Selective CO₂ reduction to C₃ and C₄ oxyhydrocarbons on nickel phosphides at overpotentials as low as 10 mV†


We introduce five nickel phosphide compounds as electro-catalysts for the reduction of carbon dioxide in aqueous solution, that achieve unprecedented selectivity to C₃ and C₄ products (the first such report). Three products: formic acid (C₁), methylglyoxal (C₃), and 2,3-furandiol (C₄), are observed at potentials as low as +50 mV vs. RHE, and at the highest half-reaction energy efficiencies reported to date for any C₁ product (99%). The maximum selectivity for 2,3-furandiol is 71% (faradaic efficiency) at 0.00 V vs. RHE on Ni₂P, which is equivalent to an overpotential of 10 mV, with the balance forming methylglyoxal, the proposed reaction intermediate. P content in the series correlates closely with both the total C products and product selectivity, establishing definitive structure–function relationships. We propose a reaction mechanism for the formation of multi-carbon products, involving hydride transfer as the potential-determining step to oxygen-bound intermediates. This unlocks a new and more energy-efficient reduction route that has only been previously observed in nickel-based enzymes. This performance contrasts with simple metallic catalysts that have poor selectivity between multi-carbon products, and which require high overpotentials (>700 mV) to achieve comparable reaction rates.

Introduction

The electrochemical reduction of carbon dioxide (CO₂) using water as hydrogen source has the potential to enable sustainable production of fuels, chemicals and polymers from renewable energy sources. While active and selective catalysts for CO₂ reduction to CO₁−₇ and HCOOH[8−10] have been developed over the past few years, the generation of high-value multi-carbon products is not yet sufficiently efficient. Copper and alloys thereof are the only catalysts proven to generate C₂ and C₃ alkanes, alcohols, ketones and...
aldehydes at significant rates. However, copper-based catalysts are still limited by three problems: (1) poor selectivity of the reaction produces a wide range of carbon products, (2) high overpotentials waste energy to heat, and (3) significant H₂ co-production competes with the desired organic compounds.

Norskov and co-workers have proposed a mechanism for the conversion of CO₂ to CH₄ on copper, based on density functional theory (DFT), which involves initial reduction to adsorbed CO (*CO), which blocks surface H-adsorption sites and suppresses the significant competing hydrogen evolution reaction (HER). Their proposed potential-determining step (PDS) occurs when HCO* binds parallel to the Cu surface to create bonding interactions to both C and O atoms. The calculated PDS requires an applied potential of ~0.74 V vs. RHE, which corresponds to the experimental onset of methane and ethylene production observed by Hori et al. Since the PDS involves the binding of HCO*, the theoretical overpotential for methane formation should scale with the CO binding energy for different metal surfaces. This descriptor is near the optimal value for copper, rationalizing its ranking as the best pure transition metal catalyst for reducing CO₂ beyond 2-electron reduction products.

Binary materials that favor binding the HCO* intermediate through both the carbon and oxygen atoms should break the scaling relationships obeyed by simple metals and could potentially improve catalytic activity. Both nickel and phosphorous allow for increased stabilization of oxygen-bound intermediates, potentially decreasing the overpotential for reaction. Additionally, they form multiple binary compounds that can absorb hydrogen atoms which have different hydride bond strength (hydricity). Moreover, the two principal enzymes that convert CO₂ to CO and subsequently couple C–C bonds, both utilize nickel in the active site. Both enzymes utilize sulfide + cyanide ligands to nickel, possibly to tune hydricity. Here, we approximate this ligand set using phosphorous which provides an iso-electronic replacement for the S + CN⁻ ligands. Nickel phosphides have been reported as highly active HER catalysts. Using them for CO₂RR is contrary to the belief that effective catalysts should have poor HER activity, yet still efficiently transfer adsorbed hydrogen atoms to a *CO intermediate. In contrast, other theoretical predictions by Rossmeisl et al. claim that having hydrogen binding energy near thermo-neutral is critical for predicting the ability of pure metals to generate products beyond CO, and is equally as important as the *CO binding energy. This represents a shift in dogma for CO₂RR research and underscores the importance of reversible hydrogen binding for both HER and CO₂RR activities.

Based on these various insights, we synthesized a family of five nickel phosphide compounds: Ni₅P, Ni₃P, Ni₁₂P₅, Ni₅P₄, and NiP₂, and evaluated their performance as electrocatalysts for CO₂RR. Our results demonstrate that product selectivity greatly improves with increasing P content in this series. This is the first report of the formation of methylglyoxal (C₃) and 2,3-furandiol (C₄) products, with potential applications in the polymer industry. The best nickel phosphide catalyst achieves essentially complete discrimination over the HER, and an energy efficiency of 99% with the lowest overpotential reported thus far for any C1 products.

