

Cite this: *RSC Sustainability*, 2024, 2, 445

# Efficient integration of carbon dioxide reduction and 5-hydroxymethylfurfural oxidation at high current density†

Roger Lin, ‡ Haoyan Yang, ‡ Hanyu Zheng, ‡ Mahdi Salehi, Amirhossein Farzi, Poojan Patel, Xiao Wang, Jiaxun Guo, Kefang Liu, Zhengyuan Gao, Xiaojia Li and Ali Seifitokaldani \*

Commercialization of the electrochemical CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR) is a crucial step towards carbon utilization and addressing climate change. However, its low energy efficiency in full electrolytic systems poses significant challenges. To overcome this, we investigate an integrated electrolytic cell coupling the CO<sub>2</sub>RR with the 5-hydroxymethylfurfural oxidation reaction (HMFOR), a promising alternative to the oxygen evolution reaction (OER) at the anode. The utilization of nickel-phosphorus (NiP) electrocatalysts achieves high faradaic efficiency (90%) and stability (>200 redox cycles) for the anodic HMFOR to produce 2,5-furandicarboxylic acid (FDCA) at a high current density of 100 mA cm<sup>-2</sup>. Simultaneously, Sn, Ag and Cu nanoparticles act as efficient cathode electrocatalysts in a flow cell reactor, to produce formate, syngas, C<sub>2+</sub> hydrocarbon and other CO<sub>2</sub>RR products. Compared to the OER-CO<sub>2</sub>RR, the integrated HMFOR-CO<sub>2</sub>RR system demonstrates an energy efficiency increase of more than 10%. Additionally, economic analysis shows a 45% reduction in the levelized cost of potassium formate production in an optimistic scenario. This integrated CO<sub>2</sub>RR-HMFOR electrocatalytic system holds promise for commercializing CO<sub>2</sub> reactors.

Received 20th October 2023  
Accepted 12th December 2023

DOI: 10.1039/d3su00379e

rsc.li/rscsus

## Sustainability spotlight

In the face of increasing CO<sub>2</sub> levels and their associated environmental challenges, our study focuses on a novel approach to combat climate change. We have explored an enhanced CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR) system, leveraging clean electricity to establish a carbon capture and utilization (CCU) process. Our innovation involves replacing the anodic oxygen evolution reaction (OER) with an energy-efficient upgrade of biomass-derived 5-hydroxymethylfurfural (HMF) to 2,5-furandicarboxylic acid (FDCA). This work not only aligns with the UN's Sustainable Development Goals for waste recycling (SDG 12) but also contributes to the industrialization of efficient clean energy consumption (SDG 7 and 9), thus combating climate change (SDG 13).

## Introduction

The electrochemical CO<sub>2</sub> reduction reaction (CO<sub>2</sub>RR) is one of the promising options for carbon capture and utilization (CCU). It is a catalytic reaction using electricity as the energy source to convert CO<sub>2</sub> or bicarbonate to various products including carbon monoxide, alcohols, hydrocarbons, and organic acids (shown in Fig. 1). Among these products, some might have greater market potential, such as formic acid (HCOOH),<sup>1,2</sup> methanol (CH<sub>3</sub>OH),<sup>3-5</sup> ethanol (CH<sub>3</sub>CH<sub>2</sub>OH),<sup>6,7</sup> ethylene (C<sub>2</sub>H<sub>4</sub>),<sup>8,9</sup> and carbon monoxide (CO).<sup>10,11</sup> However, challenges remain today as the applied potential across the cell is often

large, while the energy efficiency is low. This means that lab-scale projects have a high energy consumption for a required output, leading to high costs and challenges to scale up to industrial applications. There is also a limited amount of economic analysis proof for a profitable process on the industrial scale, so a financial obstacle exists when implementing the CO<sub>2</sub>RR alone. According to Jouny,<sup>12</sup> CO and formic acid are profitable under current economic conditions, but many other CO<sub>2</sub>RR products may have a promising future as profitable options. Moreover, given the research by Kibria,<sup>13</sup> it is shown that the CO<sub>2</sub>RR needs to maintain a stable operation for 80 000 hours to be economically viable. Currently, there is a trade-off between high selectivity and high market value for a single CO<sub>2</sub>RR product. Formate and CO have high selectivity with low market prices, while ethanol and ethylene are the opposite. The current methods are not yet satisfactory to the industry, and the performance must be increased in parallel with further

Department of Chemical Engineering, McGill University, Montréal, QC H3A0C5, Canada. E-mail: ali.seifitokaldani@mcgill.ca; Tel: +1-514-398-4866

† Electronic supplementary information (ESI) available. See DOI: <https://doi.org/10.1039/d3su00379e>

‡ These authors contributed equally to this work.



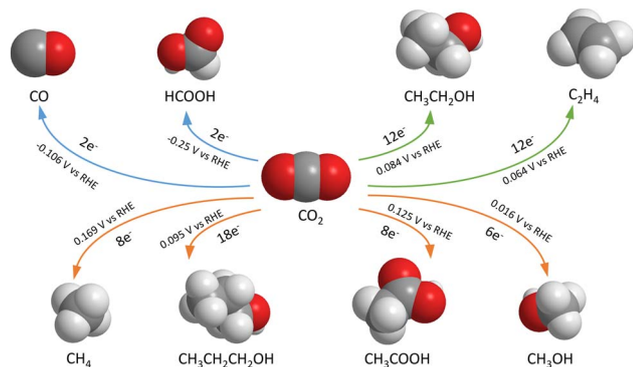


Fig. 1 Common electrochemical CO<sub>2</sub> reduction reaction products, number of electrons transferred and reaction standard potentials.<sup>14</sup>

developments in techno-economic analysis (TEA) and life-cycle assessment (LCA) studies.

A typical electrolyzer consists of the CO<sub>2</sub>RR in the cathode and the oxygen evolution reaction (OER) in the anode, which is the generation of oxygen through water electrolysis. Previous studies have developed various electrocatalysts for the OER, as shown in Table 1 below. However, the OER faces challenges such as high energy requirements due to its large overpotential and slow charge transfer. Hence, more studies are looking into replacing the OER with a reaction requiring less energy input, *i.e.*, lower applied potential, while achieving a high reaction rate.<sup>15</sup> The oxidations of methanol, ethanol, glycerol benzyl alcohol, hydroxymethyl-furfural, and HMF are some examples of the alternative anodic reactions.<sup>16–18</sup>

We propose to replace the OER with biomass upgrading. Specifically, we focus on the electrocatalytic oxidation reaction of 5-hydroxymethylfurfural (HMF), converting a biomass byproduct from the pulp and paper industry, into more valuable chemicals. Some previous studies break down and convert HMF into formic acid (FA) and maleic acid (MA),<sup>24</sup> while HMF can also be converted into 2,5-furandicarboxylic acid (FDCA).<sup>25,26</sup> FDCA serves as a crucial precursor, capable of generating polyethylene 2,5-furandicarboxylate (PEF) as a monomer, which in turn, can be utilized in the production of polyethylene terephthalate (PET), a promising contender for biomass-based plastics.<sup>27</sup> Au, Pt, Pd and various precious metals as well as molecular-based electrocatalysts have been studied to produce FDCA at low potential with outstanding performance.<sup>28–31</sup>

However, noble metals are scarce and expensive. Meanwhile, the research by Latsuzbaia showed that Ni has outstanding catalytic performance with high FDCA yield and faradaic efficiency (FE) of around 90% and 80%, respectively.<sup>32</sup> Ni also has a lower cost than noble metals, making it more feasible for industrial application. Kubota *et al.* demonstrated the possible acidic HMFOR electrolyzer in pH 1 sulfuric acid electrolyte, with a 53.8% yield of FDCA but an FE of only 33.8% due to the MA formation.<sup>33</sup> Recent studies on the HMFOR, with the hydrogen evolution reaction (HER) in the cathode, have tested various Ni-based catalysts including 3D Ni<sub>2</sub>P and Ni<sub>3</sub>S<sub>2</sub> on nickel foam,<sup>34–36</sup> Co<sub>0.4</sub>NiS@NF,<sup>37</sup> and NiCoMn-layered double hydroxides<sup>38</sup> to demonstrate advantages in voltage reduction, stability, and faradaic efficiencies. The HMFOR-CO<sub>2</sub>RR system was later studied by Choi *et al.* under near-neutral conditions, which achieved FE of 70% but only at a low current density of ~1 mA cm<sup>-2</sup>.<sup>39</sup> Recent work from Bi *et al.* has achieved a higher current density of around 100 mA cm<sup>-2</sup> by producing MA and FA from HMF in the anode and CO in the cathode with the aid of acetonitrile and [Bmim]BF<sub>4</sub>. They also showed a stable current density and CO faradaic efficiency after 10 hours of continuous electrolysis with no noticeable change in the structure and composition of the electrocatalyst. Besides, the CO<sub>2</sub>RR-HMFOR paired system reduced the overall cell voltage to 2.70 V while maintaining a considerably high yield.<sup>29</sup> In the current research, an alkaline aqueous system is more practical than the acidic one because FDCA is a weak acid and has low solubility in water at low pH, which leads to selectivity and separation issues.<sup>40</sup>

To efficiently tackle the carbon dioxide challenge, we present the combination of the CO<sub>2</sub>RR and HMFOR in an alkaline system *via* the NiP electrocatalyst on carbon felt (CF) to achieve a high current density of 100 mA cm<sup>-2</sup> and 90% FE on both the cathode and anode, with a low cell voltage of ~2 V. This work also investigates the feasibility of a combined system from experimental and techno-economic perspectives. Using this setup, CO<sub>2</sub> and HMF are transformed into products with considerable market values and demands, such as CO (0.20 USD per kg),<sup>41</sup> formate (0.90 USD per kg),<sup>42</sup> ethylene (0.49–0.94 USD per kg),<sup>43</sup> ethanol (0.77 USD per kg),<sup>43</sup> FDCA (1.53 USD per kg)<sup>44</sup> and their respective net present values (NPVs) are all expected to increase in the future. Implementation of such a system not only offers vast opportunities for scientific exploration but also presents a green solution for carbon capture and reutilization. By simultaneously advancing carbon-neutral objectives, renewable fuel

Table 1 Brief summary of OER electrocatalysts and reaction parameters compared with the HMFOR

Electrode	Reaction	Electrolyte	Voltage (V vs. RHE) at 10 mA cm <sup>-1</sup>	Tafel slope [mV dec <sup>-1</sup> ]	Ref.
α-Co(OH) <sub>2</sub> /HNFs	OER	1.0 M KOH	1.54	68.9	19
Co-Mo-N hollow polyhedra	OER	1.0 M KOH	1.53	57	20
N-doped hollow carbon spheres	OER	0.1 M KOH	1.59	182	21
Co <sub>0.5</sub> Fe <sub>0.5</sub> O <sub>0.5</sub> F <sub>1.5</sub>	OER	1.0 M KOH	1.45	27	22
Ni <sub>3</sub> ZnCo <sub>0.7</sub> @CoNiCuFe-NC	OER	1.0 M KOH	1.43	80.4	23
Commercial IrO <sub>2</sub>	OER	1.0 M KOH	1.51	105.1	23
NiP@Ni/Carbon felt (CF)	OER	1.0 M KOH	1.53	111.7	This work
NiP@Ni/CF	HMFOR	1.0 M KOH (15 mM HMF)	1.40	35.8	This work



production, and circular economy principles, this integrated approach promises significant environmental benefits.

