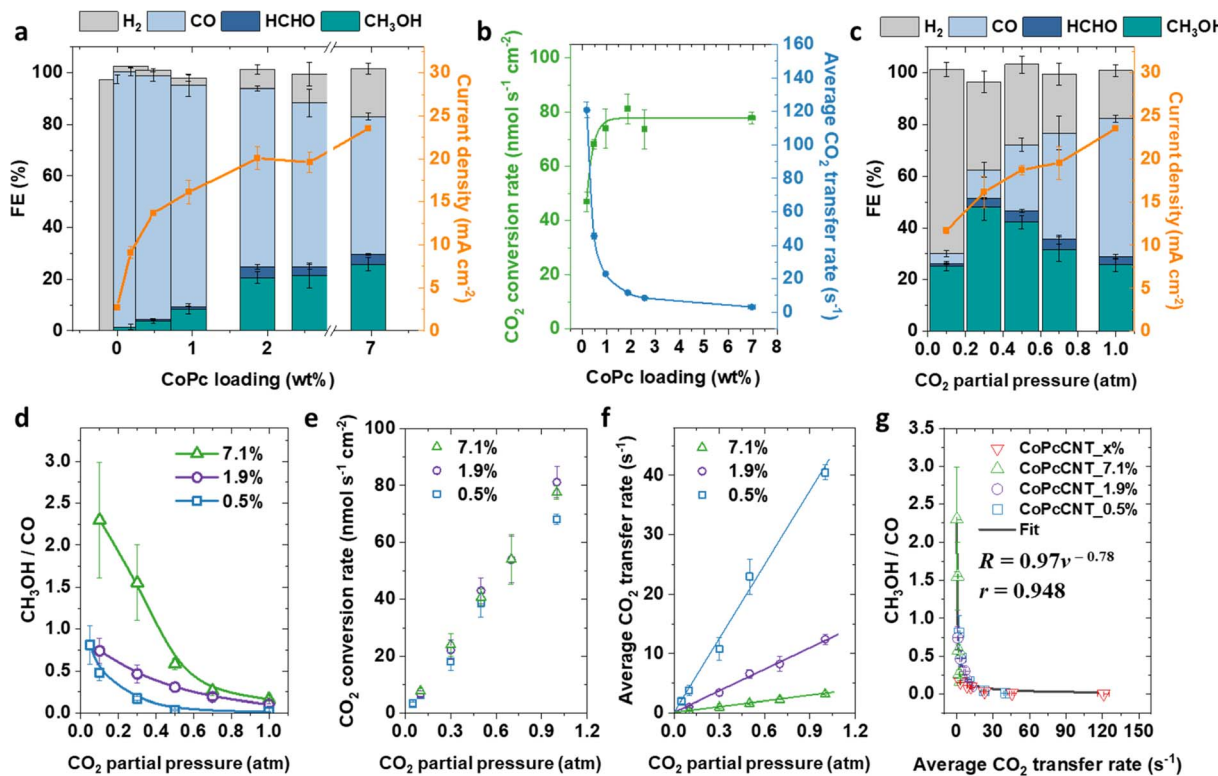
The product distribution at various CoPc loadings is shown in Fig. 3a. As the CoPc loading on the CNTs increased from 0.2 to 7.0%, the FE of methanol increased from 2 to 26% (Fig. 3a and Table S2†). The FEs of formaldehyde and H<sub>2</sub> also increased to 4 and 19%, respectively. Correspondingly, the FE of CO decreased from 99 to 53%. Electrolysis on bare CNTs produced only H<sub>2</sub>, indicating that the CNTs themselves are inactive for the CO<sub>2</sub>RR (Fig. 3a). We note that Wang and coworkers' recent study has also disclosed a positive correlation between FE<sub>methanol</sub> and CoPc loading.<sup>19</sup>

Further analysis reveals that the total CO<sub>2</sub> conversion rate increased from ~47 nmol s<sup>-1</sup> cm<sup>-2</sup> on CoPcCNT\_0.2%, and

plateaued at ~77 nmol s<sup>-1</sup> cm<sup>-2</sup> when the CoPc loading exceeded 1% (Fig. 3b; See Section S1.2.7 for a sample calculation). The plateau indicates that the CO<sub>2</sub> conversion rate is limited by CO<sub>2</sub> mass transfer to the CoPc sites. Since the limiting CO<sub>2</sub> mass transfer rate (77 nmol s<sup>-1</sup> cm<sup>-2</sup>) to the electrode should be the same for all CoPcCNT\_x% samples, an increase in CoPc loading from 0.2 to 7.0% will lower the average CO<sub>2</sub> transfer rate to each CoPc molecular complex from 121 to 3 s<sup>-1</sup>. As no new catalytic species such as Co clusters were detected, we hypothesize that the CO<sub>2</sub> mass transfer to each CoPc site must have played a significant role in its subsequent conversion to methanol or CO.

To evaluate our hypothesis on the effect of CO<sub>2</sub> mass transfer on CO<sub>2</sub>RR selectivity, we controlled the CO<sub>2</sub> mass transfer by varying the CO<sub>2</sub> partial pressures (pCO<sub>2</sub>) on three catalysts, namely CoPcCNT\_0.5%, CoPcCNT\_1.9%, and CoPcCNT\_7.0%, at around -0.7 V (corrected by the local pH, Table S6†). On all three catalysts, the FE<sub>methanol</sub> and pCO<sub>2</sub> exhibited non-monotonic correlations (Fig. 3c, S15 and Tables S3–S5†). It is also noteworthy that the maximum FE<sub>methanol</sub> was not obtained at 1 atm, but at 0.1–0.5 atm of CO<sub>2</sub>. FE<sub>CO</sub>, in contrast, increased with pCO<sub>2</sub>. It is further notable that while methanol could be generated at -0.8 V when the pCO<sub>2</sub> is 0.2 atm, no methanol could be detected at 1 atm of CO<sub>2</sub> (Fig. 2a and S16†). This phenomenon confirms that CO<sub>2</sub> mass transfer indeed affects the production of methanol, but not in a monotonic way.