Results and discussion

Catalyst crystallinity and purity

Compositional purity, crystal phase and crystal facet exposure are critical variables when comparing catalyst performance. Five different nickel phosphide compounds (Ni₅P, Ni₃P, Ni₁₂P₅, Ni₅P₄, and NiP₂) were synthesized by solid state reaction at 700 °C, in vacuum-sealed quartz tubes, using high purity elemental precursors. Comparison of the unique powder X-ray diffraction patterns to the nickel phosphide reference patterns (Fig. S1–S5, ESI†), verified that each was a single, pure phase, lacking contamination from secondary phases or amorphous material below the 2% detection limit. The nickel phosphides were intentionally synthesized at high temperature to achieve thermodynamic equilibrium among facets (i.e., polycrystallinity). Polycrystallinity was confirmed by SEM analysis, showing particles with roughly spherical morphology, lacking distinct faceting, and with sizes ranging from 1–20 µm in diameter (Fig. S6, ESI†).

Electrolysis setup

The performance of polycrystalline electrocatalysts has, to date, been limited by the ability to consistently reproduce stable catalyst/electrode interfaces from powdered catalysts supported on conductors. Our group has developed a successful protocol for preparing electrodes from nickel phosphides by mixing them with a binder and pressing them into rigid pellets. Due to the metallic nature of nickel phosphides, no addition of conductive carbon was required. To obtain electrodes with a 2 cm diameter, the different polycrystalline powders were mixed with 1% (w/w) neutral NaFion® (Sigma Aldrich 5 wt% solution in lower aliphatic alcohols and water, neutralized with 4 mg NaOH pellets per mL of solution). After grinding with a mortar and pestle until the solvent had evaporated, the mixture was transferred to an aluminum die containing an aluminum mesh for mechanical support (McMaster-Carr, 20 × 20 mesh size, 0.016” wire diameter), then pressed at 7 ton per cm². The resulting pellets were porous and had a mean thickness of 575 µm (see ESI† Fig. S7). The aluminum die was used directly as the working electrode support in a sandwich-type cell, depicting in Fig. 1. During the reaction, only the catalyst pellet was exposed to the electrolyte, and the back of the aluminum support was connected to the potentiostat. Aluminum was chosen for the support as it has been previously shown to have low activity for CO₂RR and HER.

The use of relatively large and porous electrodes can lead to substantial iR-drop and significant errors in potential determination. Resistive losses from the electrolyte were minimized by the use of a 0.5 M KHCO₃ buffer, resulting in a stable solution resistance of 6–8 ohms. Potentiostatic electrochemical impedance spectroscopy (PEIS) was performed before each experiment to measure the uncompensated resistance,
which was used for positive feedback iR compensation (Fig. S7, ESI†). The solution resistance during the reactions consistently changed by less than 0.5 ohm (<2 mV). All potentials were measured against a commercial Hg/Hg2SO4 reference electrode, and converted to the thermodynamically relevant reversible hydrogen electrode (RHE) scale.

Avoiding gaseous CO2 depletion is a concern for CO2RR.36,37 To minimize mass transport limitations, in addition to the carbonate buffer, CO2 gas was fed through the bottom of the cell via a glass frit (4–8 µm pores), generating bubbles of 50 to 150 µm (measured by optical imaging). Such small bubble sizes are sufficient to ensure CO2 saturation at operating currents lower than 10 mA cm−2, as shown in a prior study by Lobaccaro et al.37 Gas-phase products were detected by an online gas chromatograph, using both thermal conductivity and flame ionization detectors, arranged in series. The working electrode had a large surface area (3.14 cm2) to electrolyte volume (6 mL) ratio (S/V = 0.52 cm−1) to maximize the concentration of liquid phase products in the electrolyte, in accordance with recent literature recommendations.37,38 This allowed for direct product quantification by HPLC, that was further corroborated by NMR and LCMS analyses for unambiguous product assignments and yields (refer to ESI,† Fig. S12 and S13).