## Methodology

### Materials and chemicals

A flow cell was used for a continuous CO<sub>2</sub>RR experiment combined with the HMFOR. It contained three chambers: cathode gas chamber, catholyte chamber and anolyte chamber (Fig. 2). A gas diffusion layer (GDL) with an electrocatalyst coating was installed between the cathode gas chamber and the catholyte chamber (see Fig. S1, ESI<sup>†</sup>). The anode electrocatalyst NiP coated on Ni-deposited carbon felt (NiP@Ni/CF) was placed in the anode chamber. An anion exchange membrane was in place to separate the two half-cells, allowing anions to pass through while preventing product crossover. 1 M KOH solution as the electrolyte was filled in both cathode and anode chambers, where HMF was present in the anode during the HMFOR tests.

### Electrocatalyst synthesis

The anode electrocatalyst NiP@Ni/CF was prepared by electro-deposition of Ni on 2 nm-Ni-deposited carbon felt (Ni/CF) in an undivided cell (Fig. 3 and S2, ESI<sup>†</sup>). Ag, Cu, and Sn electrocatalysts were used on the cathode side. Ag/CP, Cu/CP, Ag/PTFE, and Cu/PTFE were prepared by depositing metal nanoparticles on carbon paper (CP) or PTFE (Polytetrafluoroethylene) *via* the physical vapor deposition (PVD) technique; while Sn/CP was prepared by drop-casting Sn nanoparticles onto CP (see Fig. S3, ESI<sup>†</sup>).

### Characterization

Physicochemical characterization techniques were employed to investigate the morphology and chemical composition of the electrocatalysts, as well as to confirm the presence of their active sites. These methods encompassed scanning electron microscopy (SEM), energy-dispersive X-ray spectroscopy (EDX), X-ray

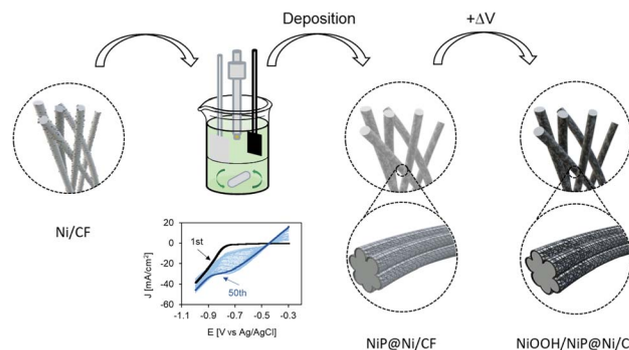


Fig. 3 Synthesis of the NiP electrocatalyst starts with Ni-deposited carbon felt, which is coated with the NiP layer by electrodeposition. Applying an oxidative potential on the electrocatalyst yields NiOOH on the coated surface layer.

diffraction (XRD), X-ray photoelectron spectroscopy (XPS) and X-ray absorption spectroscopy (XAS).

### Electrochemical measurements

Cyclic voltammetry (CV) and linear sweep voltammetry (LSV) were used to investigate the potential window of the desired reaction and activity of the electrocatalyst. In the HMFOR, the potential window is between the surface oxidation of Ni<sup>2+</sup> and the OER-dominating region, while in the CO<sub>2</sub>RR, it is between the onset potential and the hydrogen evolution reaction (HER)-dominating region. CV was also used to study the recyclability of the NiP electrocatalyst.

Chronoamperometry and chronopotentiometry experiments were conducted to assess the electrochemical performance of the electrocatalysts under reaction conditions. Electrochemical impedance spectroscopy (EIS) tests were performed using a frequency response analyzer (FRA) to determine the system resistance ( $R_s$ ) and charge transfer resistance ( $R_{ct}$ ) of the reactor setup with the electrocatalyst.

To convert the reference electrode reading to the reversible hydrogen electrode (RHE) potential under atmospheric non-standard pH conditions, eqn (1) was used:

$$E(\text{RHE}) = E_{\text{Ag}/\text{AgCl}} + 0.210 \text{ V} + 0.059 \times \text{pH} \quad (1)$$

### Product analysis

The gas and liquid products of the CO<sub>2</sub>RR and HMFOR from the chronoamperometry and chronopotentiometry experiments were analyzed using gas chromatography (GC) and high-performance liquid chromatography (HPLC), respectively.

A flow test was done before gas product sampling to determine the gas flow rate, which was used to calculate the product rate and eventually product FE.

The calculations of product yield and faradaic efficiency are given by the following eqn (2) and (3) respectively:

$$\text{yield}_{\text{product}} = \frac{\Delta N_{\text{Product}}}{N_{\text{Reactant, in}}} \times \frac{V_{\text{Reactant}}}{V_{\text{Product}}} \times 100\% \quad (2)$$

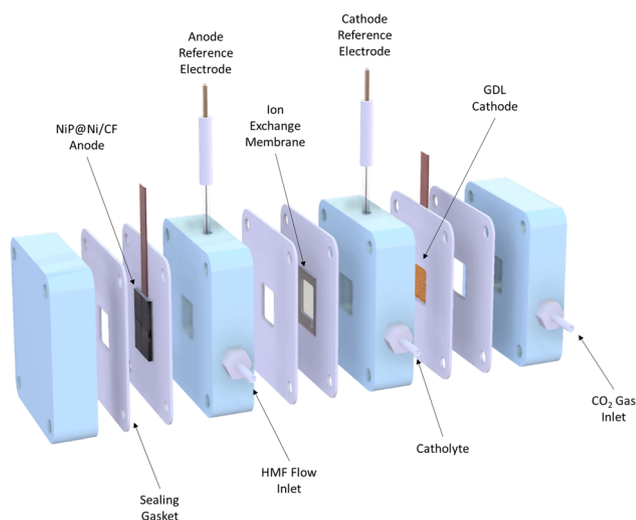


Fig. 2 Cross-sectional view of the flow cell.



$$FE_{\text{Product}} = \frac{n \times F \times N_{\text{Product}}}{Q} \times 100\% \quad (3)$$

where  $N$  is the number of moles,  $\nu$  is the stoichiometric coefficient in the balanced reaction,  $F$  is the Faraday number (96 485 C mol<sup>-1</sup>),  $n$  is the number of moles of electrons transferred per mole of product, and  $Q$  is the charge.

Applied voltages were  $iR$ -corrected with the system resistance to exclude the influence of the electrical setup.

From the standard potentials and the applied voltage, voltage efficiency (VE) is calculated:

$$VE = \frac{E_{\text{OER}}^0 - E_{\text{CO}_2\text{RR}}^0}{V_{\text{cell}}} \times 100\% \quad (4)$$

From the FE and VE, the energy efficiency (EE) of the electrolyzer is calculated for each product:

$$EE_i = VE \times FE_i \times 100\% \quad (5)$$

In Cu-based electrocatalysts where multiple products were produced, the total EE is a sum of the individual EE of detected products.

When the CO<sub>2</sub>RR is combined with the HMFOR, an additional aspect to assess energy saving is achieved by dividing the conventional CO<sub>2</sub>RR-OER voltage by the improved cell voltage when HMF is introduced at the identical current. This yields the voltage saving factor  $\eta_v$ :

$$\eta_v = \frac{\Delta E_{\text{CO}_2\text{RR-OER}}}{V_{\text{cell}}} \quad (6)$$

This parameter effectively indicates the voltage-saving capability of the electrochemical cell, where a number higher than unity means better energy saving. A theoretical maximum value of the saving on the conventional OER occurs when the cathode and the anode have zero overpotential. Then, based on the voltage saving, the faradaic efficiencies of both cathode and anode reactions are multiplied to yield the cell energy saving factor  $\eta_{\text{cell}}$ :

$$\eta_{\text{cell}} = \eta_v \times FE_{\text{cathode}} \times FE_{\text{anode}} \quad (7)$$

This parameter indicates the cell energy saving compared to the conventional CO<sub>2</sub>RR-OER system. A near-unity result indicates no significant energy saving for the full cell, while a larger number refers to a considerable saving from the energetic perspective. If the FE of one half-cell is too low,  $\eta_{\text{cell}}$  will be a relatively small number, indicating a low benefit for simultaneous production.

## Results and discussion

### Electrocatalyst characterization

SEM images in Fig. 4 show a uniform coating layer consisting of nickel- and phosphorus-rich deposits on the fiber structure of the carbon felt. The surface of the NiP layer has a typical nanosheet structure of Ni(OH)<sub>2</sub>, indicating the successful

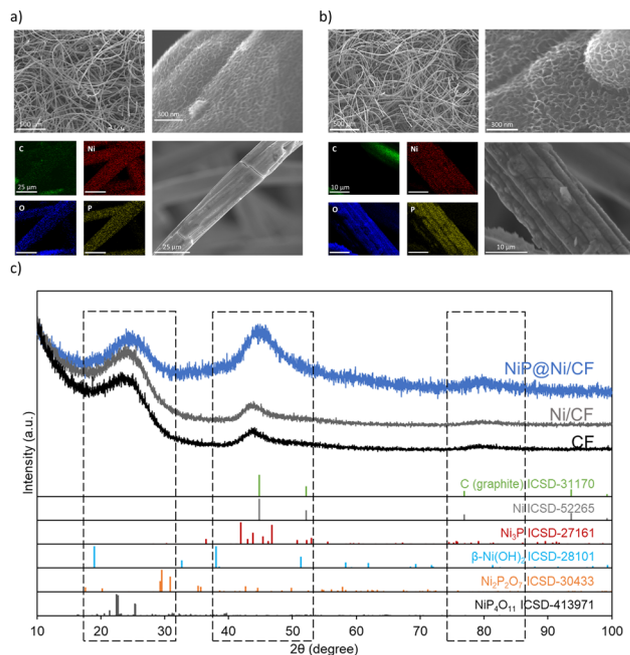


Fig. 4 Scanning electron microscopy and energy-dispersive X-ray spectroscopy of the NiP@Ni/CF anode electrocatalyst (a) as-prepared and (b) post-HMFOR; (c) X-ray diffraction patterns of NiP@Ni/CF, Ni/CF and CF.

deposition of Ni,<sup>45,46</sup> EDX results indicate the co-deposition of Ni and P (Fig. 4a, b and S4, ESI†). The carbon fibers are uniformly covered by nickel (46%), oxygen (15%) and phosphorus (10%). Similarly, EDX on NiP@Ni/CF after being used for the HMFOR (>10 cycles) shows the presence of Ni, O and P.

The XRD results in Fig. 4c demonstrate that the peaks between 20°–100° of 2\* $\theta$  are mainly from the carbon felt substrate. When we compare the Ni/CF with the CF substrate, we observe no difference, as the 2 nm of deposited Ni remains in nanoparticles, without any large crystals on the substrate. For the NiP@Ni/CF sample, signal broadening in the ranges of 20°–30°, 40°–50°, and 75°–85° indicates successful deposition of mostly amorphous or microcrystalline nickel–phosphorus active materials on the carbon felt, potentially with a broad crystal size distribution.<sup>47</sup> This is consistent with the results from SEM and EDX showing the typical nanosheet Ni(OH)<sub>2</sub> on top of the nickel and phosphorus co-deposit layer.