**Fig. 3** Faradaic efficiencies of CO<sub>2</sub>RR products and total current density as a function of the CoPc loading in the CoPcCNT catalysts at  $-1.0$  V and  $1$  atm CO<sub>2</sub>. (b) CO<sub>2</sub> conversion rate and average CO<sub>2</sub> transfer rate to each CoPc as a function of the CoPc loading. (c) FE of CO<sub>2</sub>RR products and total current density as a function of the CO<sub>2</sub> partial pressure (pCO<sub>2</sub>) on CoPcCNT\_7.0% at  $-0.7$  V (corrected based on the local pH). (d) Ratio of the production rates of CH<sub>3</sub>OH and CO versus pCO<sub>2</sub> on CoPcCNT\_0.5%, CoPcCNT\_1.9%, and CoPcCNT\_7.0%. (e) CO<sub>2</sub> conversion rate and (f) average CO<sub>2</sub> transfer rate to each CoPc as a function of pCO<sub>2</sub> on CoPcCNT\_0.5%, CoPcCNT\_1.9%, and CoPcCNT\_7.0%. (g) Ratio of the production rates of CH<sub>3</sub>OH and CO as a function of the average CO<sub>2</sub> transfer rate to each CoPc obtained from four datasets. CoPcCNT\_0.5%, CoPcCNT\_1.9%, and CoPcCNT\_7.0% denote the experiments where CO<sub>2</sub> partial pressure is controlled with the results corresponding to the three curves in (3d), respectively. CoPcCNT\_x% denotes the experiments where the CoPc loading is controlled with x% ranging from 0.2 to 7.0%, corresponding to the blue curve in (3b). 'Fit' denotes the black curve derived from the nonlinear curve fitting with the equation  $R = 0.97v^{-0.78}$ , where  $R$  is the CH<sub>3</sub>OH/CO ratio and  $v$  is the CO<sub>2</sub> transfer rate to each CoPc. 'r' denotes the correlation coefficient of the fit. All CO<sub>2</sub>RR experiments for (a–g) were conducted at  $-1.0$  V in  $0.1$  M KHCO<sub>3</sub> saturated with CO<sub>2</sub> with the pressure noted in the graphs or in this caption.

Further analysis of all three catalysts reveals a negative correlation between pCO<sub>2</sub> and the ratio of the production rates (mole s<sup>-1</sup> cm<sup>-2</sup>) of CH<sub>3</sub>OH and CO (Fig. 3d). The CH<sub>3</sub>OH/CO ratio also increased with CoPc loading. For instance, at a pCO<sub>2</sub> of  $0.3$  atm, when the CoPc loading increased from  $0.5$  to  $7.0\%$ , the value of CH<sub>3</sub>OH/CO increased  $\sim 7\times$  from  $0.2$  to  $1.6$ . These correlations imply the competitive formation of methanol and CO from the CO<sub>2</sub>RR, and this competition is modulated by both pCO<sub>2</sub> and the CoPc loading. Specifically, we believe that the CO<sub>2</sub> transfer rate to each CoPc directly influences the CH<sub>3</sub>OH/CO ratio.

To verify the validity of this conclusion, we first note that the CO<sub>2</sub> conversion rate on all three catalysts is linearly related to pCO<sub>2</sub> (Fig. 3e), which indicates that the reaction rate of the CO<sub>2</sub>RR is limited by CO<sub>2</sub> mass transfer. This in turn suggests that the total CO<sub>2</sub> transfer to the electrode has the same rate as CO<sub>2</sub> conversion. The CO<sub>2</sub> transfer rate to each CoPc complex can thus be calculated by dividing the CO<sub>2</sub> conversion rate by the CoPc loading (Fig. 3f and Section S1.2.7). We then plotted the curves of the CH<sub>3</sub>OH/CO ratio ( $R$ ) versus the CO<sub>2</sub> transfer rate to each CoPc ( $v$ ) in the same graph (Fig. 3g). Remarkably, the curves from four

distinct datasets, either by controlling CoPc loading or pCO<sub>2</sub>, follow a consistent trend, and fit in a power-law function ( $R = 0.97v^{-0.78}$ ). This behavior reveals that the CO<sub>2</sub> transfer rate to each CoPc controls the competition between methanol and CO formation. More specifically, CO<sub>2</sub> may only be reduced to methanol when the supply of CO<sub>2</sub> is low.