CO2 reduction products

Table 1 lists the reduction potentials (E0) and the number of electrons required to reduce CO2 to various products, including the three products observed in this work (formate, methylglyoxal, and 2,3-furandiol). E0 at pH 7.0 vs. RHE was calculated from tabulated39,40 thermodynamic data when available, and otherwise estimated by Mavrovouniotis’ method of individual group contributions41 (details in ESI,† Table S8). While formate is widely reported as a CO2 reduction product,3,8–10 this is the first report of the formation of methylglyoxal and 2,3-furandiol under electrochemical conditions. The E0 values reveal the latter products are thermodynamically easier to form than CO,

### Table 1 Standard electrochemical potentials at pH 7.0

<table>
<thead>
<tr>
<th>Product</th>
<th>Half-reaction</th>
<th>E0 (V vs. RHE)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hydrogen</td>
<td>2(e− + H+) ⇌ H2</td>
<td>0.00</td>
</tr>
<tr>
<td>Formic acid</td>
<td>CO2 + 2(e− + H+) ⇌ HCOOH</td>
<td>−0.02</td>
</tr>
<tr>
<td>CO</td>
<td>CO2 + 2(e− + H+) ⇌ CO</td>
<td>−0.10</td>
</tr>
<tr>
<td>Acetic acid</td>
<td>CO2 + 8(e− + H+) ⇌ H2COOH</td>
<td>+0.23</td>
</tr>
<tr>
<td>Methylglyoxal</td>
<td>3CO2 + 12(e− + H+) ⇌ C3H4O2 + 4H2O</td>
<td>+0.02</td>
</tr>
<tr>
<td>2,3-Furandiol</td>
<td>4CO2 + 14(e− + H+) ⇌ C4H4O3 + 5H2O</td>
<td>+0.01</td>
</tr>
</tbody>
</table>

Fig. 1 Scheme of the sandwich-type electrochemical cell used. The cathode is nickel phosphide supported onto a die, separated from the anode by a Nafion membrane. The counter electrode is a Pt black@platinum foil. The electrolyte is purged from the bottom with CO2 microbubbles and the headspace of the working electrode compartment is sampled by on-line gas chromatography.

Fig. 2 (A) iR-corrected linear sweep voltammetry of Ni2P at 0.5 mV s−1. In grey, argon-purged 0.5 M phosphate buffer, pH 7.5. This current corresponds solely to the HER; in blue, CO2-saturated 0.5 M KHCO3, where the current is due to CO2 reduction and HER. Binding of CO2RR intermediates partially suppresses HER. Furthermore, the current for CO2RR is seen to be larger than those attributed to HER in the phosphate buffer at low overpotentials (see inset). (B) Representative chronoamperometry measurements at different potentials for Ni2P. Due to the high porosity of the catalyst, there is an initial charging period (as previously reported in acid and base27), after which the current stabilizes. Voltammetry and chronoamperometry for all stoichiometries can be found in the ESI,† Fig. S9.
formate, and H₂, suggesting a possible approach for selectivity. To test the origin of the carbon products, isotopic labeling with \(^{13}\)CO₂ as carbon source was conducted (refer to Fig. S14 in ESI†). This confirmed that dissolved CO₂ was indeed the sole source of carbon for C\(_1\), C\(_3\), and C\(_4\) products. Control experiments using Ar-purged KHCO\(_3\) electrolyte reduced the CO₂RR currents to 20% of their previous value, confirming that dissolved CO₂, rather than ionized forms of (bi)carbonate, is the main substrate for CO₂RR on nickel phosphides.

**Current vs. potential**

Fig. 2(A) presents voltammograms for Ni\(_2\)P, obtained using Ar saturated 0.5 M sodium phosphate buffer (grey), and CO₂ saturated 0.5 M KHCO\(_3\) (blue), both at (pH 7.5). Under an argon atmosphere, the reductive current due to hydrogen evolution sharply increases with increasing overpotential. In contrast, under CO₂ saturation, the current is suppressed at all negative potentials, indicating that CO₂RR intermediates bind to some or all of the same sites that would otherwise be active for HER. Most notably, at positive potentials, the observed current increases in the presence of CO₂, indicating that CO₂RR dominates. Four of the nickel phosphides express behavior, with the exception of NiP\(_2\), which reaches open circuit potential (OCP) below 0 V vs. RHE (see ESI,† Fig. S9).

The stability of the catalyst current density was assessed by chronoamperometry, and is presented in Fig. 2(B) for Ni\(_2\)P (and for the remaining stoichiometries in the ESI,† Fig. S10).

The total current decreases in the first half hour of the experiment at all negative potentials (break-in period), due to the reduction of the surface phosphoxides, as well as the build-up of a pH gradient within the porous electrocatalyst, in agreement with our previous HER study using nickel phosphides. After the initial break-in period, the current stabilizes, and no significant loss of CO₂ current activity is observed. The total charge passed in the break-in period amounts to less than 1% of the total charge that contributes to products. To measure corrosion resistance, dissolved nickel in the solution was quantified by ICP-OES. Less than 0.023% of the nickel in the catalyst was dissolved after 2.5 hours of electrolysis (see ESI,† Table S7), equivalent to trace amounts lost during reduction of the oxidized surface.