XPS survey reveals the oxidation state of Ni sites on the NiP@Ni/CF electrocatalyst and confirms the presence of Ni, O, C, and P on the electrocatalyst surface (Fig. S5, ESI†). High

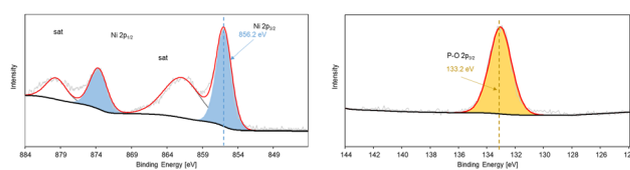


Fig. 5 High resolution XPS of (a) Ni 2p spectra and (b) P 2p spectra on as-prepared NiP@Ni/CF.



resolution XPS of Ni 2p spectra in Fig. 5 shows that Ni<sup>2+</sup> (856.2 eV) is the dominant Ni species on the surface, while P2p spectra reveal that phosphorus-oxygen bonds are also present on the as-prepared electrocatalyst surface.<sup>46,48</sup> Higher Ni oxidation state, namely Ni<sup>3+</sup> (857.7 eV), is found after the HMFOR, revealing the oxidation of Ni surface species post-HMFOR.<sup>40</sup> This surface oxidation of Ni<sup>2+</sup> is typical to the Ni(OH)<sub>2</sub>/NiOOH material as shown in Fig. S6 (ESI),<sup>†</sup> prepared by pulsed oxidation of Ni foam from our previously published work.<sup>49</sup> The redox couple of Ni<sup>2+</sup> and Ni<sup>3+</sup>, specifically in the form of NiOOH, is well known to be responsible for the HMFOR and other organic oxidation reactions targeting hydroxyl and aldehyde groups.<sup>50–53</sup> A dominant Ni–P bonding is not clearly seen likely due to the broad range of coordination possible among Ni, P and O, but the Ni 2p and P 2p spectra indicate a composition of nickel hydroxides and phosphate species, given the oxidized Ni peak of a typical Ni–O bond and the partially oxidized P peak (133.2 eV) indicating the presence of the P–O bond,<sup>54,55</sup> potentially resulting from surface oxidation of Ni–P species.<sup>48,56</sup> However, we are also aware that the presence of the P–O bond can enable further Ni oxidation under anodic conditions (Fig. S5, ESI<sup>†</sup>) and therefore generate an active Ni<sup>2+</sup>/Ni<sup>3+</sup> redox pair for the HMFOR.<sup>57</sup> The XPS results also indicate that the conversion of Ni<sup>2+</sup> to Ni<sup>3+</sup> takes place on the surface of the electrocatalyst after applying an anodic potential during the HMFOR.

XAS results indicate the existence of Ni and provide insights into its oxidation state on the electrocatalyst. Scanning is carried out under both *ex situ* conditions and *in situ* oxidative conditions at 0.44 V vs. Ag/AgCl (Fig. S7, ESI<sup>†</sup>). As observed in Fig. 6, as-prepared NiP and NiP deposited on carbon felt (NiP@Ni/CF) have a blue shift to the right side compared to the Ni foil reference. Such a shift towards Ni(II) shows that in the NiP electrocatalyst a higher oxidation state can be attributed to Ni.

From the surface-sensitive total electron yield (TEY) method on the catalytic NiOOH deposited on Ni foam, tests yield a blue shift from the Ni foil and Ni(II) oxide, showing similar edge and peak features to the oxide.<sup>58</sup> This suggests that the surface layer

of the NiOOH electrocatalysts exhibits an oxidation state higher than +2. To quantitatively investigate the oxidation states of the NiP, we conducted linear combination fitting on our experimental data. Based on Table S1 (ESI),<sup>†</sup> despite the presence of minor deviations in fitted values, it is feasible to effectively combine Ni foil and Ni(II) to fit the NiP/CF electrocatalyst. For NiP, the disparities between the *in situ* and *ex situ* oxidation states are not statistically significant. Therefore, under both *ex situ* and oxidative conditions, it can be claimed that the NiP@Ni/CF sample has an average oxidation state of about +1.2. It is in agreement with a blue shift of the edge in NiP@Ni/CF towards the Ni(II) oxide standard (Fig. 6).

SEM images of the cathode electrocatalysts Ag/CP, Cu/CP, and Sn/CP show uniform deposition of metal particles on carbon paper, (Fig. S8, ESI<sup>†</sup>). For Ag and Cu electrocatalysts on the PTFE surface, the deposited metals are shown to be uniformly coating the polymer fibers (Fig. S9, ESI<sup>†</sup>). EDX analysis of the electrocatalysts also indicates the uniform distribution of catalytic materials on the carbon paper (Fig. S10, ESI<sup>†</sup>). To verify the oxidation state of the as-prepared electrocatalysts, high resolution XPS is also used. Ag on carbon paper is shown to be in the Ag (0) state (Fig. S11, ESI<sup>†</sup>), while Cu and Sn consist of mixed metallic states and naturally occurring higher oxidation states (Cu<sup>+</sup>/Cu<sup>2+</sup> and Sn<sup>2+</sup>) (Fig. S12 and S13, ESI<sup>†</sup>).

## Electrochemical performance

**Electrocatalytic activity: LSV-CV, EIS, and tafel slope.** The electrochemical performance test of NiP@Ni/CF is first carried out *via* cyclic and linear sweep voltammetry. CV shows the active oxidation region for the HMFOR starts from 1.35 to 1.36 vs. RHE (Fig. 7a), which aligns with the oxidation of Ni<sup>2+</sup> to Ni<sup>3+</sup> (eqn (8)) in alkaline electrolyte, indicating that Ni<sup>3+</sup> is the active site. According to previous studies, two pathways exist for HMF molecule oxidation on Ni-based electrocatalysts.<sup>49,52,59</sup> Eqn (9) shows the spontaneous indirect oxidation of HMF by Ni<sup>3+</sup> species on the electrocatalyst surface, while eqn (10)–(12) are the direct oxidations of HMF on Ni<sup>3+</sup> at higher applied potentials. The reduction region in 0.9–1.3 V corresponds to the reduction of surface oxidized Ni species.<sup>60</sup> The smaller reduction peak in CV indicates that less Ni<sup>3+</sup> is available for reduction (Fig. S14, ESI<sup>†</sup>).

LSV of the anode electrocatalyst in 1 M KOH without HMF is performed twice consecutively to obtain the OER background. The exposed Ni species is completely oxidized during the first scan, and the second scan is only attributed to the OER (dash line in Fig. 7b). The onset potential of the NiP electrocatalyst for Ni oxidation and the HMFOR is ~200 mV smaller than that of the OER from the LSV tests. This indicates that the HMFOR is favoured over the OER under low applied anodic potentials, allowing higher selectivity for the HMFOR. A recyclability test using cyclic voltammetry shows stable reversibility of Ni active sites for 200 redox cycles in 1 M KOH (Fig. 7c). More importantly, the Ni active sites are shown to be stable with unchanged HMFOR activity after 200 redox cycles. In Fig. 7d, electrochemical impedance spectroscopy shows a smaller charge transfer resistance ( $R_{ct}$ ) with HMF in the electrolyte at an

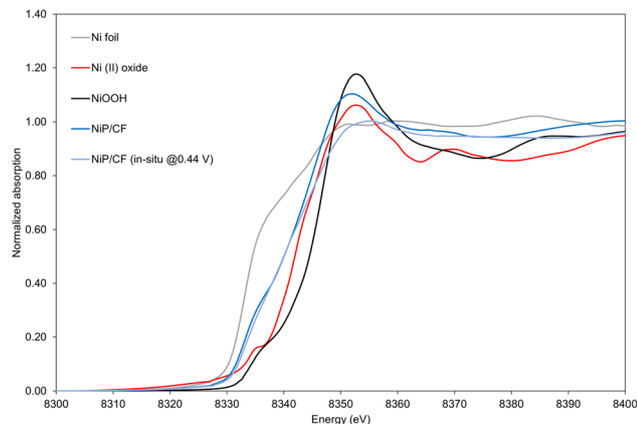


Fig. 6 X-ray absorption near edge structure (XANES) result of references (Ni foil, Ni(II) oxide and NiOOH) and samples under *ex situ* and *in situ* conditions at 0.44 V vs. Ag/AgCl (for NiP electrocatalysts).



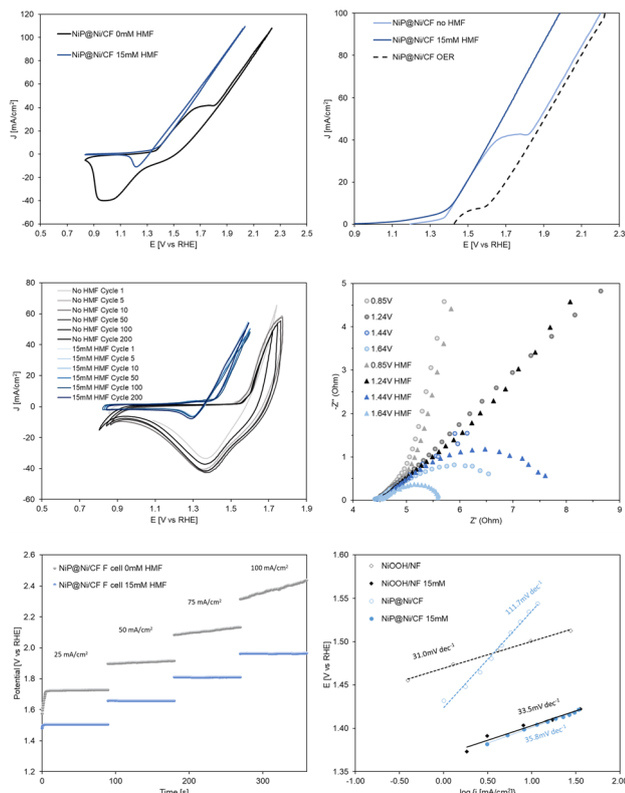
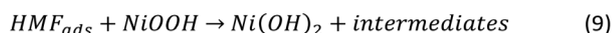
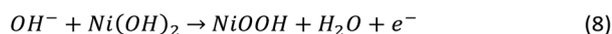


Fig. 7 (a) Cyclic voltammetry ( $10 \text{ mV s}^{-1}$ ) of NiP@Ni/CF with and without HMF; (b) linear sweep voltammetry ( $5 \text{ mV s}^{-1}$ ) of NiP@Ni/CF with and without HMF; (c) recyclability test by cyclic voltammetry ( $50 \text{ mV s}^{-1}$ ) over 200 redox cycles on the NiP@Ni/CF electrocatalyst in 1 M KOH with and without HMF; (d) electrochemical impedance spectroscopy (EIS) for NiP@Ni/CF in 1 M KOH with and without HMF (see Fig. S15–S18†); (e) chronopotentiometry of NiP@Ni/CF in 1 M KOH with and without HMF in a flow cell. The gradual increase of applied potential without HMF is due to the production of oxygen gas from the oxygen evolution reaction, which reduced the active surface area of the anode. (f) Tafel slopes of the oxidized NiP electrocatalyst for the HMFOR and OER, compared to NiOOH/Ni foam (potentials are  $iR$ -corrected).