### Mechanistic considerations and computational modelling

Thus far, we have identified CO, formaldehyde and methanol as CO<sub>2</sub>RR products on CoPcCNT (Fig. 2a). We have also electro-reduced CO on CoPcCNT\_7% and found that it can be converted to CH<sub>2</sub>O and CH<sub>3</sub>OH. CH<sub>2</sub>O itself can be reduced to methanol (Fig. S17a and Note S1†). On the other hand, the electrolysis of formate on CoPcCNT did not yield methanol (Fig. S17b†).<sup>10</sup>

We performed density-functional theory (DFT) calculations to help connect these observations, elaborate a reaction pathway that complies with all of them, and provide simple catalyst design guidelines for future studies. In line with a previous study,<sup>13</sup> we found that CoPc under negative

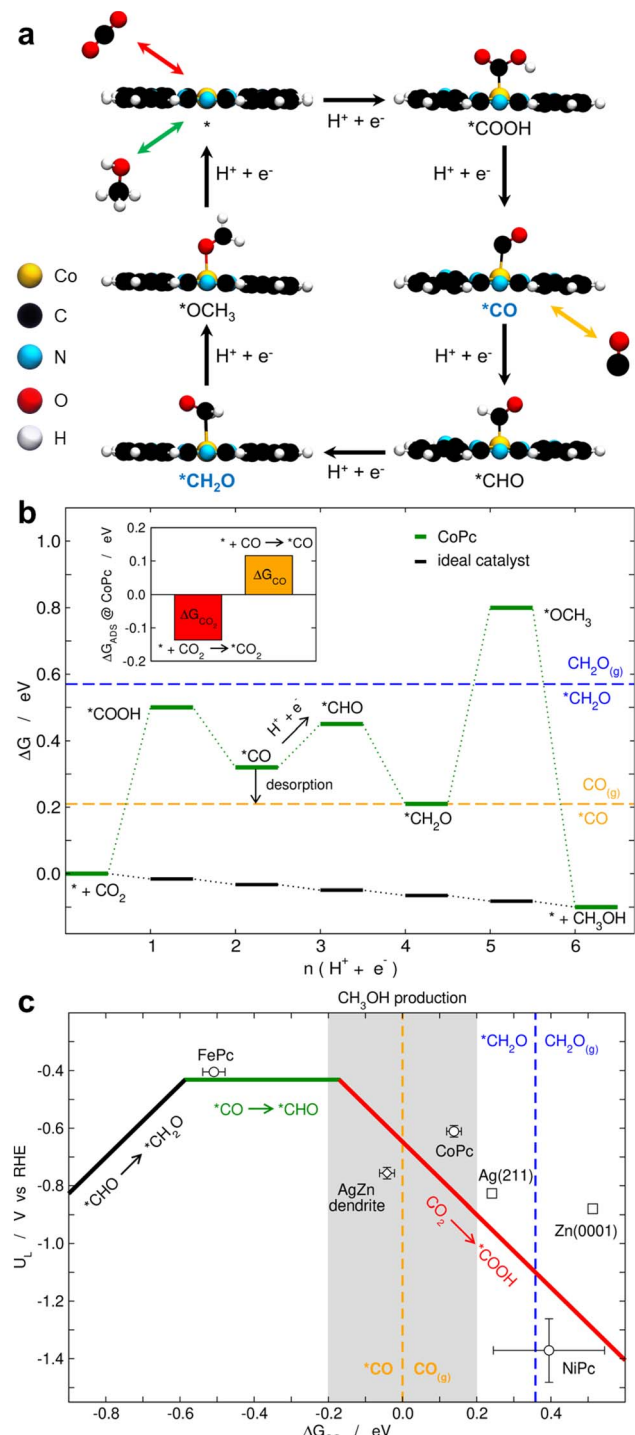


Fig. 4 (a) Side views of the lowest-energy electrochemical CO<sub>2</sub>RR intermediates to methanol on CoPc. Top views of the catalyst and the intermediates are provided in Fig. S18.† CO desorption and re-adsorption are possible upon the second proton-electron transfer, as indicated by the orange arrow. \*CO and \*CH<sub>2</sub>O appear in blue as CO(g) and CH<sub>2</sub>O(g) were shown in experiments to be reducible to methanol. (b) Free-energy diagram for CO<sub>2</sub> electroreduction to methanol on CoPc at 0 V. The equilibrium adsorption/desorption lines for CO and CH<sub>2</sub>O are shown in orange and blue, respectively. For comparison, the free-energy landscape for the ideal catalyst is also provided. Inset: Adsorption energies of CO<sub>2</sub> and CO on CoPc at 0 V. (c) Volcano plot for the CO<sub>2</sub>RR to methanol on FePc, CoPc and NiPc. Ag(211), Zn(0001) and Ag dendrites on Zn are also provided for

potentials is reduced and contains a -H ligand on one of the N atoms (Fig. S18†). CoPc may also contain an extra electron under such conditions,<sup>13</sup> but our calculations indicate that adding or removing it shifts the free energies of the intermediates on average by only  $0.06 \pm 0.03$  eV and does not modify our conclusions (Fig. S19 and Table S7†). After inspecting each electrochemical step for possible bifurcations, the lowest-energy pathway for the CO<sub>2</sub>RR to methanol is: CO<sub>2</sub> → \*COOH → \*CO → \*CHO → \*CH<sub>2</sub>O → \*OCH<sub>3</sub> → CH<sub>3</sub>OH, which agrees with previous studies (Fig. 4a and S19†).<sup>13,26</sup>

In Fig. 4b, we provide a free-energy diagram for the pathway in Fig. 4a on CoPc in which the equilibrium lines for CO (orange) and CH<sub>2</sub>O (blue) adsorption are shown. The potential-limiting step is the hydrogenation of \*CH<sub>2</sub>O to \*OCH<sub>3</sub>, and the thermodynamic potential to overcome it is  $-0.59$  V. The first step, namely, the formation of \*COOH requires a slightly lower potential ( $-0.50$  V). Upon the second proton-electron transfer, \*CO can either desorb as CO<sub>(g)</sub> or be further reduced to methanol *via* \*CHO. This indicates that the extent of \*CO adsorption/desorption is crucial in determining the selectivity of the CO<sub>2</sub>RR toward carbon monoxide or methanol. In the event that a \*CO desorbs, the chances for it to re-adsorb onto an active site and hydrogenate to methanol will decrease if a species with a stronger adsorption energy is nearby. According to our calculations, such a species is likely CO<sub>2</sub>, as its adsorption energy is  $-0.14$  eV, which is more negative than that of any other adsorbed species on CoPc, including \*CO and \*CH<sub>2</sub>O (Fig. 4b and Table S7†). In particular, the inset of Fig. 4b shows that CO<sub>2</sub> adsorption is stronger than that of CO on CoPc at 0 V.