**Selectivity vs. potential**

Fig. 3 plots the faradaic efficiency of each product as a function of potential and catalyst composition. Reduction of CO₂ to 2,3-furandiol and methylglyoxal is predominant from 0.05 V to −0.10 V vs. RHE on the more phosphorus-rich nickel phosphides (Ni\(_{12}\)P\(_5\), Ni\(_5\)P\(_4\), Ni\(_3\)P\(_4\), and NiP\(_2\)), with Ni\(_2\)P giving the highest faradaic yield at the lowest overpotential. In contrast, the low-phosphorus Ni\(_3\)P resulted in significantly less CO₂RR relative to HER and poorer selectivity, with more formic acid production than the other catalysts. The maximum selectivity of 84% for methylglyoxal was obtained on NiP\(_2\) at −0.10 V. The reaction on NiP\(_2\) was not performed at potentials more positive than −0.05 V vs. RHE because the catalyst reached OCP near 0 V, thus reducing the current and product formation below the detection limit. For 2,3-furandiol, the maximum faradaic efficiency of 71% was observed at 0 V vs. RHE on Ni\(_2\)P. Although formic acid is produced at all potentials, its faradaic efficiency never exceeds 5% for any of the catalysts. At more reductive potentials (≤−0.2 V vs. RHE), the reaction selectivity shifts to HER. This behavior is in stark contrast with what is observed on copper catalysts, where, at high overpotentials, hydrogen evolution is suppressed, and CO₂RR favored. This, along with the low overpotentials at which C–C coupling occurs, indicates that the mechanism of CO₂RR on nickel phosphides is radically different from those previously reported for simple metal catalysts.

Another important figure of merit is the CO₂RR current density that can be achieved, depicted in Fig. 4. In general,
all catalysts except Ni$_3$P show distinct profiles with peaks indicative of discrete potentials that drive CO$_2$ reduction more efficiently, albeit at different peak potentials. The maximum CO$_2$RR specific current density from NiP$_2$ is $-470$ $\mu$A cm$^{-2}$ at $-0.05$ V and a second substantial peak ($-380$ $\mu$A cm$^{-2}$) is evident at $-0.3$ V, suggestive of the population of two different electronic states. The former CO$_2$RR specific current density is twice that of polycrystalline copper for C$_3$ products at $-1.1$ V vs. RHE.$^{38}$ Only a single peak occurs on Ni$_1$P ($-330$ $\mu$A cm$^{-2}$) at $-0.40$ V vs. RHE, with currents that are ten fold lower at more positive potentials. For comparison, Ni$_3$P$_4$, which is notably the most active HER catalyst among the studied phases,$^{27,42}$ exhibits smaller CO$_2$RR currents across a broader range of potentials with peaks at $-0.4$ V ($-200$ $\mu$A cm$^{-2}$) and $+0.05$ V ($-80$ $\mu$A cm$^{-2}$). The latter peak is the highest CO$_2$RR activity among all the catalysts at this potential.

Table 2 Turnover frequency at the potential with maximum CO$_2$RR selectivity, based on electrochemical surface area

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Potential (V vs. RHE)</th>
<th>HCOO$^-$</th>
<th>MG</th>
<th>FD</th>
<th>H$_2$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni$_3$P</td>
<td>$-0.10$</td>
<td>219</td>
<td>15.3</td>
<td>24.0</td>
<td>5119</td>
</tr>
<tr>
<td>Ni$_1$P$_3$</td>
<td>0.00</td>
<td>16.4</td>
<td>201</td>
<td>175</td>
<td>1281</td>
</tr>
<tr>
<td>Ni$_2$P</td>
<td>0.00</td>
<td>14.0</td>
<td>27.4</td>
<td>127</td>
<td>0</td>
</tr>
<tr>
<td>Ni$_5$P$_4$</td>
<td>+0.05</td>
<td>14.5</td>
<td>48.5</td>
<td>30.0</td>
<td>57.3</td>
</tr>
<tr>
<td>NiP$_2$</td>
<td>$-0.10$</td>
<td>2.16</td>
<td>204</td>
<td>68.5</td>
<td>0</td>
</tr>
</tbody>
</table>

Table 3 Energy efficiency of the CO$_2$RR at the potential with maximum selectivity, considering a perfect oxygen evolution anode