Scheme 1 HMF oxidation reaction steps on the Ni-based electrocatalyst under alkaline conditions.

applied potential beyond 1.44 V (non- $iR$  corrected). This again proves that after the onset of the HMFOR, biomass oxidation is favoured over the OER, even after the onset of the OER at an applied potential of 1.64 V.

Stepwise chronopotentiometry is then performed to verify the potential required at each applied current density near the

onset of Ni oxidation (Fig. 7e). The results corroborate the previous observation that the NiP electrocatalyst can achieve equivalent current density at lower potential for the HMFOR in contrast to the OER. This illustrates that the higher activity of the HMFOR remains consistent even at higher applied current densities ( $100 \text{ mA cm}^{-2}$ ). Tafel slopes are obtained from stepwise chronoamperometry to confirm the higher activity for the HMFOR ( $35.8 \text{ mV dec}^{-1}$ ) compared to the OER ( $111.7 \text{ mV dec}^{-1}$ ) by the NiP electrocatalyst (Fig. 7f). It also shows that the oxidized NiP electrocatalyst has much smaller Tafel slope for the HMFOR compared to the OER, while a typical NiOOH on Ni foam electrocatalyst has indistinguishable HMFOR and OER Tafel slopes. This emphasizes that the intrinsically higher activity for the HMFOR demonstrated by NiP results in less applied potential to reach the same specific reaction activity.

LSV for the cathodic  $\text{CO}_2\text{RR}$  on Sn, Cu and Ag electrocatalysts on carbon paper as well as Cu and Ag on the PTFE substrate is shown in Fig. 8. The  $\text{CO}_2\text{RR}$  occurs on these electrocatalysts near  $-0.3 \text{ V vs. RHE}$ , while in the absence of  $\text{CO}_2$  (with only  $\text{N}_2$  purging in the gas chamber), the hydrogen evolution reaction takes place on the cathode near  $-0.5 \text{ V}$ . The difference in onset potential is the largest for Sn/CP, but all three electrocatalysts demonstrate the activity for the  $\text{CO}_2\text{RR}$  over the HER.

**Selectivity: cathode and anode FE.** Combined chronopotentiometry tests are then carried out with PVD Ag/CP, Cu/CP, drop-casted Sn/CP and PVD Ag/PTFE, Cu/PTFE electrocatalysts (Fig. S3, ESI†). Based on initial tests and information from existing studies, the efficiency of the  $\text{CO}_2\text{RR}$  in terms of FE remains largely consistent at a given current density, while the anode applied potential drops when HMF is present, attributed to the substitution of the OER with the HMFOR. The FE of  $\text{CO}_2\text{RR}$  and HMFOR is calculated for both  $25 \text{ mA cm}^{-2}$  (low) and  $100 \text{ mA cm}^{-2}$  (high), as shown in Fig. 9a and b (see Fig. S19–S24 for HPLC and GC, ESI†). With Sn/CP as the cathode electrocatalyst, for both low and high current densities, formate (62–80%) is the major product in the liquid phase and CO (8–17%) is the minor product in the gas phase. Ag/CP achieves  $\sim 95\%$  of CO production as a robust electrocatalyst. Cu/CP produces products

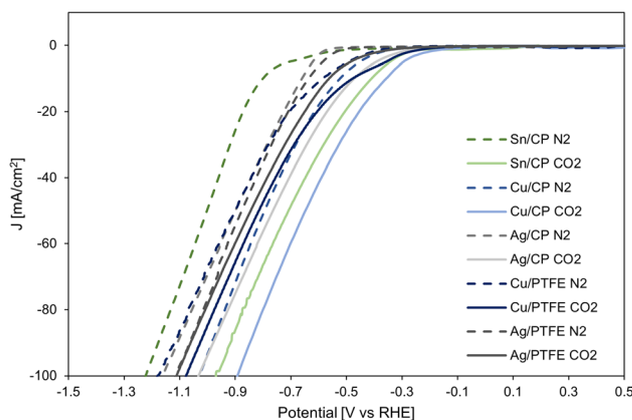


Fig. 8 Linear sweep voltammetry of cathode electrocatalysts Sn, Cu and Ag on carbon paper, Cu and Ag on PTFE under  $\text{N}_2$  (HER) and  $\text{CO}_2$  ( $\text{CO}_2\text{RR}$ ).



with different FE depending on the current density. At 25 mA cm<sup>-2</sup>, formate (~18%), acetate (~7%) and ethanol (6–16%) are in the liquid phase, and ethylene (~5%) and CO (32–41%) are in the gas phase. At 100 mA cm<sup>-2</sup>, the differences include lowered FE for formate (~8%), acetate (~1%), ethanol (5–10%), CO (22–35%), and a higher FE for ethylene (~31%). Taking 1-propanol (3%) into account, it results in 23–33% FE toward C<sub>2+</sub> products at the lower current, and 37–48% FE towards C<sub>2+</sub> at the higher current. For PTFE substrate samples, the higher current is used to compare with the carbon paper substrate results. Ag/PTFE shows similar FE to the Ag/CP, e.g., 80% FE for CO. Cu/PTFE interestingly shows a significant improvement in C<sub>2+</sub> product FE, especially ethylene (54–60%), while suppressing formate (~4%) and CO (~7%). This leads to the total FE toward C<sub>2+</sub> product to be in a high range of 75–80%. On the other half-cell, the HMFOR in the anode is consistent with ~60% FE toward FDCA, ~15% FE toward HMFA, ~12% toward FFCA, and 3–5% FE of DFF at the lower current density. At higher current density, i.e., 100 mA cm<sup>-2</sup>, 73–78% FE is toward FDCA in the flow-cell, regardless of the cathode electrocatalyst and reaction, as shown in Fig. 9b. Product analysis shows that the FE toward FDCA, HMFA, FFCA and DFF sums to 93–96%, demonstrating a high conversion of HMF to value-added products.

As shown in Fig. 9a and b, in the NiP//Sn/CP combined system with *iR* correction, the cathodic applied potential is around -0.43 V and -0.61 V at a current density of 25 and 100 mA cm<sup>-2</sup>, respectively; at high current density, the anodic applied potential decreases to ~240 mV when 15 mM of HMF is added to the anolyte. Similarly, the NiP//Cu/CP system shows a cathodic applied potential around -0.43 V at low current and -0.57 V at high current, while the anodic counterpart demonstrates a drop of ~240 mV when the HMFOR replaces the OER at 100 mA cm<sup>-2</sup>. For NiP//Ag/CP, the cathodic applied potential is -0.50 V and -0.68 V at low and high current density, while anodic potentials drop from 1.67 V to 1.45 V. In combined systems with PTFE as the cathode substrate, both systems at 100 mA cm<sup>-2</sup> show a similar drop in anode potential, while NiP//Cu/PTFE requires -0.58 V at the cathode and NiP//Ag/PTFE requires -0.76 V. A repeated test of 8 cycles is carried out on the NiP electrocatalyst in 15 mM HMF, showing a stable anode performance at 1.42 V RHE and >90% total FE for the HMFOR (Fig. 9c). See Table S2 (ESI)<sup>†</sup> for product summary data and Table S3 (ESI)<sup>†</sup> for performance comparison.

**Energy efficiency: improvement with the HMFOR.** In this study, due to the reduction in the anodic applied potential upon introducing HMF to the anode, the overall energy efficiency of

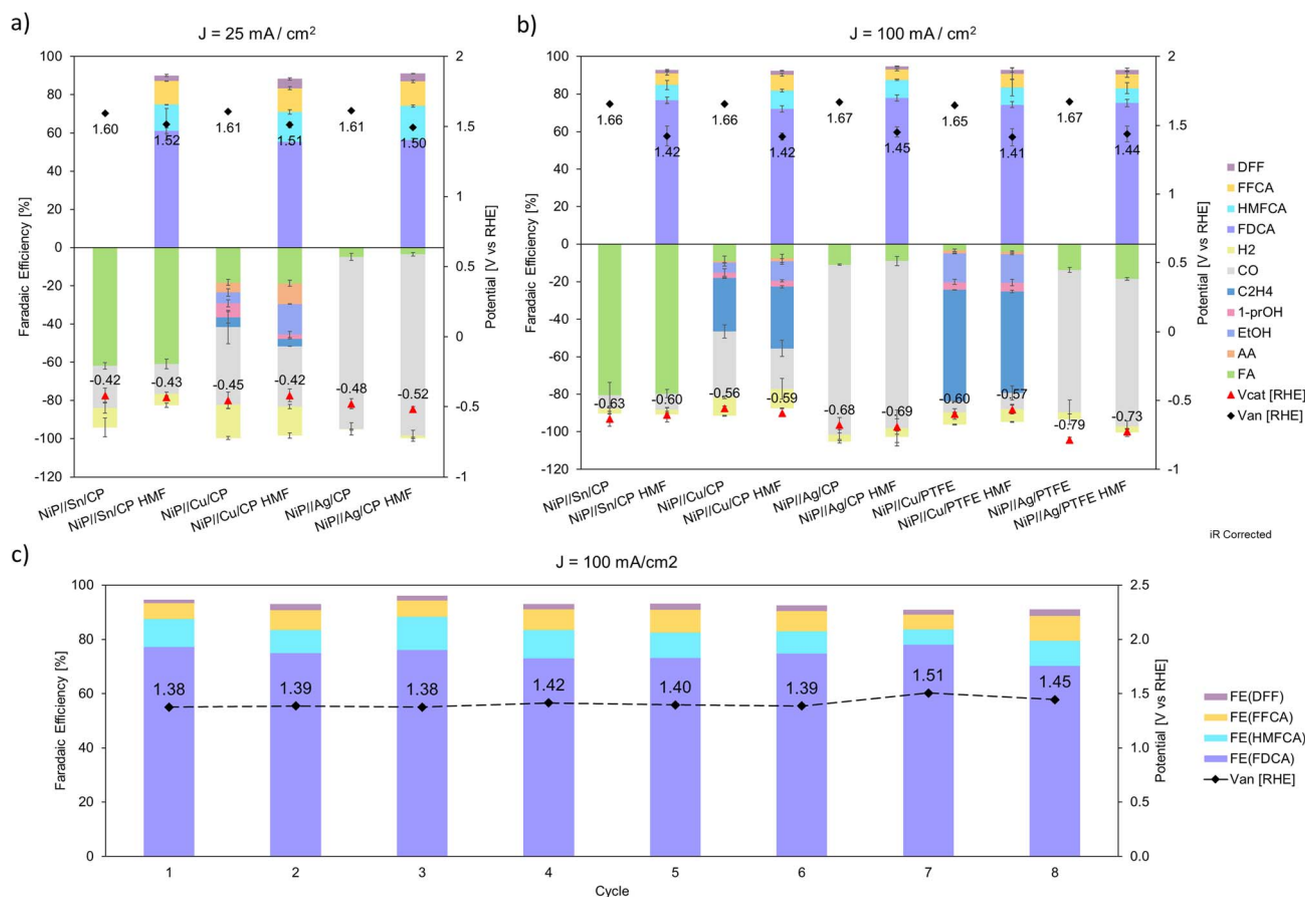


Fig. 9 Faradaic efficiencies of (a) cathode CO<sub>2</sub>RR products and anode HMFOR products at 25 mA cm<sup>-2</sup>; (b) cathode CO<sub>2</sub>RR products and anode HMFOR products at 100 mA cm<sup>-2</sup>; (c) repeated tests on NiP@Ni/CF with 15 mM HMF. The OER-paired system results are shown with cathode FE only, while the HMFOR-paired system results are shown with products from each half-cell.