Several previous studies have reached a similar conclusion to ours that CO<sub>2</sub> adsorption is stronger than that of CO on CoPc.<sup>13,26</sup> In particular, the microkinetic analysis conducted by McCrory and coworkers indicated a 3.3-fold increase in the effective equilibrium binding constant for CO<sub>2</sub>-CoPc compared to CO-CoPc.<sup>26</sup> However, it is worth noting that such an increase in the binding constant was supported by a calculated binding energy difference between CO<sub>2</sub> and CO of only  $0.03$  eV (0.6 kcal mol<sup>-1</sup>). As DFT generally has an accuracy of no less than  $\pm 0.1$  eV, even when hybrid functionals are used, the aforementioned agreement between the experiment and theory should be considered cautiously. The inaccuracies can be aggravated when gas-phase errors in *e.g.* CO<sub>2</sub>, CO, CH<sub>2</sub>O and CH<sub>3</sub>OH are not accounted for,<sup>41,42</sup> and the impact can be observed not just in the adsorption energies but also in the predicted equilibrium and onset potentials. In this case, we used gas-phase energy corrections to ensure that (i) our DFT-predicted equilibrium potentials and reaction energies match the experimental ones (see Section S1.5 and Fig. S20†), and (ii) an accurate gas-phase reference is used for the adsorption/desorption energies in the free-energy diagrams. The qualitative and quantitative consequences of omitting gas-phase corrections are illustrated in the

comparison (AgZn).<sup>40</sup> The datapoints are averages of the elementary steps on the phthalocyanines with and without extra electrons, and the average of two dendrites for AgZn. The grey area, where methanol production is enabled, corresponds to  $\Delta G_{CO} = 0 \pm 0.2$  eV.

free-energy diagrams for CoPc and the ideal catalyst in Fig. S21.† Such considerations are still often overlooked in computational models and may severely limit their predictiveness.<sup>42</sup>

If \*CO binding is to be strengthened to obtain more methanol, it is pertinent to ask by how much. To address this question and bearing in mind that previous studies found no liquid products on FePc and NiPc during the CO<sub>2</sub>RR,<sup>9,13</sup> we modelled the pathway in Fig. 4a on these two phthalocyanines by means of DFT calculations and elaborated a Sabatier-type activity plot in Fig. 4c. The figure shows the potentials required by the various limiting steps ( $U_L$ ) of the phthalocyanines as a function of their free energies of adsorption of \*CO. The phthalocyanine datapoints in Fig. 4c correspond to averages of charged and uncharged systems (Table S7†). We note that the narrow error bars indicate that the presence or absence of an extra electron at the metal-Pc sites does not change our conclusions in this case. This is also illustrated in the free-energy diagrams in Fig. S19 and in Table S7.†

Unlike the CO<sub>2</sub>RR volcano plot for the production of methane, which has a sharp summit,<sup>43</sup> we observe here a relatively wide plateau where \*CO hydrogenation is potential-limiting. The methanol activity of FePc is limited by this step, while that of NiPc is limited by \*COOH formation. As mentioned before, CoPc is limited by the formation of \*OCH<sub>3</sub>, but \*COOH formation requires a similar potential (−0.59 vs. −0.50 V). The experimental inertness of NiPc can be explained by its weak binding of the reaction intermediates,<sup>9,13</sup> which results in an average limiting potential of around −1.3 V. In turn, FePc is predicted to be active but experiments show otherwise,<sup>9</sup> suggesting that \*CO hydrogenation is kinetically impeded. This helps us formulate a hypothesis: active materials for the CO<sub>2</sub>RR to methanol ought to be found at around  $\Delta G_{CO} = 0 \pm 0.2$  eV, that is within the grey-colored zone in Fig. 4c. To the right of it, \*CO and eventually \*CH<sub>2</sub>O adsorption are too weak and the CO<sub>2</sub>RR only leads to CO(g). To the left of the grey zone,

the plateau prevents an improvement of the limiting potential when \*CO adsorption is strengthened, and \*CO hydrogenation might be kinetically impeded, such that CO(g) will also be produced.

To test this hypothesis, we turn to our previous work, where Ag and Zn were shown to reduce CO<sub>2</sub> mainly to CO, but Zn dendrites deposited on Ag foam were able to reduce CO<sub>2</sub> to methanol with an experimental faradaic efficiency of 10.5%.<sup>40</sup> We observe exactly this in Fig. 4c: Ag(211) and Zn(0001) bind \*CO too weakly to be within the methanol production (grey) zone, while AgZn dendrites bind \*CO strong enough to be within it. In perspective, our simple classification of the active sites in terms of \*CO binding is similar to other classifications for CO<sub>2</sub>RR products<sup>44,45</sup> and, certainly, its exciting predictions call for future experiments and calculations.