<table>
<thead>
<tr>
<th>Catalyst</th>
<th>Potential (V vs. RHE)</th>
<th>CO$_2$RR energy efficiency (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Ni$_3$P</td>
<td>$-0.10$</td>
<td>8</td>
</tr>
<tr>
<td>Ni$_1$P$_3$</td>
<td>0.00</td>
<td>65</td>
</tr>
<tr>
<td>Ni$_2$P</td>
<td>0.00</td>
<td>99</td>
</tr>
<tr>
<td>Ni$_5$P$_4$</td>
<td>0.05</td>
<td>83</td>
</tr>
<tr>
<td>NiP$_2$</td>
<td>$-0.10$</td>
<td>92</td>
</tr>
</tbody>
</table>

Energy efficiency

Energy efficiency ($\varepsilon_e$) is a practical metric useful when comparing CO$_2$ reduction catalysts for energy storage applications, and is defined as the ratio of the thermoneutral potential (free energy) for each product to the applied electrical energy, eqn (1)$^{43}$

$$\varepsilon_e = \sum (E^0 \times FE/E_{cell})$$  

Table 3 gives the energy efficiency for CO$_2$RR, assuming a perfect oxygen evolution catalyst at the anode. The values range from 8 to 99%.

Fig. 5 XPS spectra of Ni$_3$P catalyst before and after CO$_2$RR (from left) C 1s, Ni 2p, and P 2p with fitted spectra. Top row is the analysis of the pristine catalyst; bottom row is the catalytically cycled material.
from 8% for Ni₅P to the maximum 99% for Ni₃P. For comparison, the energy efficiency is only 23% on polycrystalline copper. 38

**Surface changes following catalysis**

The surface stability of all catalysts was further evaluated by X-ray photoelectron spectroscopy (XPS) before and after reaction. Fig. 5 presents the experimental and fitted XPS spectra for the Ni₅P catalyst (additional XPS results are shown in ESI† Fig. S20–S24). XPS spectra are internally referenced to carbon (red peak) at a binding energy of 284.8 eV (see Fig. 5), and an additional peak (blue) from partially oxidized carbon (adventitious), which appears at the binding energy characteristic of aldehydes and terminal hydroxides. 44 Post-catalysis, the carbon peaks increase in intensity, along with the appearance of carbonate species (K 2p doublets from K₂CO₃ are also observed, see ESI†). Both carbonate and potassium binding energy shifts are also in agreement with the presence of hydrated and anhydrous K₂CO₃ (electrolyte) post-catalysis. 44,45 In the post-reduced form (see ESI† for complete XPS analysis results), the catalyst composition does not change significantly in its reaction of Ni₂P, the blue C 1s peak is shifted to a binding energy that could be attributed to aromatic carbons bound to hydroxide, such as those in 2,3-furandiol (reference for 1,2-dihydroxybenzene is shown). 44 This assignment is tentative as the peak could also be attributed to adventitious carbon that was not observed in the pristine catalyst.

The Ni 2p XPS spectra of Ni₅P (Fig. 5B and E) show the characteristic 2p₁/₂ and 2p₁/₂ doublets, each with distinct satellite peaks. The Gaussian modelling shows that three distinct chemical species are present. The species are ascribed to Ni⁰⁺ from Ni₂P and Ni²⁺ nickel hydroxide and/or oxide mixture (Ni(OH)₂/NiO), as well as Ni²⁺ from Ni₃(PO₄)₂. 44,46 This is in agreement with previous studies suggesting that nickel phosphides surface-oxidize to form a partially hydrated surface phosphate on top of the pristine nickel phosphide. 27,29 The surface phosphate layer thickness will be less than 1 nm, estimated by the probe depth of XPS in Ni(s). It should be noted that the relative content (estimated by peak height) of Ni²⁺ relative to Ni²⁺ from the combined Ni(OH)₂/NiO and Ni₃(PO₄)₂ decreases upon catalytic turnover. When the catalyst is air-exposed post-catalysis, the surface re-oxidizes. The relative change indicates that the surface nickel oxide/phosphate thickness increases when oxidation occurs in the electrolyte, compared to oxidation in air post-synthesis. The latter conditions favor the formation of a hydroxylated surface phosphate.

The P 2p XPS spectrum of Ni₅P shows two sets of doublets in the 2p₁/₂ and 2p₁/₂ regions, which are ascribed to P⁰⁻ and PO₃⁻. The ratio of P⁰⁻/PO₃⁻ is seen to decrease after catalytic turnover, indicating that the surface phosphate has a higher degree of hydration post catalysis due to exposure to the electrolyte. The atomic ratio of P⁰⁻/Ni²⁺ is ~1.9 both before and after catalytic turnover, respectively, and indicates that the catalyst composition does not change significantly in its reduced form (see ESI† for complex XPS analysis results).