the combined system experiences a direct improvement of approximately 5 to 6% compared to the OER-paired system. This translates to a relative improvement of about 11 to 12% when compared to using the OER as the anodic reaction. This is true among the paired cases of NiP//Sn/CP, NiP//Cu/CP, NiP//Ag/CP, NiP//Ag/PTFE, and NiP//Cu/PTFE combined systems (Table 2). Using the NiP//Cu/CP and the NiP//Cu/PTFE systems as a case study of the EE improvement, we can see the CO<sub>2</sub>RR energy efficiency increases from 44.6% to 49.8% for the Cu/CP cathode (Fig. 10a). However the PTFE substrate provides an extra hydrophobic GDL, enhancing the production of C<sub>2+</sub> products by C–C coupling.<sup>61</sup> Therefore, when coupled with the HMFOR, the energy efficiency for CO<sub>2</sub>RR products increases from 47.0% to 52.4% (Fig. 10b). Both cases with the copper cathode yield a relative improvement of 11.5% in energy efficiency. This can become significant when a selected product, such as ethylene, is the production target, where the related operational costs can be directly reduced, including energy consumption, reagent consumption and maintenance costs. Copper as the only element that is known to produce multiple C<sub>2+</sub> products from the CO<sub>2</sub>RR, is a step closer to commercialization as a result of the energy efficiency rise.<sup>13</sup>

Experimental results from the combined system have shown that the objective of high selectivity at high current density with improved system energy efficiency is achieved by replacing the conventional OER with the HMFOR as the anode reaction. This proposed technique in combination with literature results generally gives a predicted improvement in energy efficiency of 10 to 30% (Table S4, ESI†).

Following the experimental results, theoretical calculations are carried out to emphasize the significance of system optimization for future work. The base case is the currently observed CO<sub>2</sub>RR-OER system. Table S5 (ESI)† illustrates that if the cathode operates ideally (with a cathodic potential approaching the standard potential) and the faradaic efficiency for the CO<sub>2</sub>RR products reaches 100%, the incorporation of the observed HMFOR catalyzed by NiP into the system would result

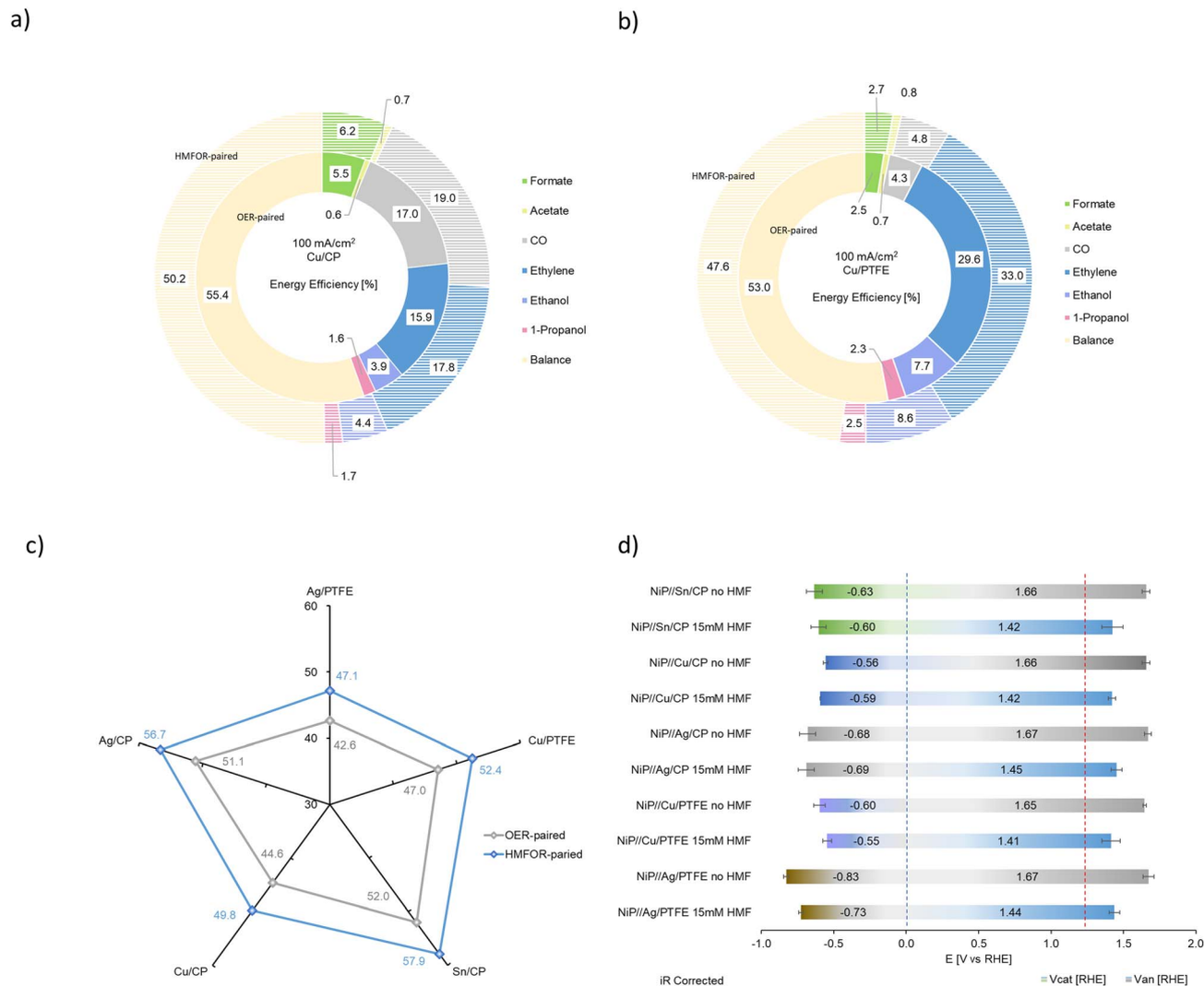
in an anticipated efficiency of around the mid to high 80% range. This would signify a notable increase of approximately 14 to 18% in comparison to when the OER is the anodic reaction, which means that the improvement by pairing with the HMFOR would be relatively better with an ideal cathode than a non-ideal cathode with respect to each product. Moreover, if the anode reaction is improved for the HMFOR (at 1.36 V vs. RHE where surface Ni<sup>2+</sup> is oxidized to Ni<sup>3+</sup>) with the current cathode performance, as shown in Table S6 (ESI),† the relative improvement on the system with the observed cathode performance would be 14 to 15%. This indicates the universal significance of anode reaction design. It can also be understood that the FE and applied potential on the cathode side greatly affect the cathode and the cell energy efficiencies, whereas the incorporation of the HMFOR would yield a relatively similar increase in cell EE on top of it. Hence, the initial focus for enhancement is on elevating the cathode FE, given its direct impact on energy efficiency. Subsequently, identifying a suitable anode reaction that substantially reduces the additional energy consumption at the anode stands out as another key step.

To illustrate the energy saving, more stringent parameters are introduced. Similar to the VE, dividing the cell voltage of the CO<sub>2</sub>RR-OER at 100 mA cm<sup>-2</sup> by the improved cell voltage demonstrates a voltage saving factor ( $\eta_v$ ) of the new setup. This is then multiplied by the FE of each half cell to obtain the cell energy saving factor,  $\eta_{cell}$ . The numerical value of  $\eta_v$ ,  $\eta_{cell}$  and the relative improvement are summarized in Table S7 (ESI).† These results show a general improvement in  $\eta_v$ , while the formate production by Sn/CP and the CO production by Ag/CP have the highest  $\eta_{cell}$ , given their robust cathodic FE. Cu/CP in contrast suffers from low FE for multiple products. When combined with the NiP, even with >90% HMFOR FE, the  $\eta_{cell}$  results are still below 1 for all the tests. In addition to optimizing the cathode FE, the combined test results indicate that further advancement at the anodic half-cell is needed. It should be noticed that the conventional OER has by default 100% FE,

**Table 2** System energy efficiencies and savings improved by a combination of the HMFOR with observed cathode FE and potential at 100 mA cm<sup>-2</sup>. Variables followed by the prime (') symbol indicate the proposed combined system with the NiP electrocatalyst for the HMFOR, same for all following tables. Equations for the calculation of FE and EE are shown in Appendices. Potentials are *iR* corrected

Product	Catalyst	V <sub>cat</sub> RHE	V <sub>an'</sub> RHE	EE (%)	EE' (%)	Improv. %
Formate	Sn/CP	-0.62	1.43	52.0	57.9	11.28
Formate	Cu/CP	-0.58	1.43	5.5	6.2	11.52
Acetate	Cu/CP	-0.58	1.43	0.6	0.7	11.52
CO	Cu/CP	-0.58	1.43	17.0	19.0	11.52
Ethylene	Cu/CP	-0.58	1.43	15.9	17.8	11.52
Ethanol	Cu/CP	-0.58	1.43	3.9	4.4	11.52
1-Propanol	Cu/CP	-0.58	1.43	1.6	1.7	11.52
CO	Ag/CP	-0.69	1.43	51.1	56.7	10.92
Formate	Cu/PTFE	-0.58	1.43	2.5	2.7	11.48
Acetate	Cu/PTFE	-0.58	1.43	0.7	0.8	11.48
CO	Cu/PTFE	-0.58	1.43	4.3	4.8	11.48
Ethylene	Cu/PTFE	-0.58	1.43	29.6	33.0	11.48
Ethanol	Cu/PTFE	-0.58	1.43	7.7	8.6	11.48
1-Propanol	Cu/PTFE	-0.58	1.43	2.3	2.5	11.48
CO	Ag/PTFE	-0.76	1.43	42.6	47.1	10.56





**Fig. 10** CO<sub>2</sub>RR-HMFOR system in comparison to the conventional CO<sub>2</sub>RR-OER system: (a) energy efficiency breakdown by product on the Cu/CP cathode electrocatalyst; (b) energy efficiency breakdown by product on the Cu/PTFE cathode electrocatalyst; (c) energy efficiency of the electrolyzer with different cathode electrocatalysts paired to the OER or HMFOR via the NiP anode electrocatalyst at 100 mA cm<sup>-2</sup>; (d) half-cell potentials on the cathode and anode at 100 mA cm<sup>-2</sup> with various cathode electrocatalysts and the NiP anode electrocatalyst, blue and red dotted lines indicating thermodynamic potential window for the HER and OER.

so the only way of increasing the anode energy saving is to further lower the applied anode potential while keeping a high FE. Since the HMFOR is active after the presence of Ni<sup>3+</sup> (1.36 V vs. RHE), the theoretical maximum  $\eta_{\text{cell}}$  for the CO<sub>2</sub>RR-HMFOR system via Ni<sup>3+</sup>-based electrocatalysts will occur when the anode potential is 1.36 V, which gives 19 to 24% energy saving improvement on the current CO<sub>2</sub>RR-OER system, as shown in Table S8 (ESI).<sup>†</sup> This leads to the understanding that developing a better electrocatalyst for the HMFOR at lower applied potential is the key to increasing the energy saving for the coupled CO<sub>2</sub>RR-HMFOR system.