While we have no direct experimental evidence supporting the stronger CO<sub>2</sub> adsorption on CoPc as compared to CO, our *in situ* Raman spectroscopy and CVs of CoPcCNT in CO<sub>2</sub>-, CO- and N<sub>2</sub>-saturated electrolytes indicate a different interaction of CO<sub>2</sub> and CO with CoPc. At −0.2 V, the Raman bands of CoPc in CO<sub>2</sub>-saturated electrolyte were more attenuated than those in CO- and N<sub>2</sub>-saturated electrolytes (Fig. 5a and S22, and Note S2†). In the CVs of CoPcCNT\_7.0%, the oxidation waves (A) and (B) at around 0.1 and 0.9 V, respectively, were suppressed in CO<sub>2</sub>-saturated electrolyte, relative to that in CO- and N<sub>2</sub>-saturated electrolytes (Fig. 5b and S23, and Note S3†). To directly show the competing binding of CO<sub>2</sub> and CO and evaluate the dwelling time of \*CO, more advanced electrochemical spectroscopic methods, such as *in situ* time-resolved infrared spectroscopy, should be used to quantify the binding strength of CO<sub>2</sub> and CO on CoPc. These measurements may also reveal more microkinetic details and the physical model behind the power-law function as shown in Fig. 3g.

On the basis of the above results, to enhance methanol production, it is necessary to moderately enhance the \*CO adsorption energy of CoPc to prevent CO desorption, but still

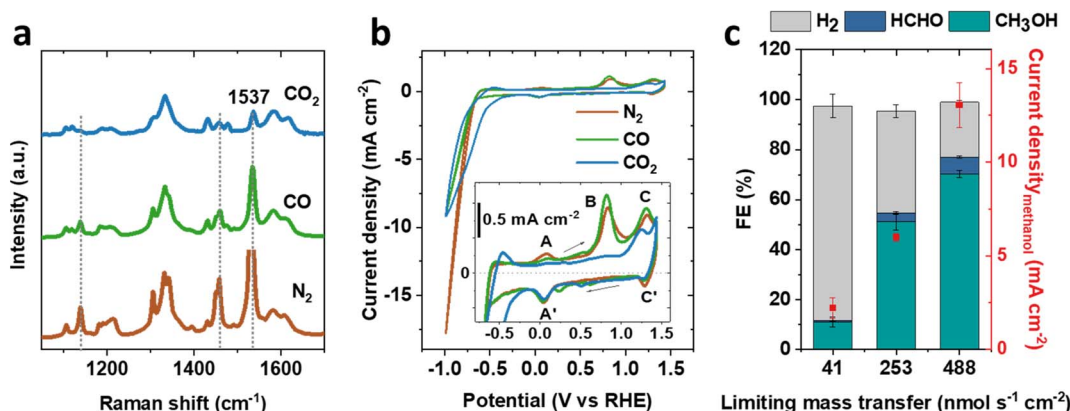


Fig. 5 (a) *In situ* Raman spectra of CoPcCNT\_7.0% measured at −0.2 V in phosphate buffers (pH 7) saturated by N<sub>2</sub>, CO, or CO<sub>2</sub>. The 1537 cm<sup>−1</sup> peak in the spectrum measured in N<sub>2</sub>-saturated electrolyte is truncated as its signal is too intense (see the full spectra in Fig. S22†). (b) Cyclic voltammograms of CoPcCNT\_7.0% in 0.1 M KClO<sub>4</sub> solution saturated by N<sub>2</sub>, CO, or CO<sub>2</sub>. The A and A' peaks correspond to CoPc(II)/CoPc(I) and the C and C' peaks to CoPc(II)/CoPc(III).<sup>37</sup> The scan rate was 50 mV s<sup>−1</sup>. (c) Faradaic efficiencies of the CORR products and partial current density of methanol as a function of the limiting CO mass transfer to the electrode at −0.86 V. The electrolyte used was 0.1 M KHCO<sub>3</sub> saturated by 1 atm of CO.



allow its hydrogenation. It is also necessary to reduce the competitive adsorption of  $\text{CO}_2$ , so that  $^*\text{CO}$  could be reduced to methanol. This can be controlled by decreasing the mass transfer of the  $\text{CO}_2$  reactant to the CoPcCNT catalysts as shown in Fig. 3.

Further directions for optimizing the selectivity of CoPc towards methanol can be obtained by comparing its free-energy diagram to that of the ideal catalyst. As shown in Fig. 4b, such a hypothetical catalyst has energy-symmetric electrochemical steps consecutively separated by 16.4 meV, which would allow it to operate with no energy losses at the equilibrium potential, in principle. A comparison of the two energy landscapes in Fig. 4b indicates that CoPc is relatively efficient for methanol production but is far from ideal. In particular, the first and fifth electrochemical steps are rather energy-intensive and the free energies of the adsorbates are visibly weaker than those of the ideal catalyst. For example, the ideal  $^*\text{COOH}$  level with respect to  $\text{CO}_2$  should be at  $-0.02$  eV, but it is  $0.50$  eV on CoPc.