After reaction, bulk changes were also evaluated by powder X-ray diffraction (ESI† Fig. S1–S5). For Ni₃P, Ni₅P₃, Ni₅P, and Ni₅P₆, no detectable crystalline impurity was formed after catalysis (<2%). However, NiP₃, the most active catalyst, originally a pure monoclinic phase, partially converts (9%) to the cubic NiP₃ phase. Additionally, four minor peaks appear that could not be assigned based on XRD.

**Reaction mechanism on nickel phosphides**

Because all three reduction products are oxygenates, the C-C coupling step presumably occurs before the two carbon–oxygen bonds in CO₂ are broken. Additionally, the predominant formation of C-C coupling products implies that key reaction intermediates are bound to the catalyst by oxygen atom(s) rather than by the carbon atom, in contrast to the proposed mechanisms on catalysts that form formate as major product. 47

Because the formation of all three products takes place at near-equilibrium potential, it is helpful to consider both thermodynamic and kinetic constraints on the possible reaction pathways to C-C coupling products. We consider the 2-electron reduction of CO₂ to formate first.

On formate-forming metals, where larger overpotentials are common, it is hypothesized that CO₂ binds through the oxygen atoms to the catalyst surface, upon the transfer of a single electron in a bent configuration followed by a proton-coupled electron transfer (PCET) to yield formate. 48 However, the equilibrium potential for the single electron transfer to form the radical anion is ~1.45 V vs. RHE in aqueous media, 49 which is prohibitive for product formation in this study. More recently, it has been suggested that on metals such as tin, the first step of CO₂ reduction to formate is PCET, 50 as opposed to the single electron transfer suggested above. However, transition metals that are believed to operate through this mechanism still require strongly reducing potentials (~0.7 to ~1.0 V vs. RHE).

Only a few catalysts are able to reduce CO₂ to HCOO⁻ at near thermoneutral potential: the formate dehydrogenase enzyme, 51 thought to operate through hydride transfer (CO₂ + H⁺ HCOO⁻) 52 and palladium-based materials, 8,10 which are also known to form active hydrides. DFT calculations of the hydrogen evolution reaction on Ni₅P, Ni₃P, and Ni₅P₂ indicate the presence of multiple types of hydride sites comprised of both Ni and P atoms at relevant potentials for HER and CO₂RR. 28,53–55 Notably, P sites are considered the most active for HER. These factors, together with the low potential at which the reaction operates, points to a hydride transfer mechanism for the initial step. This pathway is particularly favorable because the two-electron mechanism avoids the formation of high-energy radicals, both anionic CO₂⁻ and electro-neutral COOH. We next examine possible C-C coupling reactions that could generate methylglyoxal and 2,3-furandiol. Fig. 6 highlights the standard Gibbs free energy (ΔG°) of a few possible reactions, calculated from tabulated values of ΔG° of formation 59–41 (see ESI† Section S17). Standard free energy changes may guide the prediction of a suitable pathway even though the values may differ for non-standard conditions. The ΔG° for C-C coupling reactions becomes increasingly unfavorable in the sequence: reductive carboxylation of alcohols (~50 kJ mol⁻¹ for methanol) < self-condensation of aldehydes (~25 kJ mol⁻¹ for formaldehyde) < reductive CO coupling (~70 kJ mol⁻¹) < the reduction of β-ketocarboxylic acids to ketoaldehyde (~100 kJ mol⁻¹) <
carboxylation of carboxylic acids (+210 kJ mol\(^{-1}\) for acetic acid). Reductive coupling of CO units, while shown to be important for the formation of ethylene on copper at strongly reducing potentials,\(^\text{48}\) is unlikely on nickel phosphides at low applied overpotentials, as the catalyst is highly oxophilic and selectively generates formate, not carbon monoxide. It should be noted that while the reductive carboxylation of methanol is highly exergonic, alcohols are kinetically very unreactive.\(^\text{56}\) Therefore, the most energetically favored pathway for carbon–carbon coupling, under mild conditions in a bicarbonate buffer, is aldehyde self-condensation.

The literature on formaldehyde self-condensation to form trioses and tetroses suggests that the reaction is catalyzed by Lewis acids in the presence of water.\(^\text{57}\) Binding of the carbonyl group of formaldehyde to a Lewis acid significantly lowers the barrier for proton abstraction from the C–H bond of formaldehyde, allowing C–C bond formation and producing glycolaldehyde. Nickel phosphides have Lewis acid character due to the partial positive charge on the nickel atoms, as shown in the XPS measurements (see Fig. 5B and E), and could catalyze this aldehyde condensation. CO\(_2\) itself can also catalyze this condensation via carbonylation of nucleophilic oxides and phosphides.

One significant finding is that acetate is not formed, despite being thermodynamically favored (Table 1). This supports the aldehyde condensation pathway proposed, since forming C\(_3\) products is both kinetically and thermodynamically favored.