As shown in Fig. 10d, the distribution of applied potentials after *iR* correction demonstrates that with the HMFOR replacing the OER, there is still room to improve the applied voltage on both the cathode and the anode. Obviously, on the cathode, the thermodynamic potential depends on the type of the carbon

dioxide reduction product formed, but as the applied potential approaches the thermodynamic potential of the HER at 0 V vs. RHE (blue dotted line), it is more likely to achieve higher faradaic efficiency, hence higher energy efficiency, while suppressing hydrogen evolution. On the other hand, since the thermodynamic potential for the OER is 1.23 V (red dotted line), operating at a potential lower than 1.23 V for the anodic reaction will yield even higher energy saving in the CO<sub>2</sub>RR system. Besides the energy analysis, one should also consider the economic benefits of the HMFOR.

### Economic analysis

The economic analysis is performed to evaluate the NPV and LCP (levelized cost of production) for the process. To estimate NPV and LCP, the capital expenditure (CAPEX), the operating expenditure (OPEX), and the annual profit are evaluated. A



39.8% tax rate and 10% internal rate of return (IRR) are assumed. All cash flows/prices are calculated as of 2023. The end-of-life NPV is calculated using the following eqn (13):

$$NPV = \sum_{i=1}^n \frac{CF_i}{(1 + IRR)^i} \quad (13)$$

where  $i$  is the number of years,  $n$  is the project lifetime, and  $CF$  is the annual cash flow.

A base case scenario is set based on current experiment and literature data, with current market values. Then a pessimistic case and an optimistic case are set to compare by considering the performance in reality and possible improvement in the future, with volatility in the market. The model of the conventional CO<sub>2</sub>RR-OER system is based on techno-economic parameters as shown in Table 3, while the typical HMFOR-HER setup is modeled using parameters shown in Table 4.

**Costs of formate production.** The most economically feasible products in the CO<sub>2</sub>RR are formate/formic acid and carbon monoxide.<sup>12</sup> While some products such as ethylene and ethanol could potentially become economical to produce, the costs associated with production need to be lowered. Large scale production of products such as methane, methanol and 1-propanol, still needs to demonstrate higher selectivity to compete with cheaper production methods in a limited market size.<sup>13</sup>

The electrochemical reduction of carbon dioxide to formic acid often faces a challenge in downstream separation costs due to the relatively low concentration of the product.<sup>62</sup> To set up a simpler model, we look at potassium formate as the final

CO<sub>2</sub>RR product, which is a result of formate ions produced combining with potassium ions in the electrolyte. However, in this alkaline AEM system, the drawback is obvious – CO<sub>2</sub> consumption can be ~30% in the electrolyte forming bicarbonate.<sup>63</sup> It is therefore assumed that CO<sub>2</sub> would be reacting with KOH to produce a certain amount of bicarbonate, which is the major impurity. The removal of bicarbonate is expected in a crystallizer. The bleed from the crystallizer containing a higher concentration of potassium formate will then be for sale in the form of solution.

The selling price of potassium formate (50 wt%) is estimated to be 898 \$ per ton, with an average annual inflation of 5%.<sup>42</sup> LCP of the formate is shown to be reasonable as the current market price lies between the base case (964.49 \$ per ton) and the optimistic case (638.81 \$ per ton), closer to the base case (Fig. 11a). This means in the optimistic case, there is a larger room to adjust the selling price to account for extra costs or to generate more revenue. The economic analysis estimated for the plant is based on potassium formate production throughput of 100 tonnes per day for a project lifetime of 20 years with 90% operation factor. The CAPEX of a CO<sub>2</sub>RR-OER system considering the process areas (excluding storage warehouse, buildings, utility, site development *etc.*) is estimated to be 13 M\$ while the OPEX is much higher (62 M\$ per y) due to the electrolyte consumption (KOH, 80%), as shown in Fig. 11.

**Costs of FDCA production.** As a biomass-derived product, conventional production of FDCA is carried out through thermocatalysis and requires heating and precious metal electrocatalysts such as gold and platinum.<sup>64,65</sup> However

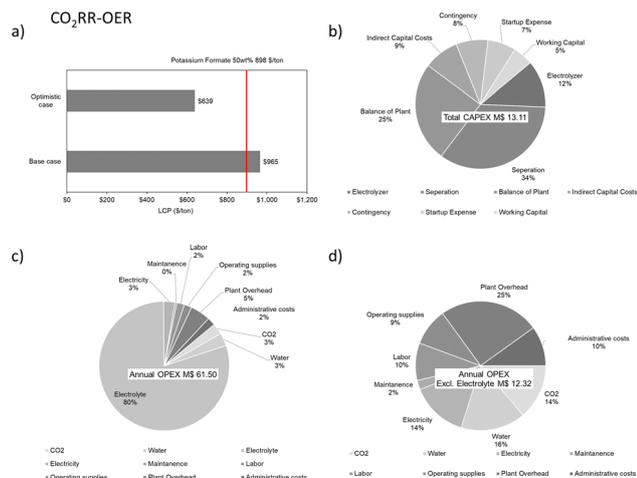
Table 3 CO<sub>2</sub>RR-OER system economic analysis parameters in 3 case scenarios

CO <sub>2</sub> RR-OER parameter	Unit	Pessimistic case	Base case	Optimistic case
Cell voltage	V	4.4	2.2	1.76
Electricity price	\$ per kW h	0.04	0.03	0.02
Electrolyzer unit cost	\$ per m <sup>2</sup>	920	460	230
Current density	A cm <sup>-2</sup>	0.05	0.1	0.15
Formate FE	%	70	80	90
CO <sub>2</sub> price	\$ per kg	0.06	0.04	0.02
Water price	\$ per kg	0.004	0.003	0.002
KOH price	\$ per kg	1.59	1.33	1.06
Single pass conversion	%	30	50	70

Table 4 HMFOR-HER system economic analysis parameters in 3 case scenarios

HMFOR-HER Parameter	Unit	Pessimistic case	Base case	Optimistic case
Cell voltage	V	4	2	1.6
Electricity price	\$ per kW h	0.04	0.03	0.02
Electrolyzer unit cost	\$ per m <sup>2</sup>	920	460	230
Current density	A cm <sup>-2</sup>	0.05	0.1	0.15
FDCA FE	%	85	90	95
HMF price	\$ per kg	1.28	1.07	0.86
Water price	\$ per kg	0.004	0.003	0.002
KOH price	\$ per kg	1.59	1.33	1.06
H <sub>2</sub> SO <sub>4</sub> price	\$ per kg	0.52	0.26	0.13





**Fig. 11** Economic analyses of the CO<sub>2</sub>RR-OER system for potassium formate production including (a) levelized cost of production of potassium formate in the base case and optimistic case; (b) total CAPEX; (c) annual OPEX and (d) annual OPEX excluding electrolyte costs.

through the electrocatalytic route, the levelized cost of FDCA is shown to be feasible in recent studies given the current market price.<sup>17,66</sup>

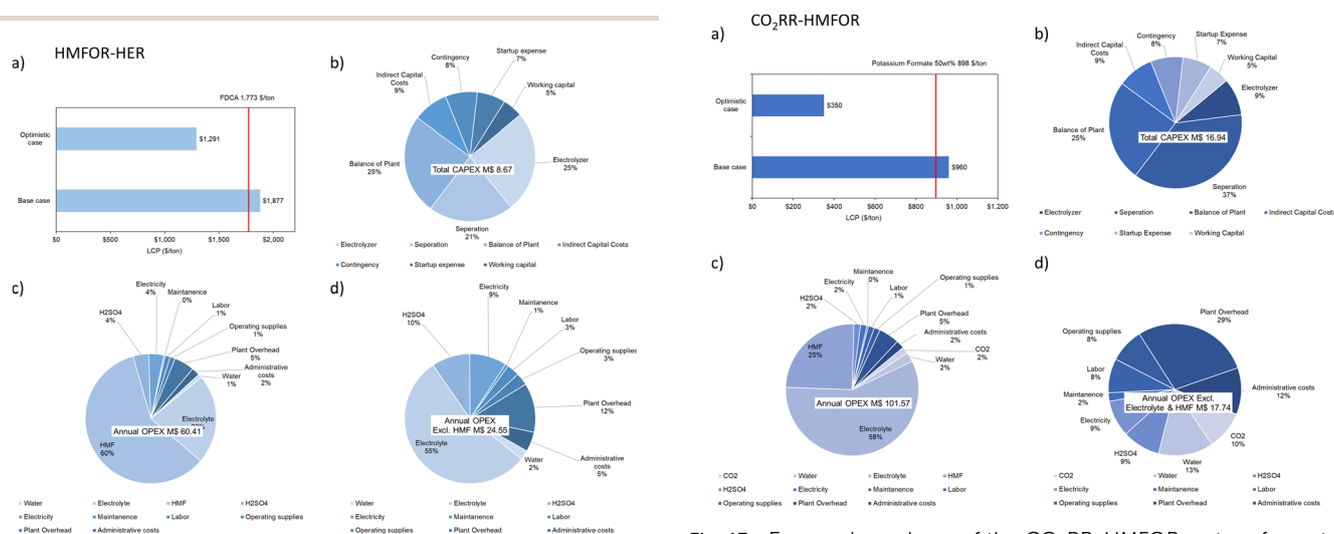
In earlier literature, the selling price of FDCA was found to be 3175 \$ per ton.<sup>67</sup> Kim *et al.* later estimated that the minimum selling price for FDCA to be 1532 \$ per ton in 2020.<sup>44</sup> Hence, we estimated the selling price to be 1773 \$ per ton in 2023 with an average annual inflation of 5%. In the optimistic case, the LCP of FDCA can be much lower, giving room to account for extra costs or to generate more revenue, as shown in Fig. 12a. The economic analysis estimated for the plant is based on FDCA production throughput of 100 tonnes per day for a project lifetime of 20 years with 90% operation factor. The CAPEX of

a HMFOR-HER system considering the process areas (excluding storage warehouse, buildings, utility, site development *etc.*) is estimated to be 9 M\$. Similar to the CO<sub>2</sub>RR-OER case, the OPEX is also much higher (60 M\$ per y) due to the feed material (HMF, 60%) and electrolyte consumption (KOH, 22%), as shown in Fig. 12.