Our results indicate that using CO as a feedstock is a more straightforward approach to boost the selectivity toward methanol, as no  $\text{CO}_2$  could then compete with CO for the active sites. Thus, we evaluated the CORR on CoPcCNT\_7.0% under different mass transfer conditions (Fig. 5c, S1 and S2†). The selectivity and activity of methanol and formaldehyde production now positively correlate with the mass transfer of CO, unlike that of  $\text{CO}_2$ . Faradaic efficiencies of 70% for methanol and 7% for formaldehyde could be achieved from the CORR at  $-0.86$  V and at a total current density of  $-19$  mA  $\text{cm}^{-2}$  in  $0.1$  M  $\text{KHCO}_3$  electrolyte. The stability of CoPcCNT for the CORR at  $-0.86$  V in a flow cell was relatively poor, as evidenced by the increase in  $\text{H}_2$  FE from 20 to 36% and the decrease in the cathodic current density from around 13 to 10 mA  $\text{cm}^{-2}$  during a 2 h electrolysis (Fig. S24†). The instability of the CORR could be caused by the detachment of CoPc from the CNT support or by the decomposition of CoPc complexes, as observed in Wang's work.<sup>9</sup> The catalyst stability could be improved by adding a functional group or ligand that enhances the anchoring of CoPc, or by optimizing the diameter of the CNT support.<sup>9,24,46</sup> Moreover, unlike metal catalysts mounted in flow cells that often exhibit current densities larger than 100 mA  $\text{cm}^{-2}$ ,<sup>47</sup> the CORR on CoPcCNT in a flow cell still exhibits a rather small current density of  $\sim 13$  mA  $\text{cm}^{-2}$ . This is likely due to the easy wetting of CoPcCNT by the aqueous electrolyte that results in only a small fraction of CoPc active sites being accessible to CO.<sup>48</sup> The microenvironment in the catalyst layer may be optimized to mitigate this issue and, hence, increase the current density.<sup>23,48</sup>

Overall, we have elucidated in this work that (1) Co(i)Pc is the likely catalytically active species under  $\text{CO}_2$  reduction conditions, and (2) the  $\text{CO}_2$  transfer rate to each CoPc together with the overpotential inclines the  $\text{CO}_2$ RR toward either methanol or CO. The underlying reason is that  $\text{CO}_2$  mass transfer modulates the coverage of  $^*\text{CO}_2$  and  $^*\text{CO}$ . These findings, which align with those from several other recent studies,<sup>13,19,26</sup> suggest two approaches to modulate the selectivity of methanol and CO from the  $\text{CO}_2$ RR on CoPc: first, by adjusting the CoPc loading in the electrode, and second, by regulating the mass transfer of

$\text{CO}_2$ . We note that similar mass-transfer effects have previously been reported on the copper-catalyzed CORR, showing that the competitive production of ethylene, acetate, and n-propanol may be correlated with the mass transfer of CO.<sup>49,50</sup> However, the underlying mechanism still remains a matter of debate. Herein, our findings highlight the crucial role of adsorption intermediates from a new perspective, as on CoPc an excess or lack of  $\text{CO}_2$  around the active sites suffices to prevent or enable CO adsorption and hydrogenation.

## Conclusions

In this work, by means of *in situ* XANES and *in situ* Raman spectroscopies, we determined that reduced CoPc with Co(i) centers are the active sites for the  $\text{CO}_2$ RR. We found that the selectivity to methanol is positively correlated with the CoPc loading. The negative correlation between the ratio of the  $\text{CH}_3\text{OH}/\text{CO}$  production rate and the  $\text{CO}_2$  partial pressure further led us to identify the  $\text{CO}_2$  transfer rate to each CoPc as a key factor for methanol production relative to CO. This rate is crucial in view of the competitive adsorption of  $\text{CO}_2$  and CO on CoPc, inferred from experiments and corroborated by DFT calculations. Inspired by this finding, we tuned the mass transfer of CO to each CoPc molecule, which enabled us to reach a 70% faradaic efficiency and a partial current density of  $13$  mA  $\text{cm}^{-2}$  for methanol production from the CORR. Computational modelling based on the Sabatier principle revealed that the  $\text{CO}_2$ RR on CoPc can be enhanced by strengthening the binding of  $^*\text{CO}$  under the condition that  $-0.2$  eV  $< \Delta G_{\text{CO}} < 0.2$  eV, and by making the free-energy diagram more energetically symmetric along the entire  $\text{CO}_2$ RR catalytic pathway. This body of work provides valuable guidelines and inspiration for developing efficient metal complex-based catalysts, as well as for identifying the reaction conditions that optimize their activity and selectivity.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

The authors thank the National Research Foundation of Singapore (Urban Solutions and Sustainability, Industry Alignment Fund (Pre-Positioning) Programme, A-0004543-00-00), and Shell Global Solutions International B.V. (A-0004543-01-00) for financial support of this project. FCV received financial support from grants PID2021-127957NB-I00 and TED2021-132550B-C21, which are funded by MCIN/AEI/10.13039/501100011033 and by the European Union. FCV also acknowledges financial support through grant IT1453-22 "Grupos Consolidados UPV/EHU del Gobierno Vasco" and thanks the Red Española de Supercomputación for computational resources (grants QHS-2023-2-0007 and QHS-2023-3-0006). The use of supercomputing facilities at SURFsara was sponsored by NWO Physical Sciences, with financial support by NWO. The authors thank Dr Jia Liu from



the Department of Chemistry, National University of Singapore, for useful scientific discussions.