Based on these steps, we propose that CO\(_2\) reduction on nickel phosphides proceeds through the mechanism depicted in Fig. 7. In step 1, CO\(_2\) inserts into a surface hydride bond to generate an adsorbed formate species, \(\ast\text{HCOO}\). This is believed to be the potential-determining step (PDS) because the Tafel slopes for all three observed products are roughly the same (see ESI,\(^\dagger\) Fig. S19). We note that all three products are preceded by hydride exchange reactions with the surface, in steps 1, 2, and 10, and thus, the PDS for each product may be similar although chemically distinct steps. In step 2, formate is protonated and attacked by a second hydride, forming formaldehyde (H\(_2\)CO\(_*\)) upon elimination of hydroxide. Although formaldehyde is not detected, it is highly reactive and presumably surface-bound to nucleophilic phosphide, whereupon two successive, energetically favored, aldehyde self-condensation reactions occur to generate glyceraldehyde. Step 6, the keto–enol tautomerization of an unactivated methyl group, is predicted to have the highest energy barrier, and thus accounts for the accumulation of the methylglyoxal precursor. This step is followed by another energetically favorable self-condensation of aldehyde with formaldehyde on the catalyst. The cyclization in step 8 forms a more stable five-membered ring by intramolecular condensation of an alcohol and an aldehyde. The hydride abstraction in step 10, the terminal product-forming reaction, is driven by the stability of the aromatic furan ring. There is precedent in literature for the hydride abstraction by nickel phosphides, as this is believed to be the mechanism for the thermally activated hydrodeoxygenation reaction that they are
known to catalyze. The proposed mechanism was validated by reduction of selected intermediates (formate, formaldehyde, methylglyoxal) as individual starting reagents in the absence of CO₂. In all cases, the resulting product distribution matched the expected end products in precisely the same stoichiometries observed when starting from CO₂ (refer to ESI† Table S10).

The foregoing mechanism may account for the observed preference for P-rich nickel phosphides in forming C₃ and C₄ products, as these contain more of the nucleophilic P sites for binding both CO₂ and reactive hydride formation, the kind that exhibit nearly thermoneutral binding energy. Such sites are favored to undergo CO₂ addition in the initial PDS, step 1. Surface reconstruction may contribute to the formation of additional P adatoms. In particular, the theoretically predicted reconstruction of Ni₂P[001] produces a P-rich termination that is calculated to be highly nucleophilic.

Conclusions

This study demonstrates for the first time the use of transition metal phosphides for CO₂ reduction. Transition metal phosphides are the first class of materials, other than enzymes, that are able to convert CO₂ to C₃ and C₄ products in aqueous media at a near-thermoneutral potential with high selectivity, making them the best available electrocatalysts for forming >C₂ products. Copper is the only other non-biological catalyst that is able to produce multicarbon products with more than 1% faradaic efficiency. Five different nickel phosphide compounds examined here exceed this value, with NiP₂ the largest at 100%. When the kinetically facile HER reaction is discriminated against by using low overpotentials, the lowest energy C₄ products appear. A strong structure-selectivity relationship favoring higher MW C₄ products emerges among the five nickel phosphide catalysts as P content increases (NiP₂ most selective and Ni₃P least selective). Likewise, a strong structure–activity relationship between the integrated current producing C₃ products and P content emerges. Each catalyst exhibits a different current–potential profile to form C₄ products with distinct peaks. This is indicative of the population of discrete electronic states that form the key intermediates which produce these products. These relationships differ dramatically from pure metallic electrodes, notably copper. This study proposes a reaction pathway for the energy-efficient synthesis of multi-carbon chemicals from CO₂, via formate and formaldehyde intermediates, without the carbon monoxide intermediate formed when using pure metallic electrodes. Future work will focus on expansion of the mechanistic understanding of this reaction, as well as electrode engineering and catalyst development to improve current densities to industrially relevant values.

Experimental

Catalyst synthesis

Nickel metal powder (Sigma Aldrich, 99.99%, <150 μm) was mixed with stoichiometric amounts, plus 1.5% molar excess, of red phosphorus (Alfa Aesar, 98.9%, 100 mesh). The powders were ground with an agate mortar and pestle for 10 min, transferred to a quartz tube, then flushed with argon and evacuated to less than 100 mTorr three times. The evacuated quartz tubes contained batches of ~5 grams of sample, which were sealed and heated at a rate of 0.5 °C min⁻¹ stepwise (350 °C, 450 °C, and 550 °C) to 700 °C. The temperature was maintained for 6 hours at each intermediate step, and 24 h at the final temperature (to avoid hotspot formation due to the exothermic reaction). The powders were then analyzed by PXRD and, if not phase-pure, excess phosphorus or nickel was added and the procedure repeated as many times as necessary. The synthesized Ni₃P contained excess metallic nickel, which was removed by stirring with 10% HCl under nitrogen for 12 hours, and by washing with copious amounts of water. The acid wash was repeated as many times as necessary for complete removal of Ni, verified by PXRD.