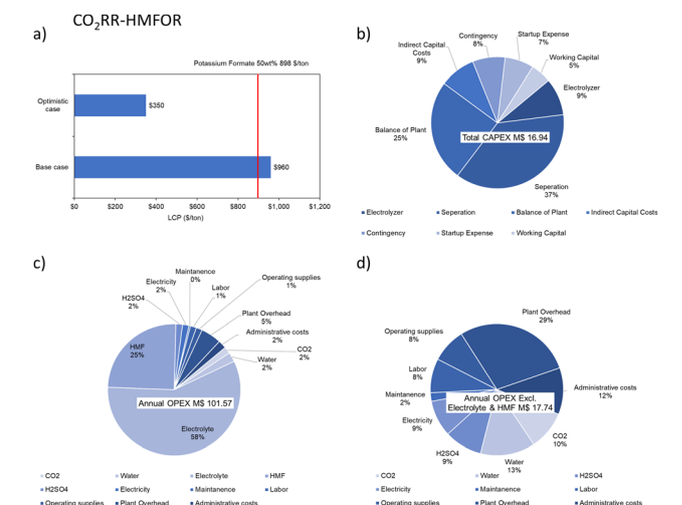
**Improved NPV on the combined system.** In this study, formate is used as an example to illustrate the economic feasibility of the CO<sub>2</sub>RR-HMFOR system. Potassium formate is modeled as an example product from the cathode reaction. Potassium formate (50 wt% to 75 wt%) has a current market price of 0.9 to 1.0 USD per kg,<sup>42</sup> and a market size of 579.25 M\$ in 2020, which is expected to grow with 5.11% CAGR till 2028.<sup>68</sup>

With the same basis on potassium formate production throughput, project lifetime and operation factor, while using the HMFOR in the anode, the economic analysis shows a general improvement. In the combined system, the LCP of the formate becomes lower in both the base case (959.73 \$ per ton) and the optimistic case (350.05 \$ per ton), giving a 55% decrease compared to the optimistic LCP (638.81 \$ per ton) from the conventional system. The CAPEX considering the process areas (excluding storage warehouse, buildings, utility, and site development, *etc.*) is estimated to be 17 M\$ while the OPEX is 102 M\$ per y due to the electrolyte consumption (KOH, 58%) and feed material (HMF, 25%), as shown in Fig. 13. This indicates a heavier front investment and higher operation costs but a greater revenue to make up the costs compared to the CO<sub>2</sub>RR-OER case.

**Sensitivity.** The sensitivity test for the conventional CO<sub>2</sub>RR-OER system and the combined CO<sub>2</sub>RR-HMFOR system is carried out to visualize the impact of changes to techno-economic parameters, as shown in Table 5. By varying the CO<sub>2</sub>RR parameters, the conventional OER-paired system shows an NPV of -M\$34 for the base case and M\$106 for the optimistic



**Fig. 12** Economic analyses of the HMFOR-HER system for FDCA production including (a) levelized cost of production of FDCA in the base case and optimistic case; (b) total CAPEX; (c) annual OPEX and (d) annual OPEX excluding HMF costs.



**Fig. 13** Economic analyses of the CO<sub>2</sub>RR-HMFOR system for potassium formate and FDCA production including (a) levelized cost of production of potassium formate in the base case and optimistic case; (b) total CAPEX; (c) annual OPEX and (d) annual OPEX excluding electrolyte and HMF costs.



Table 5 CO<sub>2</sub>RR-HMFOR system economic analysis parameters in 3 case scenarios

CO <sub>2</sub> RR-HMFOR parameter	Unit	Pessimistic case	Base case	Optimistic case
Cell voltage	V	4	2	1.6
Electricity price	\$ per kW h	0.04	0.03	0.02
Electrolyzer unit cost	\$ per m <sup>2</sup>	920	460	230
Current density	A cm <sup>-2</sup>	0.05	0.1	0.15
Formate FE	%	70	80	90
CO <sub>2</sub> price	\$ per kg	0.06	0.04	0.02
Water price	\$ per kg	0.004	0.003	0.002
KOH price	\$ per kg	1.59	1.33	1.06
Single pass conversion	%	30	50	70
HCOOK throughput	t day <sup>-1</sup>	100	100	120
FDCA FE	%	85	90	95
HMF price	\$ per kg	1.28	1.07	0.86
H <sub>2</sub> SO <sub>4</sub> price	\$ per kg	0.52	0.26	0.13
FDCA selling price	t day <sup>-1</sup>	1418	1773	2128

case, whereas the HMFOR-paired system has a better base case and optimistic case NPVs, -M\$31 and M\$120, respectively (Fig. 14). The difference is due to the production of FDCA in the latter system which can bring in additional revenue. Then for the CO<sub>2</sub>RR-HMFOR combined system, optimizing the techno-economic parameters on both reactions yields a more improved NPV, *i.e.*, M\$225 in Fig. 15.

Given the top influencers being CO<sub>2</sub> single pass conversion and the chemical prices (KOH, HMF and FDCA), the

recommended priority for improvement is optimization of the CO<sub>2</sub> half-cell as well as securing feedstock/product prices.

## Conclusions

The successful synthesis of the NiP electrocatalyst by facile electrodeposition generates Ni<sup>2+</sup>/Ni<sup>3+</sup> active sites in the catalytic surface layer characterized by SEM-EDX, XRD, XPS and XAS. We demonstrated its robust activity in the 5-HMF oxidation reaction thanks to the synergistic incorporation of phosphorus in the electrodeposited nickel catalyst, which makes it a feasible alternative to the OER in a conventional CO<sub>2</sub> reduction electrolyzer. High selectivity (>90%) and stability (>200 CV cycles) of the HMFOR are achieved in a flow cell electrolyzer paired with the cathodic CO<sub>2</sub>RR at a current density of 100 mA cm<sup>-2</sup> for the first time. The combined system is shown to simultaneously produce FDCA and value-added CO<sub>2</sub>RR products such as CO, formate, and C<sub>2+</sub> products with consistent cathode FE and a global improvement of 11–12% energy efficiency relative to the conventional OER-paired system. Through a techno-economic analysis, the combined system has shown economic benefits including lower electricity costs, an additional NPV gain from FDCA production and a potential > M\$100 NPV gain with anode optimization. Sensitivity analysis emphasizes the heavy dependence of NPV on CO<sub>2</sub> conversion and chemical prices. It is concluded that the combination of an anode reaction which simultaneously lowers the energy requirement and produces a valuable product is beneficial to the current CO<sub>2</sub>RR electrolyzer. Recommendations for the future study include electrocatalyst development, flow-cell design (potential membraneless setup), process optimization, feed composition and temperature effect.

## Author contributions

Conceptualization: RL, HY, and HZ. Data curation: RL, HY, HZ, MS, JG, and KL. Investigation: RL, HY, HZ, MS, AF, and PP. Software analysis: MS and AF. Visualization: ZG, XL, and RL.

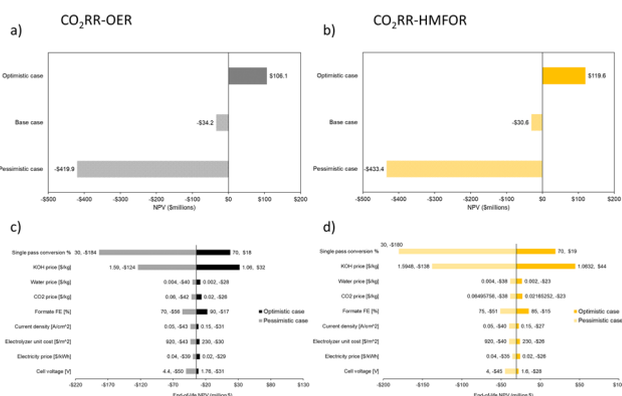


Fig. 14 Three cases of varying CO<sub>2</sub>RR parameters generate NPV results on (a) CO<sub>2</sub>RR-OER and (b) CO<sub>2</sub>RR-HMFOR; followed by (c), (d) sensitivity analyses done using the same CO<sub>2</sub>RR parameters on the two systems respectively.

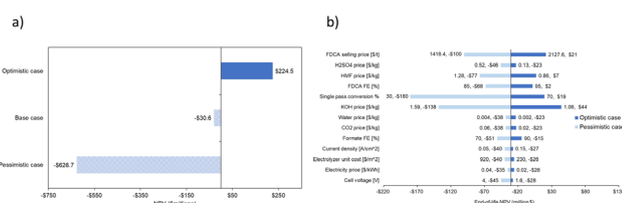


Fig. 15 Three cases of varying CO<sub>2</sub>RR and HMFOR parameters generate (a) NPV results on the CO<sub>2</sub>RR-HMFOR; followed by (b) sensitivity analyses done using the same parameters on the same system.



Writing: RL, HY, and HZ. Review & editing: PP, XW, JG, MS, XL, and AS. Supervision: AS.

## Conflicts of interest

There are no conflicts to declare.

## Acknowledgements

This work was partly supported by Fonds de recherche du Québec – Nature et technologies (FRQNT) Master Scholarship, FRQNT New Researchers Fund, Eugénie Ulmer Lamothe (EUL) Summer Scholarship, NSERC Discovery Grant, Canada Research Chair, and CFI-JELF. The synchrotron XAS was done at the Canadian Light Source (CLS) in Saskatchewan, Canada.