## References

- P. De Luna, C. Hahn, D. Higgins, S. A. Jaffer, T. F. Jaramillo and E. H. Sargent, *Science*, 2019, **364**, eaav3506.
- S. Sarp, S. Gonzalez Hernandez, C. Chen and S. W. Sheehan, *Joule*, 2021, **5**, 59–76.
- H. Choi, D.-K. Lee, M.-K. Han, G. Janani, S. Surendran, J. H. Kim, J. K. Kim, H. Cho and U. Sim, *J. Electrochem. Soc.*, 2020, **167**, 164503.
- Y. Y. Birdja, E. Pérez-Gallent, M. C. Figueiredo, A. J. Göttle, F. Calle-Vallejo and M. T. M. Koper, *Nat. Energy*, 2019, **4**, 732–745.
- S. Nitopi, E. Bertheussen, S. B. Scott, X. Liu, A. K. Engstfeld, S. Horch, B. Seger, I. E. L. Stephens, K. Chan, C. Hahn, J. K. Nørskov, T. F. Jaramillo and I. Chorkendorff, *Chem. Rev.*, 2019, **119**, 7610–7672.
- Q. H. Low and B. S. Yeo, *J. Electrochem. Energy Convers. Storage*, 2020, **17**, 040802.
- I. E. L. Stephens, K. Chan, A. Bagger, S. W. Boettcher, J. Bonin, E. Boutin, A. K. Buckley, R. Buonsanti, E. R. Cave, X. Chang, S. W. Chee, A. H. M. da Silva, P. de Luna, O. Einsle, B. Endrődi, M. Escudero-Escribano, J. V. F. de Araujo, M. C. Figueiredo, C. Hahn, K. U. Hansen, S. Haussener, S. Hunegnaw, Z. Huo, Y. J. Hwang, C. Janáky, B. S. Jayathilake, F. Jiao, Z. P. Jovanov, P. Karimi, M. T. M. Koper, K. P. Kuhl, W. H. Lee, Z. Liang, X. Liu, S. Ma, M. Ma, H.-S. Oh, M. Robert, B. R. Cuenya, J. Rossmeisl, C. Roy, M. P. Ryan, E. H. Sargent, P. Sebastián-Pascual, B. Seger, L. Steier, P. Strasser, A. S. Varela, R. E. Vos, X. Wang, B. Xu, H. Yadegari and Y. Zhou, *JPhys Energy*, 2022, **4**, 042003.
- Y. Liu, F. Li, X. Zhang and X. Ji, *Curr. Opin. Green Sustainable Chem.*, 2020, **23**, 10–17.
- Y. Wu, Z. Jiang, X. Lu, Y. Liang and H. Wang, *Nature*, 2019, **575**, 639–642.
- E. Boutin, M. Wang, J. C. Lin, M. Mesnage, D. Mendoza, B. Lassalle-Kaiser, C. Hahn, T. F. Jaramillo and M. Robert, *Angew. Chem., Int. Ed.*, 2019, **58**, 16172–16176.
- Y. Wu, G. Hu, C. L. Rooney, G. W. Brudvig and H. Wang, *ChemSusChem*, 2020, **13**, 6296–6299.
- E. Boutin, A. Salamé, L. Merakeb, T. Chatterjee and M. Robert, *Chem.-Eur. J.*, 2022, **28**, e202200697.
- X. Ren, J. Zhao, X. Li, J. Shao, B. Pan, A. Salamé, E. Boutin, T. Groizard, S. Wang, J. Ding, X. Zhang, W.-Y. Huang, W.-J. Zeng, C. Liu, Y. Li, S.-F. Hung, Y. Huang, M. Robert and B. Liu, *Nat. Commun.*, 2023, **14**, 1–10.
- S. Yang, Y. Yu, X. Gao, Z. Zhang and F. Wang, *Chem. Soc. Rev.*, 2021, **50**, 12985–13011.
- Z. Yue, C. Ou, N. Ding, L. Tao, J. Zhao and J. Chen, *ChemCatChem*, 2020, **12**, 6103–6130.
- D. Grammatico, A. J. Bagnall, L. Riccardi, M. Fontecave, B.-L. Su and L. Billon, *Angew. Chem., Int. Ed.*, 2022, **61**, e202206399.
- Q. Feng, Y. Sun, X. Gu and Z. Dong, *Electrocatalysis*, 2022, **13**, 675–690.
- S. Zhang, X. Jing, Y. Wang and F. Li, *ChemNanoMat*, 2021, **7**, 728–736.
- C. L. Rooney, M. Lyons, Y. Wu, G. Hu, M. Wang, C. Choi, Y. Gao, C.-W. Chang, G. W. Brudvig, Z. Feng and H. Wang, *Angew. Chem., Int. Ed.*, 2024, **63**, e202310623.
- X. Wu, J. W. Sun, P. F. Liu, J. Y. Zhao, Y. Liu, L. Guo, S. Dai, H. G. Yang and H. Zhao, *Adv. Funct. Mater.*, 2022, **32**, 2107301.
- H. Xu, H. Cai, L. Cui, L. Yu, R. Gao and C. Shi, *Nano Res.*, 2023, **16**, 3649–3657.
- Q. Wu, D. Si, J. Liang, Y. Huang and R. Cao, *Appl. Catal., B*, 2023, **333**, 122803.
- J. Li, B. Shang, Y. Gao, S. Cheon, C. L. Rooney and H. Wang, *Nat. Synth.*, 2023, **2**, 1194–1201.
- J. Su, C. B. Musgrave, Y. Song, L. Huang, Y. Liu, G. Li, Y. Xin, P. Xiong, M. M.-J. Li, H. Wu, M. Zhu, H. M. Chen, J. Zhang, H. Shen, B. Z. Tang, M. Robert, W. A. Goddard and R. Ye, *Nat. Catal.*, 2023, **6**, 818–828.
- J. Ding, Z. Wei, F. Li, J. Zhang, Q. Zhang, J. Zhou, W. Wang, Y. Liu, Z. Zhang, X. Su, R. Yang, W. Liu, C. Su, H. B. Yang, Y. Huang, Y. Zhai and B. Liu, *Nat. Commun.*, 2023, **14**, 6550.
- L. Yao, K. E. Rivera-Cruz, P. M. Zimmerman, N. Singh and C. C. L. McCrory, *ACS Catal.*, 2024, **14**, 366–372.
- Y. Liu, A. Deb, K. Y. Leung, W. Nie, W. S. Dean, J. E. Penner-Hahn and C. C. L. McCrory, *Dalton Trans.*, 2020, **49**, 16329–16339.
- W. Hu, D. Wang, Q. Ma, B. J. Reinhart, X. Zhang and J. Huang, *J. Photochem. Photobiol.*, 2022, **11**, 100132.
- N. Li, W. Lu, K. Pei and W. Chen, *RSC Adv.*, 2015, **5**, 9374–9380.
- D. Masheder and K. P. J. Williams, *J. Raman Spectrosc.*, 1987, **18**, 391–398.
- F. Evangelista, A. Ruocco, R. Gotter, A. Cossaro, L. Floreano, A. Morgante, F. Crispoldi, M. G. Betti and C. Mariani, *J. Chem. Phys.*, 2009, **131**, 174710.
- B. J. Palys, G. J. Puppels, D. van den Ham and D. Feil, *J. Electroanal. Chem.*, 1992, **326**, 105–112.
- B. J. Palys, D. M. W. van den Ham, W. Briels and D. Feil, *J. Raman Spectrosc.*, 1995, **26**, 63–76.
- Z. Weng, Y. Wu, M. Wang, J. Jiang, K. Yang, S. Huo, X.-F. Wang, Q. Ma, G. W. Brudvig, V. S. Batista, Y. Liang, Z. Feng and H. Wang, *Nat. Commun.*, 2018, **9**, 415.
- S. Ren, E. W. Lees, C. Hunt, A. Jewlal, Y. Kim, Z. Zhang, B. A. W. Mowbray, A. G. Fink, L. Melo, E. R. Grant and C. P. Berlinguette, *J. Am. Chem. Soc.*, 2023, **145**, 4414–4420.
- S. Jiang, Z. Chen, X. Chen, D. Nguyen, M. Mattei, G. Goubert and R. P. Van Duyne, *J. Phys. Chem. C*, 2019, **123**, 9852–9859.
- J. Zagal, M. Páez, A. A. Tanaka, J. R. dos Santos and C. A. Linkous, *J. Electroanal. Chem.*, 1992, **339**, 13–30.
- M. C. Biesinger, B. P. Payne, A. P. Grosvenor, L. W. M. Lau, A. R. Gerson and R. St. C. Smart, *Appl. Surf. Sci.*, 2011, **257**, 2717–2730.
- W. Luo and S. Zafeiratos, *J. Phys. Chem. C*, 2016, **120**, 14130–14139.