Powder X-ray diffraction

Powder X-ray diffraction was conducted at room temperature on a Philips Xpert system, spinning at 100 rpm, in a Bragg-Brentano geometry, Cu K-alpha 0.15418 nm, calibrated daily with a Si standard. The step size used for the diffraction patterns was 0.02°, and the scan speed was 0.013° s⁻¹. The sample holder was 3 mm deep and ½” in diameter.

Electrochemistry

Each CO₂RR faradaic efficiency value reflects the average of at least 3 replicates. The standard deviation between HPLC measurements was smaller than 2%. The cell was used was a custom-made glass-reinforced nylon-6,6 electrochemical cell, with silicon O-rings and PEEK fittings (IDEX HS). The working electrode was separated from the counter electrode by a Nafion 115 membrane (Fuel Cell Store). Platinum black deposited on Pt foil (Alfa Aesar, 99.9%) was utilized as the counter electrode. The Hach Hg/Hg₂SO₄ reference electrode was calibrated daily against a pristine Accumet SCE electrode. This SCE was periodically calibrated against a freshly flame-annealed Pt electrode in 0.5 M H₂SO₄ under 1 atm H₂ to calibrate to the RHE scale. The working electrode was prepared by mixing 1.400 g of the catalyst with 1% (w/w) neutralized Nafion suspension and was then pressed at 22 ton onto an aluminum die. The die, containing the pressed catalyst pellet was employed directly as the working electrode and current collector, with only the nickel phosphide exposed to the electrolyte. Aluminum was selected as a support because it is inert for CO₂RR. CO₂ (Air Gas, instrument grade, with a Supelco hydrocarbon trap) was supplied through the bottom of the cell to both the working and counter electrodes at a flow rate of 5 sccm (certified MKS P4B mass flow controllers). The headspace of the working electrode compartment was sampled every 30 minutes for gas chromatography.

Electrochemical measurements were performed with a Gamry 5000E potentiostat. Before each electrolysis, the electrolyte (0.5 M KHCO₃, Chelex treated) was pre-saturated with CO₂ for at least an hour. Then, a chromatograph was taken to ensure that no air was present in the headspace. An electrochemical
impedance spectrum at the open circuit from 1 Hz to 1 MHz was taken to find out the uncompensated resistance (typically between 6 and 11 ohm). Chronoamperometry was then performed for 3 hours with positive feedback IR compensation. Between experiments, the electrochemical cell was rinsed with Millipore water and the working electrode catalyst pellet was lightly polished with a fine-grit silicon carbide polishing pad (BASI) before being re-used for multiple experiments at all potentials. In doing this, the longevity of the electrodes was confirmed, with no significant difference in product distribution observed as the electrodes were re-used. Additional replicas were made using fresh electrodes at all potentials to ensure that the product distribution was not affected across the investigated potential region.

Gas chromatography
Detection and quantification of possible headspace products (hydrogen, carbon monoxide, carbon dioxide, methane, ethane, and ethylene) was performed by an auto-sampling online HP 5890 Series II GC, with a 500 µL sample loop. The GC was fitted with a 6' packed HayeSep D, and a 6' packed MoleSieve 13× column, with thermal conductivity and flame ionization detectors connected in series. Samples were taken before reaction to check for air presence, and then every 30 minutes thereafter. Calibration curves were constructed from certified gas standards (Gasco) by CO2 dilution using mass flow controllers (MFCs). The hydrogen calibration was done with in situ generated gas through electrolysis of water on platinum, under argon (supplied by an MFC), and diluted post-reaction with CO2.

High-performance liquid chromatography (UV/RID)
Liquid products were identified and quantified on a Perkin-Elmer Flexar HPLC equipped with an auto-sampler, refractive index (RID) and UV-vis detector. An HPX 87H Aminex column, with thermal conductivity and flame ionization detectors connected in series. Samples were taken before reaction to check for air presence, and then every 30 minutes thereafter. Calibration curves were constructed from certified gas standards (Gasco) by CO2 dilution using mass flow controllers (MFCs). The hydrogen calibration was done with in situ generated gas through electrolysis of water on platinum, under argon (supplied by an MFC), and diluted post-reaction with CO2.

Conflicts of interest
There are no conflicts to declare.

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