## References

- 1 S. Ma, Y. Lan, G. M. J. Perez, S. Moniri and P. J. A. Kenis, *ChemSusChem*, 2014, **7**, 866–874.
- 2 A. Löwe, M. Schmidt, F. Bienen, D. Kopljar, N. Wagner and E. Klemm, *ACS Sustain. Chem. Eng.*, 2021, **9**, 4213–4223.
- 3 W. Guo, S. Liu, X. Tan, R. Wu, X. Yan, C. Chen, Q. Zhu, L. Zheng, J. Ma, J. Zhang, Y. Huang, X. Sun and B. Han, *Angew Chem. Int. Ed. Engl.*, 2021, **60**, 21979–21987.
- 4 W. J. Teh, O. Piqué, Q. H. Low, W. Zhu, F. Calle-Vallejo and B. S. Yeo, *ACS Catal.*, 2021, **11**, 8467–8475.
- 5 H. Yang, Y. Wu, G. Li, Q. Lin, Q. Hu, Q. Zhang, J. Liu and C. He, *J. Am. Chem. Soc.*, 2019, **141**, 12717–12723.
- 6 X. Wang, Z. Wang, F. P. García de Arquer, C.-T. Dinh, A. Ozden, Y. C. Li, D.-H. Nam, J. Li, Y.-S. Liu, J. Wicks, Z. Chen, M. Chi, B. Chen, Y. Wang, J. Tam, J. Y. Howe, A. Proppe, P. Todorović, F. Li, T.-T. Zhuang, C. M. Gabardo, A. R. Kirmani, C. McCallum, S.-F. Hung, Y. Lum, M. Luo, Y. Min, A. Xu, C. P. O'Brien, B. Stephen, B. Sun, A. H. Ip, L. J. Richter, S. O. Kelley, D. Sinton and E. H. Sargent, *Nat. Energy*, 2020, **5**, 478–486.
- 7 J. E. Huang, F. Li, A. Ozden, A. Sedighian Rasouli, F. P. García de Arquer, S. Liu, S. Zhang, M. Luo, X. Wang, Y. Lum, Y. Xu, K. Bertens, R. K. Miao, C.-T. Dinh, D. Sinton and E. H. Sargent, *Science*, 2021, **372**, 1074–1078.
- 8 C. T. Dinh, T. Burdyny, M. G. Kibria, A. Seifitokaldani, C. M. Gabardo, F. P. G. de Arquer, A. Kiani, J. P. Edwards, P. De Luna, O. S. Bushuyev, C. Q. Zou, R. Quintero-Bermudez, Y. J. Pang, D. Sinton and E. H. Sargent, *Science*, 2018, **360**, 783–787.
- 9 F. P. García de Arquer, C.-T. Dinh, A. Ozden, J. Wicks, C. McCallum, A. R. Kirmani, D.-H. Nam, C. Gabardo, A. Seifitokaldani, X. Wang, Y. C. Li, F. Li, J. Edwards, L. J. Richter, S. J. Thorpe, D. Sinton and E. H. Sargent, *Science*, 2020, **367**, 661–666.
- 10 C. M. Gabardo, A. Seifitokaldani, J. P. Edwards, C.-T. Dinh, T. Burdyny, M. G. Kibria, C. P. O'Brien, E. H. Sargent and D. Sinton, *Energy Environ. Sci.*, 2018, **11**, 2531–2539.
- 11 C.-T. Dinh, F. P. García de Arquer, D. Sinton and E. H. Sargent, *ACS Energy Lett.*, 2018, **3**, 2835–2840.
- 12 M. Jouny, W. Luc and F. Jiao, *Ind. Eng. Chem. Res.*, 2018, **57**, 2165–2177.
- 13 M. G. Kibria, J. P. Edwards, C. M. Gabardo, C. T. Dinh, A. Seifitokaldani, D. Sinton and E. H. Sargent, *Adv. Mater.*, 2019, **31**, e1807166.
- 14 J. Qiao, Y. Liu and J. Zhang, *Electrochemical Reduction of Carbon Dioxide: Fundamentals and Technologies*, CRC Press, Taylor & Francis Group, Boca Raton, 2016.
- 15 W. Li, D. Xiong, X. Gao and L. Liu, *Chem. Commun.*, 2019, **55**, 8744–8763.
- 16 S. Verma, S. Lu and P. J. A. Kenis, *Nat. Energy*, 2019, **4**, 466–474.
- 17 J. Na, B. Seo, J. Kim, C. W. Lee, H. Lee, Y. J. Hwang, B. K. Min, D. K. Lee, H. S. Oh and U. Lee, *Nat. Commun.*, 2019, **10**, 5193.
- 18 A. Vass, A. Kormanyos, Z. Koszo, B. Endrodi and C. Janaky, *ACS Catal.*, 2022, **12**, 1037–1051.
- 19 H. Liu, D. Guo, W. Zhang and R. Cao, *J. Mater. Res.*, 2017, **33**, 568–580.
- 20 H. Chu, D. Zhang, B. Jin and M. Yang, *Appl. Catal., B*, 2019, **255**, 117744.
- 21 J. Tong, W. Ma, L. Bo, T. Li, W. Li, Y. Li and Q. Zhang, *J. Power Sources*, 2019, **441**, 227166.
- 22 Z. Gohari-Bajestani, X. Wang, A. Guet, R. Moury, J.-M. Grenèche, A. Hémon-Ribaud, Y. Zhang, D. Chartrand, V. Maisonneuve, A. Seifitokaldani, N. Kornienko and J. Lhoste, *Chem Catal.*, 2022, **2**, 1114–1127.
- 23 X. Du, L. Yin, W. Zhang, M. Zhang, K. Su and Z. Li, *J. Mater. Sci. Technol.*, 2023, **135**, 26–33.
- 24 J. Lan, J. Lin, Z. Chen and G. Yin, *ACS Catal.*, 2015, **5**, 2035–2041.
- 25 P. Hauke, T. Merzdorf, M. Klingenhof and P. Strasser, *Nat. Commun.*, 2023, **14**, 4708.
- 26 S. Barwe, J. Weidner, S. Cychy, D. M. Morales, S. Dieckhöfer, D. Hiltrop, J. Masa, M. Muhler and W. Schuhmann, *Angew. Chem., Int. Ed.*, 2018, **57**, 11460–11464.
- 27 E. de Jong, M. A. Dam, L. Sipos and G. J. M. Gruter, in *Biobased Monomers, Polymers, and Materials*, American Chemical Society, 2012, ch. 1, vol. 1105, pp. 1–13.
- 28 N. Heidary and N. Kornienko, *Chem. Commun.*, 2019, **55**, 11996–11999.
- 29 J. Bi, Q. Zhu, W. Guo, P. Li, S. Jia, J. Liu, J. Ma, J. Zhang, Z. Liu and B. Han, *ACS Sustain. Chem. Eng.*, 2022, **10**, 8043–8050.
- 30 J. Wu, Z. Kong, Y. Li, Y. Lu, P. Zhou, H. Wang, L. Xu, S. Wang and Y. Zou, *ACS Nano*, 2022, **16**, 21518–21526.
- 31 J. Guo, M. Abdinejad, A. Farzi, M. Salehi and A. Seifitokaldani, *Energy Adv.*, 2023, **2**, 877–885.
- 32 R. Latsuzbaia, R. Bisselink, A. Anastasopol, H. van der Meer, R. van Heck, M. S. Yagüe, M. Zijlstra, M. Roelands, M. Crockatt, E. Goetheer and E. Giling, *J. Appl. Electrochem.*, 2018, **48**, 611–626.
- 33 S. R. Kubota and K.-S. Choi, *Chemsuschem*, 2018, **11**, 2138–2145.
- 34 B. You, N. Jiang, X. Liu and Y. Sun, *Angew. Chem., Int. Ed.*, 2016, **55**, 9913–9917.
- 35 B. You, N. Jiang, M. Sheng, M. W. Bhushan and Y. Sun, *ACS Catal.*, 2016, **6**, 714–721.



- 36 B. You, X. Liu, N. Jiang and Y. Sun, *J. Am. Chem. Soc.*, 2016, **138**, 13639–13646.
- 37 Y. Sun, J. Wang, Y. Qi, W. Li and C. Wang, *Adv. Sci.*, 2022, **9**, e2200957.
- 38 B. Liu, S. Xu, M. Zhang, X. Li, D. Decarolis, Y. Liu, Y. Wang, E. K. Gibson, C. R. A. Catlow and K. Yan, *Green Chem.*, 2021, **23**, 4034–4043.
- 39 S. Choi, M. Balamurugan, K.-G. Lee, K. H. Cho, S. Park, H. Seo and K. T. Nam, *J. Phys. Chem. Lett.*, 2020, **11**, 2941–2948.
- 40 Y. Yang and T. Mu, *Green Chem.*, 2021, **23**, 4228–4254.
- 41 Z. Huang, R. G. Grim, J. A. Schaidle and L. Tao, *Energy Environ. Sci.*, 2021, **14**, 3664–3678.
- 42 K. Oen, *Master*, Norwegian University of Science and Technology 2017.
- 43 ChemAnalyst, *Ethylene Price Trend and Forecast*, <https://www.chemanalyst.com/Pricing-data/ethylene-40>, (accessed July 15, 2023).
- 44 H. Kim, J. Choi, J. Park and W. Won, *Green Chem.*, 2020, **22**, 7070–7079.
- 45 W. Li, L. Xin, X. Xu, Q. Liu, M. Zhang, S. Ding, M. Zhao and X. Lou, *Sci. Rep.*, 2015, **5**, 9277.
- 46 K. Wang, Y. Si, Z. Lv, T. Yu, X. Liu, G. Wang, G. Xie and L. Jiang, *Int. J. Hydrogen Energy*, 2020, **45**, 2504–2512.
- 47 S. Enzo, G. Fagherazzi, A. Benedetti and S. Polizzi, *J. Appl. Crystallogr.*, 1988, **21**, 536–542.
- 48 F. S. Zhang, J. W. Wang, J. Luo, R. R. Liu, Z. M. Zhang, C. T. He and T. B. Lu, *Chem. Sci.*, 2018, **9**, 1375–1384.
- 49 R. Lin, M. Salehi, J. Guo and A. Seifitokaldani, *iScience*, 2022, **25**, 104744.
- 50 M. Fleischmann, K. Korinek and D. Pletcher, *J. Electroanal. Chem. Interfacial Electrochem.*, 1971, **31**, 39–49.
- 51 M. S. E. Houache, K. Hughes, A. Ahmed, R. Safari, H. Liu, G. A. Botton and E. A. Baranova, *ACS Sustain. Chem. Eng.*, 2019, **7**, 14425–14434.
- 52 M. T. Bender, Y. C. Lam, S. Hammes-Schiffer and K. S. Choi, *J. Am. Chem. Soc.*, 2020, **142**, 21538–21547.
- 53 J. van Drunen, T. W. Napporn, B. Kokoh and G. Jerkiewicz, *J. Electroanal. Chem.*, 2014, **716**, 120–128.
- 54 Z. Li, X. Dou, Y. Zhao and C. Wu, *Inorg. Chem. Front.*, 2016, **3**, 1021–1027.
- 55 J. Masa, S. Barwe, C. Andronesco, I. Sinev, A. Ruff, K. Jayaramulu, K. Elumeeva, B. Konkena, B. Roldan Cuenya and W. Schuhmann, *ACS Energy Lett.*, 2016, **1**, 1192–1198.
- 56 B. Zhang, H. Fu and T. Mu, *Green Chem.*, 2022, **24**, 877–884.
- 57 N. Zhang, Y. Zou, L. Tao, W. Chen, L. Zhou, Z. Liu, B. Zhou, G. Huang, H. Lin and S. Wang, *Angew. Chem., Int. Ed.*, 2019, **58**, 15895–15903.
- 58 V. Chakarian and Y. U. Idzerda, *J. Appl. Phys.*, 1997, **81**, 4709.
- 59 P. Zhang, X. Sheng, X. Chen, Z. Fang, J. Jiang, M. Wang, F. Li, L. Fan, Y. Ren, B. Zhang, B. J. J. Timmer, M. S. G. Ahlquist and L. Sun, *Angew. Chem. Int. Ed.*, 2019, **58**, 9155–9159.
- 60 Y. Zhu, C. Cao, S. Tao, W. Chu, Z. Wu and Y. Li, *Sci. Rep.*, 2014, **4**, 5787.
- 61 T.-T. Zhuang, Z.-Q. Liang, A. Seifitokaldani, Y. Li, P. De Luna, T. Burdyny, F. Che, F. Meng, Y. Min, R. Quintero-Bermudez, C. T. Dinh, Y. Pang, M. Zhong, B. Zhang, J. Li, P.-N. Chen, X.-L. Zheng, H. Liang, W.-N. Ge, B.-J. Ye, D. Sinton, S.-H. Yu and E. H. Sargent, *Nat. Catal.*, 2018, **1**, 421–428.
- 62 C. Kim, Y. Lee, K. Kim and U. Lee, *Catalysts*, 2022, **12**.
- 63 H. Shin, K. U. Hansen and F. Jiao, *Nat. Sustain.*, 2021, **4**, 911–919.
- 64 S. Bello, P. Méndez-Trelles, E. Rodil, G. Feijoo and M. T. Moreira, *Sep. Purif. Technol.*, 2020, 233.
- 65 C. Chen, L. Wang, B. Zhu, Z. Zhou, S. I. El-Hout, J. Yang and J. Zhang, *J. Energy Chem.*, 2021, **54**, 528–554.
- 66 P. Patel, D. Schwartz, X. Wang, R. Lin, O. Ajao and A. Seifitokaldani, *ACS Sustain. Chem. Eng.*, 2022, **10**, 4206–4217.
- 67 C. Triebel, V. Nikolakis and M. Ierapetritou, *Comput. Chem. Eng.*, 2013, **52**, 26–34.
- 68 V. M. Research, *Potassium Formate Market Size and Forecast*, <https://www.verifiedmarketresearch.com/product/potassium-formate-market/>, (accessed October 23, 2022).