40 Q. H. Low, N. W. X. Loo, F. Calle-Vallejo and B. S. Yeo, *Angew. Chem., Int. Ed.*, 2019, **58**, 2256–2260.

41 R. Urrego-Ortiz, S. Builes, F. Illas and F. Calle-Vallejo, *EES Catal.*, 2024, **2**, 157–179.

42 L. P. Granda-Marulanda, A. Rendón-Calle, S. Builes, F. Illas, M. T. M. Koper and F. Calle-Vallejo, *ACS Catal.*, 2020, **10**, 6900–6907.

43 A. A. Peterson and J. K. Nørskov, *J. Phys. Chem. Lett.*, 2012, **3**, 251–258.

44 A. Bagger, W. Ju, A. S. Varela, P. Strasser and J. Rossmeisl, *ChemPhysChem*, 2017, **18**, 3266–3273.

45 A. Bagger, *Curr. Opin. Electrochem.*, 2023, **40**, 101339.

46 S. C. Jesudass, S. Surendran, G. Janani, T.-H. Kim and U. Sim, *Adv. Energy Mater.*, 2023, **13**, 2301918.

47 J.-J. Lv, M. Jouny, W. Luc, W. Zhu, J.-J. Zhu and F. Jiao, *Adv. Mater.*, 2018, **30**, 1803111.

48 S. Cheon, J. Li and H. Wang, *J. Am. Chem. Soc.*, 2024, **146**, 16348–16354.

49 J. Li, Z. Wang, C. McCallum, Y. Xu, F. Li, Y. Wang, C. M. Gabardo, C.-T. Dinh, T.-T. Zhuang, L. Wang, J. Y. Howe, Y. Ren, E. H. Sargent and D. Sinton, *Nat. Catal.*, 2019, **2**, 1124–1131.

50 J. Jin, J. Wicks, Q. Min, J. Li, Y. Hu, J. Ma, Y. Wang, Z. Jiang, Y. Xu, R. Lu, G. Si, P. Papangelakis, M. Shakouri, Q. Xiao, P. Ou, X. Wang, Z. Chen, W. Zhang, K. Yu, J. Song, X. Jiang, P. Qiu, Y. Lou, D. Wu, Y. Mao, A. Ozden, C. Wang, B. Y. Xia, X. Hu, V. P. Dravid, Y.-M. Yiu, T.-K. Sham, Z. Wang, D. Sinton, L. Mai, E. H. Sargent and Y. Pang, *Nature*, 2023, **617**, 724–729.

